

F-segments of Arabidopsis dehydrins show cryoprotective activities for lactate dehydrogenase depending on the hydrophobic residues

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1	ORIGINAL PAPER
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3	Title
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26	
27	Abstract
28	Although dehydrins show cryoprotective activities for freeze-sensitive enzymes, the underlying
29	mechanism is still under investigation. Here, we report that F-segments conserved in some dehydrins
30	cryoprotected lactate dehydrogenase (LDH) as well as K-segments, which were previously identified

31 as cryoprotective segments of dehydrins. The cryoprotective activity levels of four F-segments of

32Arabidopsis dehydrins were similar to that of a typical K-segment. Amino acid substitution experiments indicated that the activity of the F-segment of Arabidopsis COR47 (designated as Fseg) 33 34depended on the hydrophobic residues (L, F, and V). Intriguingly, when all the amino acids other than the hydrophobic residues were changed to glycine, the cryoprotective activity did not change, 35suggesting that the hydrophobic amino acids were sufficient for Fseg activity. Circular dichroism 36 37 analysis indicated that Fseg was mainly disordered in aqueous solution as well as Fseg  $\Phi/T$ , in 38which the hydrophobic residues of Fseg were changed to T. This suggested that the hydrophobic 39interaction might be related to the cryoprotective activities of Fseg.

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#### 41

42 Keywords Arabidopsis thaliana (Brassicaceae); protein function; late embryogenesis abundant
 43 (LEA) proteins; dehydrins

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### 46 **1. Introduction**

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48When plants are exposed to stresses, a series of late embryogenesis abundant (LEA) proteins are 49expressed (Hundertmark and Hincha, 2008; Hand et al., 2011). Expression of dehydrins, which are 50group 2 LEA proteins, is a major response to abiotic stresses such as cold, drought, and high salinity (Eriksson and Harryson, 2011; Graether and Boddington, 2014; Banerjee and Roychoudhury, 2016) 5152as well as to biotic stresses (Hanin et al., 2011). Because the amino acid sequences of dehydrins are 53dissimilar to those of any previously identified proteins, functional studies of them are still under 54way; nevertheless it has been believed that dehydrins may function as protectants for stressed plants 55according to the results of transgenic studies (Banerjee and Roychoudhury, 2016). Dehydrins are 56detected in various tissues and subcellular compartments, such as cytoplasm, nucleus, plastid, 57mitochondrion, endoplasmic reticulum, and plasma membrane (reviews cited above). The ubiquitous 58distribution of dehydrins indicates that they protect whole cells from damage caused by stresses.

Dehydrins are classified as intrinsically disordered proteins, since the proportion of hydrophilic residues is far greater than that of hydrophobic residues in the sequences. The flexibility of dehydrins has been observed with various analytical methods such as circular dichroism (CD), Fourier-transform infrared spectroscopy (FTIR), and nuclear magnetic resonance (NMR) (Findlater and Graether, 2009; Rahman et al., 2011; Ágoston et al., 2011; Clarke et al., 2015). Dehydrins have conserved sequences, e.g., K-, Y-, and S-segments (Close, 1997). K-segments (e.g., EKKGIMEKIKEKLPG) are essential for identifying dehydrins. Y-segments (e.g., DEYGNP) and 66 S-segments (e.g., LHRSGSSSSSSEDD) are frequently found in some dehydrins. By using the 67 shorthand notation of K, Y, and S, dehydrins are conventionally classified as SKn, FSKn, KnS, Kn, 68 YnSKn, YnKn, etc. Besides them,  $\Phi$ -segments, which are rich in G and polar amino acids (Close, 69 1997), PK (polylysine)-segments (Hara et al., 2009), and F-segments (DRGLFDFLGKK or extended 70ones) (Strimbeck, 2017; Wei et al., 2019) have been proposed. Many studies have reported that such 71segments had corresponding functions in vitro. K-segments showed cryoprotective activities for 72freeze-sensitive enzymes such as lactate dehydrogenase (LDH, EC 1.1.1.27) (Hughes et al., 2013; 73Hara et al., 2017). K-segments are also involved in the binding to negatively charged lipids (Koag et 74al., 2009). The membrane binding of dehydrins might maintain the fluidity of membranes at low 75temperature (Eriksson et al., 2011; Clarke et al. 2015). S-segments are likely Ca<sup>2+</sup> binding sites of 76dehydrins when the segments are phosphorylated (Alsheikh et al., 2003). The polar  $\Phi$ -segments 77might be involved in the great flexibility of dehydrins. The PK-segment and K-rich area of dehydrins 78are related to the DNA binding (Hara et al., 2009; Boddington and Graether, 2019). Although 79specific segments have not been identified, the H residues and H-rich regions were proposed to bind 80 to transition metals (Hara et al., 2005), reduce the generation of reactive oxygen species (Hara et al., 81 2013), and regulate the lipid binding of dehydrins (Eriksson et al., 2011). However, the functions of 82 F-segments, which are well conserved in cold-responsive FSKn dehydrins, are still unknown.

83 Since cold is a general environmental cue for dehydrin expression in plants, the physiological 84 roles of dehydrins in cold-stressed plants have been intensively studied. Although diverse transgenic 85 experiments revealed that the high expression levels of dehydrin genes enhanced the cold tolerance 86 of plants (e.g., Hara et al., 2003; Puhakainen et al., 2004; Peng et al., 2008; Xing et al., 2011; 87 Ochoa-Alfaro et al., 2012; Zhang et al., 2018), how dehydrins act during the establishment of cold 88 tolerance in planta has not been confirmed but is now under investigation. Many studies have 89 investigated the cryoprotection of freeze-sensitive enzymes by dehydrins (e.g., Hara et al., 2001; 90 Bravo et al., 2003; Hughes and Graether, 2011; Drira et al., 2013). It was demonstrated that size is 91 one of the most important factors in the cryoprotective activities of dehydrins (Hughes et al. 2013). 92In addition, truncation experiments indicated that K-segments of ERD10, RcDhn5, TaDHN-5, and 93WZY2 were necessary to exhibit the full inhibitory activities of the dehydrins for the cold 94denaturation of LDH (Reyes et al., 2008; Drira et al., 2013; Yang et al., 2015). As described above, 95K-segments even alone inhibited freezing damage to LDH, suggesting that K-segments might be 96 related to the cryoprotective activities of dehydrins. However, no cryoprotective sites other than 97K-segments have been determined in dehydrins.

Recently, it was reported that when the hydrophobic amino acids of a K-segment were changed to polar uncharged Ts, the cryoprotective activity of the K-segment was remarkably reduced (Hara et al., 2017). In this case, at least three hydrophobic residues were needed for the cryoprotective activity. This supported the idea that the segments containing multiple hydrophobic residues might 102 show cryoprotective activities. From this point of view, we predicted that F-segments may have 103 cryoprotective activities like K-segments, because F-segments possessed hydrophobic core regions 104 (e.g., LFDFL) in their sequences. Here we found that four F-segments of *Arabidopsis* FSKn 105 dehydrins showed potent cryoprotective activities for LDH and that the segment's hydrophobic 106 amino acids were necessary for the activities. The putative cryoprotective mechanisms of 107 F-segments were discussed on the basis of hydrophobic interaction.

- 108 109
- 110 **2. Results**
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112 2.1. Cryoprotective activities of F-segments

F-segments are conserved in FSKn dehydrins. *Arabidopsis* possesses four FSKn dehydrin genes, i.e., *At1g20440* (*COR47*, FSK3), *At1g20450* (*ERD10*, FSK3), *At1g76180* (*ERD14*, FSK2), and *At4g38410* (FSK2), all of which are expressed by cold stress (Supplemental Fig. 1). There was one F-segment per FSKn dehydrin (Strimbeck, 2017). Four F-segments, COR47Fseg, ERD10Fseg, ERD14Fseg, and At4g38410Fseg, were tested (Fig. 1A). The sequence lengths were adjusted to 15 amino acids because we compared their cryoprotective activities to the positive standard: the typical K-segment (TypK) possessing 15 amino acids (Hara et al., 2017).

121To estimate the cryoprotective activities, we measured the inhibition of both cryoinactivation and 122cryodenaturation for LDH (Fig. 1B, C). LDH has been widely used as a model enzyme for 123cryoprotection studies because it is sensitive to low temperature. Cryoinactivation was determined 124by the enzymatic activities of LDH before and after the freeze and thaw treatments (Supplemental 125Fig. 2A, B). Cryodenaturation was analyzed by monitoring the hydrophobicity of LDH during the 126freeze and thaw cycles by 8-anilino-1-naphthalene sulfonic acid (ANS), which is a detector of the 127surface hydrophobicity of proteins (Supplemental Fig. 2C). The F-segment peptides effectively 128inhibited the cryoinactivation and cryodenaturation of LDH by the same magnitude as TypK (Fig. 1291B, C). The activity levels were similar to each other. This demonstrated that the F-segment peptides 130had efficient cryoprotective activities for LDH. The following experiments were performed by using 131COR47Fseg because its sequence was identical to that of the F-segment defined in the previous 132report (Strimbeck, 2017). COR47Fseg is designated simply as Fseg in the text below.

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134 2.2. Role of hydrophobic residues

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We previously reported that the cryoprotective activity of TypK for LDH depended on the hydrophobic residues in the TypK sequence (Hara et al., 2017). This implies that, also in the case of 138Fseg, hydrophobic residues may be crucial for cryoprotective activity. To test this, we examined a 139series of Fseg-altered peptides with various amino acid substitutions (Fig. 2). A hydrophobic 140 residues-to-T substitution peptide (Fseg  $\Phi/T$ ), a negatively charged residues-to-T substitution 141 peptide (Fseg DE/T), and a positively charged residues-to-T substitution peptide (Fseg KR/T) were 142prepared. The cryoprotective analyses indicated that Fseg  $\Phi/T$  totally lost its inhibition of LDH 143cryoinactivation and cryodenaturation, but Fseg DE/T and Fseg KR/T still efficiently inhibited 144 cryoinactivation and cryodenaturation, to the same degree as Fseg. When hydrophobic residues were 145changed to E (Fseg  $\Phi/E$ ) and K (FSeg  $\Phi/K$ ), their inhibition activities diminished. These results 146 suggested that the hydrophobic residues played an important role in the cryoprotective activity of 147Fseg.

148Since it was revealed that hydrophobic residues of Fseg were necessary for cryoprotective activity, 149we changed the other amino acids to G (pG Fseg  $\Phi$ ) in order to clarify whether the hydrophobic 150residues were sufficient for the activity (Fig. 3). Surprisingly, the pG Fseg  $\Phi$  peptide showed 151similar activity to Fseg in the inhibition of both LDH cryoinactivation and cryodenaturation. In 152addition, the pG Fseg  $\Phi$ -related peptides, which contained negatively charged residues 153(pG Fseg  $\Phi DE$ ) or positively charged residues (pG Fseg  $\Phi KR$ ), also showed apparent activities. 154On the other hand, altered peptides in which G was substituted for amino acids other than charged 155residues (pG Fseg DE, pG Fseg KR, and pG Fseg DEKR) showed lower activity levels. The 156present results demonstrated that the hydrophobic residues were necessary and sufficient to exhibit 157the cryoprotective activity of Fseg.

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159 2.3. Structural features of Fseg

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161 In order to elucidate the mode of action underlying the cryoprotective activity of Fseg, we 162 compared the secondary structures between Fseg (the active peptide) and Fseg\_ $\Phi/T$  (the inactive 163 one) by using CD. We collected CD data on the solutions of Fseg and Fseg\_ $\Phi/T$  in 10 mM Tris 164 buffer pH 7.5 (Fig. 4A). Both peptides showed typical disordered states, which are expressed by 165 very negative values around 200 nm, whereas the sign of disorder in Fseg was weaker than that in 166 Fseg  $\Phi/T$ .

167 It has been documented that K-segments in solution showed a disorder-to-helix transition by the 168 addition of sodium dodecyl sulfate (SDS), and this transition may be related to the molecular 169 function of dehydrins (Koag et al., 2009). When we added SDS to the Fseg and Fseg\_ΦT solutions, 170 little transition was observed except that the  $\alpha$ -helix and  $\beta$ -strand in Fseg were increased slightly but 171 significantly by the addition of a high concentration of SDS (10 mM) (Fig. 4A, B). This suggested 172 that the cryoprotective activity of Fseg was not correlated with the capability of a structural 173 transition induced by SDS. Moreover, it is likely that Fseg did not interact with the membrane.

- 174We obtained more information on the structural difference between Fseg and Fseg  $\Phi$ T by using 175de novo structure prediction system PEP-FOLD3 (Lamiable et al., the 2016) 176(http://bioserv.rpbs.univ-paris-diderot.fr/services/PEP-FOLD3/). Because this system was designed 177for the structured peptides, we have to consider that the images represented one of the structures that 178the peptides could have. The results showed that the 3D structures were clearly different between 179Fseg and Fseg  $\Phi$ T. Fseg had a hydrophobic space in the middle of the structure (Supplemental Fig. 180 3A). The four hydrophobic amino acids ( $L_4$ ,  $F_5$ ,  $F_7$ , and  $L_8$ ) each formed a hydrophobic area that was 181 exposed to the solvent. On the other hand, Fseg  $\Phi T$  did not have any hydrophobic area and the 182whole structure was disordered for the most part (Supplemental Fig. 3B). We also built predictive 183 structures of other cryoprotective peptides used in this study, i.e., three F-segments (ERD10Fseg, 184ERD14Fseg, and At4g38410Fseg), pG Fseg  $\Phi$ , and TypK (Supplemental Fig. 3C), and found that 185all of them possessed hydrophobic spaces as Fseg did. Intriguingly, their hydrophobic areas were 186 made by four hydrophobic amino acids and were exposed to the solvent, even though their amino 187acid sequences were different.
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### 190 **3. Discussion**

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192Although the cryoprotective activities of K-segments have been well documented, it has not been 193 investigated whether other segments also show such activities. Previous studies have reported that 194 FSKn-type dehydrin COR47 was related to the cold tolerance of Arabidopsis. The COR47 gene was 195expressed and the protein was accumulated in the whole Arabidopsis plant with a specific response 196 to cold stress (Nylander et al., 2001). Overexpression of the COR47 gene enhanced the freezing 197tolerance of Arabidopsis by coexpression with the RAB18 gene (Puhakainen et al., 2004). 198OpsDHN1, a cold-responsive FSKn-type dehydrin of Opuntia streptacantha (Ochoa-Alfaro et al., 199 2012), effectively protected LDH during the freeze and thaw process (Hernández-Sánchez et al., 2002014). Recently, FSKn-type-specific conserved segments, i.e., F-segments, have been identified 201(Strimbeck, 2017; Wei et al., 2019). The sequence conservation of F-segments was higher than that 202of K-segments in the COR47 orthologues from angiosperms (Supplemental Fig. 4), implying that 203F-segments have been conserved during evolution, probably due to their important functions. Here 204we report that the F-segments of four FSKn-type dehydrins showed the LDH cryoprotective activity 205of the same potency as K-segments (Fig. 1).

A previous report demonstrated that the cryoprotective activity of a K-segment depended on the hydrophobic residues of the peptide (Hara et al., 2017). Therefore, we tested whether the hydrophobic residues of Fseg contributed to the cryoprotective activities. The experiments with amino acid substitutions indicated that hydrophobic amino acids were the determining factors for the 210 cryoprotective activity of Fseg. Since the changes in positively charged amino acids to T somewhat 211 but significantly reduced cryoprotective activities (Fig. 2B, C), the positively charged amino acids

212 might be partially related to those activities.

It was confirmed that amino acid length is a major determinant of the cryoprotective activities of dehydrins (Hughes et al. 2013). Moreover, the altered dehydrins, whose sequences were scrambled, exhibited considerable activities (Palmer et al. 2019). On the other hand, K-segments alone showed apparent cryoprotective activities and hydrophobic amino acids of K-segments were related to those activities. Taken together these findings indicate that, although the size of the disordered region is essentially important for cryoprotective activity, the amino acid sequences of segments may also be responsible for the activities of dehydrins.

220Here, the discussion moves to the mechanism underlying the LDH cryoprotective activities of 221F-segments. In general, a molecular shield model (many reviews, e.g., Chakrabortee et al., 2012) has 222been proposed to explain how dehydrins show cryoprotective activity (Hughes et al., 2013). The 223aggregation of target proteins can be inhibited by the molecular shields of dehydrins with large 224hydrodynamic radii due to the intrinsically disordered nature of those dehydrins. F-segments might 225also function as molecular shields, because Fseg was in the disordered state determined by CD. 226However, it is still unknown why hydrophobic amino acids were necessary for the cryoprotective 227 activity of Fseg. As shown in Supplemental Fig. 3, Fseg was postulated to have a hydrophobic area 228that was exposed to the solvent. Similar hydrophobic areas were found in not only other F-segments 229but also pG Fseg  $\Phi$  and TypK, all of which showed cryoprotective activities, suggesting that such 230hydrophobic areas might be crucial for cryoprotective peptides. In line with this idea, a scheme 231describing the cryoprotective mechanism of Fseg is shown in Fig. 5. During the freeze and thaw 232process, freeze-sensitive enzymes such as LDH lose their activities by aggregating together through 233the hydrophobic patches on protein surfaces (Zhang et al., 2011) (Fig. 5A). The hydrophobic area of 234Fseg might intervene in the hydrophobic interaction between target proteins. Since the hydrophobic 235area of Fseg was not structurally complemented with the hydrophobic patches of proteins, the 236interference by hydrophobic interaction could be transient. While Fseg inhibited aggregation, 237hydrophobic hydration (Davis et al., 2012) might cover the patches of protein surfaces. Thus, LDH 238is successfully kept active after the freeze and thaw process (Fig. 5B). It has been hypothesized that 239group 4 LEA proteins and chaperones might interact with the target proteins (Cuevas-Velazquez et al. 2402017; Tompa and Csermely, 2004), suggesting that the transient hydrophobic interactions might 241function also in the case of the group 4 LEA proteins and the chaperones. On the other hand, 242Fseg  $\Phi/T$ , which does not possess a hydrophobic area, cannot efficiently intervene in the 243hydrophobic interaction between the proteins. As a result, Fseg  $\Phi/T$  might be excluded from the 244hydrophobic space, after which the proteins would be aggregated due to the formation of water-tight 245seals between the proteins (Fig. 5C).

246Since the hydrophobic areas of F-segments consisted of L(M)FDFL, the sequences were certainly 247thought to be the cores of the cryoprotective activities. It is noteworthy that the sequence of 248L(M)FDFL was similar to the FXF motifs (e.g., the FNFL segment) related to the hydrophobic 249interaction between mitogen-activated protein kinases and their partner proteins (Mace et al., 2013; 250Liu et al., 2016). This implies that L(M)FDFL can affect the hydrophobic interactions between 251proteins. Moreover, it is interesting that the configurations of hydrophobic areas were similar among 252the cryoprotective peptides (Supplemental Fig. 3). In the case of F-segments, two Fs were positioned 253like spreading arms and two Ls (or an L and an M) were located between the F arms. This formed 254horizontally long hydrophobic areas. On the other hand, TypK was postulated to form a helix-like 255structure. The four hydrophobic amino acids that were located on one side of the helix made a long 256horizontal hydrophobic region, as did the F-segments. Moreover, exposure of the hydrophobic areas 257to the solvent may enhance accessibility to hydrophobic patches on the surface of proteins. This may 258facilitate the transient intervention for the hydrophobic interaction between proteins. It is suggested 259that if the peptides were all the same size, the cryoprotective activities were determined mainly by 260the configurations of the hydrophobic areas rather than by the primary and secondary structures of 261the peptides.

In conclusion, we found that F-segments had cryoprotective activities as much as K-segments did. This indicates that K-segments and F-segments may contribute to the cryoprotective activities of cold-responsive FSKn-type dehydrins. Although dehydrins possess small amounts of hydrophobic amino acids, the hydrophobic residues were key factors in the unique functions of dehydrins such as protein cryoprotection. Further studies on the functions of dehydrins may provide useful information on the physiological mechanisms underlying cold tolerance in plants and for the development of high-performance cryoprotective agents.

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# **4. Experimental**

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# 272 *4.1. Chemicals*

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274Rabbit muscle LDH (recombinant) and nicotine adenine dinucleotide (NADH) were purchased from 275Oriental Yeast (Tokyo, Japan). ANS was obtained from Sigma (Tokyo, Japan). Peptides were 276prepared by using an automated solid phase peptide synthesizer (Tetras, Advanced ChemTech, 277Louisville, KY, USA). The peptides were purified by chromatography (UFLC-20AB, Shimadzu, 278Kyoto, Japan) using a C18 reversed-phase column (AlltimaTM 4.6 x 250 mm) with a linear gradient 279of acetonitrile (from 5% to 95%) in 0.05% trifluoroacetic acid solution over 25 min. After 280identification by mass spectrometry (LCMS-2020, Shimadzu), the peptides were lyophilized for 281storage.

# 283 4.2. Cryoprotective tests for LDH

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The cryoprotective activities of peptides for LDH were estimated by the inhibition of cryoinactivation and cryodenaturation. The methods were described previously (Hara et al., 2017) with slight modifications.

288For the cryoinactivation test, 30 µL of each peptide solution (0, 0.033, 0.33, 3.3, 8.3, 17, 33, 83, 289170, and 500 µM) in 10 mM Tris-HCl buffer pH 7.5 was mixed with 20 µL of LDH solution (0.34 290 $\mu$ M as a monomer) in 1.5-mL plastic tubes. The tubes were frozen in liquid N<sub>2</sub> for 1 min and then 291transferred to a water bath at  $25\pm2$  °C for 3 min to thaw the ice. These freeze-thaw cycles were done 292three times. The LDH activities of the thawed samples were measured. The LDH reaction was 293initiated by adding peptide-containing LDH solution (4 µL) to a reaction solution (196 µL, 9.5 mM 294Tris-HCl pH 7.5, 0.58 mM sodium pyruvate, and 60 µM NADH). The reaction was monitored by 295measuring absorbance at 340 nm in a microplate reader (Varioskan Flash, Thermo Fisher Scientific, 296Tokyo, Japan) at 25 °C. Usually, the three freeze-thaw cycles reduced the LDH activity to 15 to 20% 297of the initial activity. One-hundred-percent cryoinactivation means a decreased degree of LDH activity by the freeze and thaw process. The inhibition activities of peptides for cryoinactivation 298299were evaluated as a 50% protection dose (PD<sub>50</sub>). In order to calculate PD<sub>50</sub>, we routinely used the 300 data of relative cryoinactivation at 100 µM for the maximal protection if two requirements were 301 fulfilled: 1) the relative cryoinactivation at 100  $\mu$ M was less than 20% and 2) the relative 302 cryoinactivation at 100  $\mu$ M ranged from 50% to 100% of the value at 50  $\mu$ M. When the relative 303 cryoinactivation was higher than 50% even at 300 µM, PD<sub>50</sub> was represented as more than 300 µM.

304 The cryodenaturation of LDH was analyzed by using the fluorescence of ANS, which can detect 305hydrophobic regions on a protein surface. Samples (250  $\mu$ L) containing ANS (10  $\mu$ M), LDH (4  $\mu$ M), 306 and peptides (0, 2, 5, 10, 20, 60, and 100 µM) in 10 mM sodium phosphate buffer pH 7.0 were 307 prepared in 1.5-ml plastic tubes. After the three freeze-thaw cycles as described above, fluorescence 308 at Ex 350 nm/Em 470 nm was determined (Varioskan Flash). The samples with no peptide (0  $\mu$ M) 309 showed remarkable fluorescence after the three freeze-thaw cycles. The increment of fluorescence in 310 the sample without peptide was 100%. A 50% protection dose (PD<sub>50</sub>) was used to represent the 311inhibition of cryodenaturation. When the relative cryodenaturation at 100 µM was less than 20% and 312 the relative cryodenaturation at 100  $\mu$ M ranged from 50% to 100% of the value at 50  $\mu$ M, the value 313 at 100 µM was used for the maximal protection. If the relative cryodenaturation was higher than 314 50% even at 60  $\mu$ M, PD<sub>50</sub> was represented as more than 60  $\mu$ M.

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316 4.3. Circular dichroism (CD)

318	Secondary structures of peptides (Fseg and Fseg_ $\Phi/T$ ) were analyzed by CD. Samples containing
319	the peptides (60 $\mu$ M), SDS (0.1, 1, and 10 mM) and Tris-HCl buffer pH 7.5 (10 mM) were subjected
320	to a spectropolarimeter (J-820, Jasco, Tokyo, Japan). The measurement conditions were as follows:
321	scan range from 195 to 250 nm, scan speed 100 nm min <sup>-1</sup> , resolution 1 nm, and cell width 2 mm.
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323	4.4. Peptide structures
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325	Predicted structures of peptides were produced by using the online server PEP-FOLD3
326	(http://bioserv.rpbs.univ-paris-diderot.fr/services/PEP-FOLD3/) (Lamiable et al., 2016). The
327	structures were built as in the neutral solution. The most relevant model with the best TM score was
328	chosen. The results are shown in Supplemental Fig. 3.
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330	4.5. Statistical analysis
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332	Data for $P$ values were analyzed by Dunnett's test at a significance level of 0.05.
333	
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338	
339	References
340	Ágoston, B.S., Kovács, D., Tompa, P., Perczel, A., 2011. Full backbone assignment and dynamics of
341	the intrinsically disordered dehydrin ERD14. Biomol. NMR Assign. 5, 189-193.
342	
343	Alsheikh, M.K., Heyen, B.J., Randall, S.K., 2003. Ion binding properties of the dehydrin ERD14 are
344	dependent upon phosphorylation. J. Biol. Chem. 278, 40882-40889.
345	
346	Banerjee, A., Roychoudhury, A., 2016. Group II late embryogenesis abundant (LEA) proteins:
347	structural and functional aspects in plant abiotic stress. Plant Growth Regul. 79, 1-17.
348	
349	Boddington, K.F., Graether, S.P., 2019. Binding of a Vitis riparia Dehydrin to DNA. Plant Science,
350	110172.
351	
352	Bravo, L.A., Gallardo, J., Navarrete, A., Olave, N., Martinez, J., Alberdi, M., Close, T.J., Corcuera,

L.J., 2003. Cryoprotective activity of a cold-induced dehydrin purified from barley. Physiol. Plant.118, 262-269.

355

Chakrabortee, S., Tripathi, R., Watson, M., Schierle, G.S., Kurniawan, D.P., Kaminski, C.F., Wise,
M.J., Tunnacliffe, A., 2012. Intrinsically disordered proteins asmolecular shields. Mol. Biosyst. 8,
210-219.

- 359
- Clarke, M.W., Boddington, K.F., Warnica, J.M., Atkinson, J., McKenna, S., Madge, J., Barker, C.H.,
  Graether, S.P., 2015. Structural and functional insights into the cryoprotection of membranes by the
  intrinsically disordered dehydrins. J. Biol. Chem. 290, 26900-26913.
- 363
- Close, T.J., 1997. Dehydrins: a commonalty in the response of plants to dehydration and low
   temperature. Physiol. Plant. 100, 291-296.
- 366
- Cuevas-Velazquez, C.L., Reyes, J.L., Covarrubias, A.A., 2017. Group 4 late embryogenesis
  abundant proteins as a model to study intrinsically disordered proteins in plants. Plant signal. behav.
  12, 10893-10903.
- 370
- Davis, J.G., Gierszal, K.P., Wang, P., Ben-Amotz, D., 2012. Water structural transformation at
   molecular hydrophobic interfaces. Nature 491, 582-585.
- 373

374 Drira, M., Saibi, W., Brini, F., Gargouri, A., Masmoudi, K., Hanin, M., 2013. The K-segments of the 375 wheat dehydrin DHN-5 are essential for the protection of lactate dehydrogenase and  $\beta$ -glucosidase 376 activities in vitro. Mol. Biotech. 54, 643-650.

- 377
- Eriksson, S.K., Harryson, P., 2011. Dehydrins: Molecular Biology, Structure and Function, in: Lüttge,
  U., Beck, E., Bartels, D. (Eds.), Plant Desiccation Tolerance. Ecological Studies (Analysis and
  Synthesis), vol 215. Springer, Berlin, Heidelberg, pp. 289-305.
- 381
- Eriksson, S.K., Kutzer, M., Procek, J., Gröbner, G., Harryson, P., 2011. Tunable membrane binding
  of the intrinsically disordered dehydrin Lti30, a cold-induced plant stress protein. Plant Cell 23,
  2391-2404.

385

Findlater, E.E., Graether, S.P., 2009. NMR assignments of the intrinsically disordered K 2 and YSK
2 dehydrins. Biomol. NMR Assign. 3, 273-275.

- Graether, S.P., Boddington, K.F., 2014. Disorder and function: a review of the dehydrin protein
  family. Front. Plant Sci. 5, 576.
- 391
- Hand, S.C., Menze, M.A., Toner, M., Boswell, L., Moore, D., 2011. LEA proteins during water
  stress: not just for plants anymore. Annu. Rev. Physiol. 73, 115-134.
- 394
- Hanin, M., Brini, F., Ebel, C., Toda, Y., Takeda, S., 2011. Plant dehydrins and stress tolerance:
  versatile proteins for complex mechanisms. Plant Signal. Behav. 6, 1503-1509.
- 397
- Hara, M., Endo, T., Kamiya, K., Kameyama, A., 2017. The role of hydrophobic amino acids of
  K-segments in the cryoprotection of lactate dehydrogenase by dehydrins. J. Plant Physiol. 210,
  18-23.
- 401
- Hara, M., Fujinaga, M., Kuboi, T., 2005. Metal binding by citrus dehydrin with histidine-rich
  domains. J. Exp. Bot. 56, 2695-2703.
- 404
- Hara, M., Kondo, M., Kato, T., 2013. A KS-type dehydrin and its related domains reduce
  Cu-promoted radical generation and the histidine residues contribute to the radical-reducing
  activities. J. Exp. Bot. 64, 1615-1624.
- 408
- Hara, M., Shinoda, Y., Tanaka, Y., Kuboi, T., 2009. DNA binding of citrus dehydrin promoted byzinc ion. Plant Cell Environ. 32, 532-541.
- 411
- Hara, M., Terashima, S., Fukaya, T., Kuboi, T., 2003. Enhancement of cold tolerance and inhibition
  of lipid peroxidation by citrus dehydrin in transgenic tobacco. Planta 217, 290-298.
- 414
- Hara, M., Terashima, S., Kuboi, T., 2001. Characterization and cryoprotective activity of
  cold-responsive dehydrin from *Citrus unshiu*. J. Plant Physiol. 158, 1333-1339.
- 417
- 418 Hernández-Sánchez, I.E., Martynowicz, D.M., Rodríguez-Hernández, A.A., Pérez-Morales, M.B.,
- Graether, S.P., Jiménez-Bremont, J.F., 2014. A dehydrin-dehydrin interaction: the case of SK3 from *Opuntia streptacantha*. Front Plant. Sci. 5, 520.
- 421
- 422 Hughes, S., Graether, S.P., 2011. Cryoprotective mechanism of a small intrinsically disordered 423 dehydrin protein. Protein Sci. 20, 42-50.

- 425 Hughes, S.L., Schart, V., Malcolmson, J., Hogarth, K.A., Martynowicz, D.M., Tralman-Baker, E.,
- 426 Patel, S.N., Graether, S.P., 2013. The importance of size and disorder in the cryoprotective effects of427 dehydrins. Plant Physiol. 163, 1376-1386.
- 428
- 429 Hundertmark, M., Hincha, D.K., 2008. LEA (late embryogenesis abundant) proteins and their 430 encoding genes in *Arabidopsis thaliana*. BMC Genom. 9, 118.
- 431
- Koag, M.C., Wilkens, S., Fenton, R.D., Resnik, J., Vo, E., Close, T.J., 2009. The K-segment of maize
  DHN1 mediates binding to anionic phospholipid vesicles and concomitant structural changes. Plant
  Physiol. 150, 1503-1514.
- 435

Lamiable, A., Thévenet, P., Rey, J., Vavrusa, M., Derreumaux, P., Tufféry, P., 2016. PEP-FOLD3:
faster de novo structure prediction for linear peptides in solution and in complex. Nucl. Acid. Res. 44,
W449-W454.

439

Liu, X., Zhang, C.S., Lu, C., Lin, S.C., Wu, J.W., Wang, Z.X., 2016. A conserved motif in
JNK/p38-specific MAPK phosphatases as a determinant for JNK1 recognition and inactivation.
Nature Commun. 7, 10879.

443

Mace, P.D., Wallez, Y., Egger, M.F., Dobaczewska, M.K., Robinson, H., Pasquale, E.B., Riedl, S.J.,
2013. Structure of ERK2 bound to PEA-15 reveals a mechanism for rapid release of activated
MAPK. Nature Commun. 4, 1681.

- 447
- Nylander, M., Svensson, J., Palva, E.T., Welin, B., 2001. Stress-induced accumulation and tissue
  specific localization of dehydrins in *Arabidopsis thaliana*. Plant Mol. Biol. 45, 263-279.
- 450
- 451 Ochoa-Alfaro, A.E., Rodríguez-Kessler, M., Pérez-Morales, M.B., Delgado-Sánchez, P.,
  452 Cuevas-Velazquez, C.L., Gómez-Anduro, G., Jiménez-Bremont, J.F., 2012. Functional
  453 characterization of an acidic SK 3 dehydrin isolated from an *Opuntia streptacantha* cDNA library.
  454 Planta 235, 565-578.

- Palmer, S.R., De Villa, R., Graether, S.P., 2019. Sequence composition versus sequence order in the
  cryoprotective function of an intrinsically disordered stress-response protein. Protein Sci. 28,
  1448-1459.
- 459
- 460 Peng, Y., Reyes, J.L., Wei, H., Yang, Y., Karlson, D., Covarrubias, A.A., Krebs, S.L., Fessehaie, A.,

- 461 Arora, R., 2008. RcDhn5, a cold acclimation-responsive dehydrin from *Rhododendron catawbiense*462 rescues enzyme activity from dehydration effects in vitro and enhances freezing tolerance in
  463 RcDhn5 overexpressing *Arabidopsis* plants. Physiol. Plant. 134, 583-597.
- 464
- Puhakainen, T., Hess, M.W., Mäkelä, P., Svensson, J., Heino, P., Palva, E.T., 2004. Overexpression
  of multiple dehydrin genes enhances tolerance to freezing stress in *Arabidopsis*. Plant Mol. Biol. 54,
  743-753.
- 468
- Rahman, L.N., Smith, G.S., Bamm, V.V., Voyer-Grant, J.A., Moffatt, B.A., Dutcher, J.R., Harauz, G.,
  2011. Phosphorylation of *Thellungiella salsuginea* dehydrins TsDHN-1 and TsDHN-2 facilitates
- 471 cation-induced conformational changes and actin assembly. Biochemistry 50, 9587-9604.472
- Reyes, J.L., Campos, F., Wei, H.U.I., Arora, R., Yang, Y., Karlson, D.T., Covarrubias, A.A., 2008.
  Functional dissection of hydrophilins during in vitro freeze protection. Plant Cell Environ. 31,
  1781-1790.
- 476
- 477 Strimbeck, G.R., 2017. Hiding in plain sight: the F segment and other conserved features of seed478 plant SKn dehydrins. Planta 245, 1061-1066.
- 479
- Tompa, P., Csermely, P., 2004. The role of structural disorder in the function of RNA and proteinchaperones. FASEB J. 18, 1169-1175.
- 482
- Wei, H., Yang, Y., Himmel, M.E., Tucker, M.P., Ding, S.Y., Yang, S., Arora, R., 2019. Identification
  and characterization of five cold stress-related rhododendron dehydrin genes: spotlight on a
  FSK-type dehydrin with multiple F-segments. Front. Bioeng. Biotech. 7, 30.
- 486
- Xing, X., Liu, Y., Kong, X., Liu, Y., Li, D., 2011. Overexpression of a maize dehydrin gene,
  ZmDHN2b, in tobacco enhances tolerance to low temperature. Plant Growth Regul. 65, 109-118.
- 489
- Yang, W., Zhang, L., Lv, H., Li, H., Zhang, Y., Xu, Y., Yu, J., 2015. The K-segments of wheat
  dehydrin WZY2 are essential for its protective functions under temperature stress. Front Plant Sci
  6:406.
- 493

Zhang, A., Qi, W., Singh, S.K., Fernandez, E.J., 2011. A new approach to explore the impact of
freeze-thaw cycling on protein structure: hydrogen/deuterium exchange mass spectrometry
(HX-MS). Pharm. Res. 28, 1179-1193.

Zhang, H., Shi, Y., Liu, X., Wang, R., Li, J., Xu, J., 2018. Transgenic creeping bentgrass plants
expressing a *Picea wilsonii* dehydrin gene (PicW) demonstrate improved freezing tolerance. Mol.
Biol. Rep. 45, 1627-1635.

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502 Figure legends

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Fig. 1. Cryoprotective activities of typical K-segment (TypK) and *Arabidopsis* F-segments. (A) Peptide names, corresponding dehydrins, and amino acid sequences are shown. Inhibition of cryoinactivation (B) and that of cryodenaturation (C) of lactate dehydrogenase (LDH) by the peptides are shown as PD<sub>50</sub>. Values shown are the means of three experiments. Columns and bars represent means  $\pm$  SD. No significant difference (p < 0.05) from TypK.

509

510Fig. 2. Cryoprotective activities of Fseg and its related peptides (amino acid substitutions). (A) Amino acid sequences of the peptides. Fseg means COR47Fseg in Fig. 1.  $\Phi$ , +, and - represent 511512hydrophobic, positively charged, and negatively charged amino acids, respectively. Letters of 513hydrophobic amino acids are shaded. Inhibition of cryoinactivation (B) and that of cryodenaturation 514(C) were evaluated as  $PD_{50}$ . Values shown are the means of three experiments. Columns and bars 515represent means  $\pm$  SD. Asterisks indicate significant differences (p < 0.05) from Fseg. When the 516PD<sub>50</sub> values were more than 300 µM in B and 60 µM in C, the tops of the columns were shredded. In 517these cases, asterisks are not labeled.

518

519Fig. 3. Cryoprotective activities of Fseg and its related peptides (G substitutions). (A) Amino acid sequences of the peptides. Fseg means COR47Fseg in Fig. 1.  $\Phi$ , +, and - represent hydrophobic, 520521positively charged, and negatively charged amino acids, respectively. Letters of hydrophobic amino 522acids are shaded. Inhibition of cryoinactivation (B) and that of cryodenaturation (C) were evaluated 523as PD<sub>50</sub>. Values shown are the means of three experiments. Columns and bars represent means  $\pm$  SD. 524Asterisks indicate significant differences (p < 0.05) from Fseg. When the PD<sub>50</sub> values were more 525than 300  $\mu$ M in B and 60  $\mu$ M in C, the tops of the columns are shredded. In these cases, asterisks are 526not labeled.

527

528 Fig. 4. Analyses of secondary structures of Fseg and Fseg  $\Phi/T$ . (A) Circular dichroism analysis for

529Fseg and Fseg  $\Phi/T$ . Effects of sodium dodecyl sulfate (SDS) on their secondary structures are 530shown. Gray broken lines, gray solid lines, black broken lines, and black solid lines represent 0, 0.1, 5311, and 10 mM SDS, respectively. (B) Predictive quantification of the secondary structure 532compositions in Fseg and Fseg  $\Phi/T$ by using K2D2 533(http://cbdm-01.zdv.uni-mainz.de/~andrade/k2d2/). The CD data in A were used. Black, gray, and white columns show  $\alpha$ -helix,  $\beta$ -strand, and others (random structures), respectively. Values shown 534535are the means of three experiments. Bars represent SD. An asterisk indicates a significant difference 536(p < 0.05) from the control sample (0 mM SDS).

537

538Fig. 5. Putative mechanisms for cryoprotective activity of Fseg. When target proteins (e.g., LDH) 539are frozen and thawed, hydrophobic patches appear on the surface of the proteins, and then 540hydrophobic interaction is generated between the patches (A). Fseg intervenes in the hydrophobic 541interaction via its hydrophobic area. While Fseg inhibits hydrophobic binding, hydrophobic 542hydration may cover the patches. Finally, the native state of proteins was maintained (B). Besides 543that, Fseg  $\Phi/T$  cannot intervene in the hydrophobic interaction because it does not have a 544hydrophobic area. A water-tight seal forms between the hydrophobic patches of the target proteins 545after exclusion of Fseg  $\Phi/T$ . As a result, the proteins are aggregated (C).

546

# 547 Supplemental Figure legends

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Supplemental Fig. 1. Arabidopsis FSKn dehydrins. (A) Amino acid sequences of four Arabidopsis
FSKn dehydrins. F-segments are highlighted in green. Red letters indicate K-segments. (B)
Information on the four FSKn dehydrins. Cold-responsive expressions were determined according to
the data from the Arabidopsis eFP Browser (http://bar.utoronto.ca/efp/cgi-bin/efpWeb.cgi).

553

**Supplemental Fig. 2.** Cryoprotective tests for lactate dehydrogenase (LDH). (A) Scheme of the assay. Samples containing LDH, peptides, and buffer were treated with three freeze and thaw cycles, after which enzyme activities and denaturation levels were determined. For peptides, Fseg (COR47Fseg) and Fseg\_ $\Phi/T$  (COR47Fseg\_ $\Phi/T$ ) were used. (B) Relative cryoinactivation of LDH (%). Values and bars represent means  $\pm$  SD (three experiments). (C) Relative cryodenaturaion of LDH (%). The denaturation levels were monitored by fluorescence (ex 350 nm, em 470 nm) from 560 8-anilino-1-naphthalene sulfonic acid (ANS). Values and bars represent means  $\pm$  SD (three 561 experiments).

562

563 **Supplemental Fig. 3A, B.** Predicted peptide structures of F-segments. The structures were built 564 using PEP-FOLD3 (http://bioserv.rpbs.univ-paris-diderot.fr/services/PEP-FOLD3/). Structures of 565 Fseg (A) and Fseg\_ $\Phi/T$  (B) are shown. Hydrophobic areas are indicated by yellow elliptical 566 shadows. N and C refer to the N- and C-termini, respectively. Single-letter codes with numbers are 567 positions of amino acid residues in the peptides.

568

**Supplemental Fig. 3C.** Predicted peptide structures of F-segments. The structures were built by PEP-FOLD3 (http://bioserv.rpbs.univ-paris-diderot.fr/services/PEP-FOLD3/). Structures of ERD10Fseg, ERD14Fseg, At4g38410Fseg, pG\_Fseg\_ $\Phi$ , and TypK are shown. Hydrophobic areas are indicated by yellow elliptical shadows. N and C refer to the N- and C-termini, respectively. Single-letter codes with numbers are positions of amino acid residues in the peptides.

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Supplemental Fig. 4. An alignment of amino acid sequences of *Arabidopsis* COR47 and the
orthologues from other plant species. All five dehydrins are classified as FSK3 types. *Arabidopsis, Rhododendron, Malus, Nicotiana,* and *Theobroma* represent COR47 from *Arabidopsis,* dehydrin 2
(AGI36547) from *Rhododendron catawbiense,* dehydrin COR47-like (NP\_001315732) from *Malus domestica,* dehydrin (BAD13499) from *Nicotiana tabacum,* and dehydrin 2 (EOY15190) from *Theobroma cacao,* respectively. Asterisks are identical amino acids between species. Black, gray,
and white columns represent F-, S-, and K-segments, respectively.

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Name	Dehydrin	Amino acid sequence
ТурК	-	EKKGIMEKIKEKLPG
COR47Fseg	COR47	D <sub>31</sub> RG <u>LF</u> D <u>FL</u> GKKEEE <u>V</u> 45
ERD10Fseg	ERD10	E <sub>29</sub> RG <u>MF</u> D <u>FL</u> KKKEE <u>V</u> K <sub>43</sub>
ERD14Fseg	ERD14	V <sub>25</sub> TDRG <u>LF</u> D <u>FL</u> GKKKD <sub>39</sub>
At4g38410Fseg	At4g38410	G <sub>20</sub> CG <u>MF</u> D <u>FL</u> KKKPED <u>V<sub>34</sub></u>





Fig. 1 Ohkubo et al.

А



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A	1	2	3	4	5	6	7	8	9	1	1	1	1	1	1
	-	+		Φ	Φ	-	Φ	Φ		0	1	2	3	4	5
										+	+	-	-	-	φ
Fseg	D	R	G	L	F	D	F	L	G	к	К	Е	Е	Е	۷
Fseg_Ф/T	D	R	G	Т	Т	D	Т	Т	G	к	к	Е	Е	Е	Т
Fseg_DE/T	т	R	G	L	F	Т	F	L	G	к	к	Т	Т	Т	V
Fseg_KR/T	D	Т	G	L	F	D	F	L	G	Т	Т	Е	Е	Е	V
Fseg_Ф/E	D	R	G	Е	Е	D	Е	Е	G	к	К	Е	Е	Е	Е
Fseg_Ф/K	D	R	G	к	к	D	к	к	G	к	к	Е	Е	Е	к





Fig. 2 Ohkubo et al.



Fig. 3 Ohkubo et al.



Fig. 4 Ohkubo et al.





A

#### At1g20440 COR47 (FSK3)

MAEEYKNNVPEHETPTVATEESPATTTEVT DRGLEDFLGKKEEEVKPQETTTLESEFDHKAQISEPEL AAEHEEVKENKITLLEELQEKTEEDEENKPSVIEKLHRSNSSSSSSSDEEGEEKKEKKKKIVEGEEDKKG LVEKIKEKLPGHHDKTAEDDVPVSTTIPVPVSESVVEHDHPEEKKGLVEKIKEKLPGHHDEKAEDSPA VTSTPLVVTEHPVEPTTELPVEHPEEKKGILEKIKEKLPGYHAKTTEEEVKKEKESDD

#### At1g20450 ERD10 (FSK3)

MAEEYKNTVPEQETPKVATEESSAPEIK**ERGMEDELKKKEEVK**PQETTTLASEFEHKTQISEPESFVAK HEEEEHKPTLLEQLHQKHEEEEENKPSLLDKLHRSNSSSSSSDEEGEDGEKKKKEKKKKIVEGDHVK TVEEENQGVMDRIKEKFPLGEKPGGDDVPVVTTMPAPHSVEDHKPEEEEKKGFMDKIKEKLPGHSKK PEDSQVVNTTPLVETATPIADIPEEKKGFMDKIKEKLPGYHAKTTGEEEKKEKVSD

### ♦At1g76180 ERD14 (FSK2)

MAEEIKNVPEQEVPKVATEESSAE<mark>VTDRGLFDFLGKKKD</mark>ETKPEETPIASEFEQKVHISEPEPEVKHESL LEKLHRSDSSSSSSEEEGSDGEKRKKKEKKKPTTEVEVKEE<mark>EKKGFMEKLKEKLPG</mark>HKKPEDGSA VAAAPVVVPPPVEEAHPVEKKGILEKIKEKLPGYHPKTTVEEEKKDKE

#### At4g38410 (FSK2)

MADHPRSTEQQEADAAASK GCGMFDFLKKKPEDVHSSENARVTKEPKEEEKPSLAERFHLSDSSSSD EEAGENGEKKEKKKKKKNEVAEDQCETEEKIPAGIGHEDGKEKGFMEKIKDKLPGGHNGKPEAEPH NDKAKEKGFMEKIKEKLPGHTNDEKKKET

## В

Dehydrin	Gene ID	Dehydrin type	Size (amino acids)	Expression
COR47	At1g20440	FSK3	265	Cold
ERD10	At1g20450	FSK3	260	Cold
ERD14	At1g76180	FSK2	185	Cold
At4g38410	At4g38410	FSK2	163	Cold

Supplemental Fig. 1. Arabidopsis FSKn dehydrins. (A) Amino acid sequences of four Arabidopsis FSKn dehydrins. F-segments are highlighted in green. Red letters indicate K-segments. (B) Information on the four FSKn dehydrins. Cold-responsive expressions were determined according to the data from the Arabidopsis eFP Browser (http://bar.utoronto.ca/efp/cgi-bin/efpWeb.cgi).

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Arabidopsis	M	1
Rhododendron	MRWPPVAPAETKDRGLFDFLGKKEEEETVVPAKEDKSKDQYGEVDEVAPVAPAGTKDRGL	60
Nicotiana	M	1
Theobroma	M	1
Arabidopsis	AEEYKNNVPEHETPTVATEESPATTTEVTDRGLFDFLGKKEEEVKP	47
Rhododendron	FDFLGKKEEEETAVPAKENKSKDQYGEVDEVAPVAPGETKDRGLFDFLGKKEEEEHKKPQ	120
Malus	AEEYNKKSDEHEYERKTGDYEEGSGAGETKDRGLFDFLGKKEEEKPTPYQ	51
Nicotiana		39
meobrollia	AEENIISKAFEGESIIVAGEGAVEIKDRGLFDFLGKKEEKKFG	42
Arabidopsis	QETTTLESEFDHKAQISEPELAAEHEEVKENKITLLEELQEKTEEDEENKPSVIEKLHRS	107
Rhododendron	DEEEVIVTEFEK-VKVSEPETKEFKEEEKKDGLLTK-	155
Malus	QGDQVNVAEFDEKVKISDHHDQHASSYNKVEEEEDKEKKHETLLQKLHRS	101
Nicotiana		80
Ineoproma		85
Arabidopsis	NSSSSSSDEEGEEKKEKKKIVEGEEDKKGLVEKIKEK-LPGHHDKTAE	156
Rhododendron	SSSSSSSEEGEGGEKKKKKKGLNEKIEEIKIAGDREEEQEAKIDKQEEKD	206
Malus	ESSSSSSDEEEDEEKKKKRKEKKGLTDKIKEK-ISGDEHKEEGYHKEED	150
Nicotiana	SSSSSSSDEEEEIGEDGQKIKKKKKKGLKDKIKDK-ISGEHKEEEKAGED	130
Theobroma	DSSSSSSDEEEGEGEEKKKKKKKKKKKKKKKKKKKKKKGLKEKIEEK-LEGEKKEEEMKHED	138
	*****	
Arabidopsis	DDVPVSTTIPVPVSESVVEHDHPEEEKKGLVEKIKEKLPGHHDEKAEDS	205
Rhododendron	KLVPVEEYEEVYEEAAVTTPPAEEKKGFLEKIKQKLPGKNKKTEEVP	253
Malus	TAVPVEKVYEEEHHHPAPAPAPVVHYHEEPTDSPTEEKKGFLEKIKEKLPGHKKTEEVPV	210
Nicotiana	TAVPVEKYEETEEKKGFLDKIKEKLPGGGQKKTEEV	166
Theobroma	TSVPVEKCDEPVVQAEPEKKGFLEK1KEKLPGQYKKAEEVP	179
	*** ****	
Arabidopsis	PAVTSTPLVVTEHPVEPTTELPVEHPEEKKGILEKIKEKLPGYHAKTTEEEVKKEKE	262
Rhododendron	PPSTPPPPS-DVEYVEPEPKEKKGILEKIKEKIPGHKTEEEKQRQKES	300
Malus	GAASHEQHSDDKHAAEPPVAASYEAGEEPKEKKGILEKIKEKLPGYHSKPEEDHKDIKEK	270
Nicotiana	APPPPPAAEHEAEGKEKKGFLDKIKEKLPGYHSKTEEKKKKK	208
Theobroma	PPPPPPPAECVAAEPHEGAEAKEKKGILEKIKEKLPGYHSKTEEEKEKEKES	231
Arabidanaia	**** * ***** ** 265	
Rhododendron	TDN 203	
Malus	FKDTPSY 277	
Nicotiana	208	
Theobroma	ASH 234	

Supplemental Fig. 4. An alignment of amino acid sequences of Arabidopsis COR47 and the orthologues from other plant species. All five dehydrins are classified as FSK3 types. Arabidopsis, Rhododendron, Malus, Nicotiana, and Theobroma represent COR47 from Arabidopsis, dehydrin 2 (AGI36547) from Rhododendron catawbiense, dehydrin COR47-like (NP\_001315732) from *Malus domestica*, dehydrin (BAD13499) from *Nicotiana tabacum*, and dehydrin 2 (EOY15190) from *Theobroma cacao*, respectively. Asterisks are identical amino acids between species. Black, gray, and white columns represent F-, S-, and K-segments, respectively. **Title**: F-segments of *Arabidopsis* dehydrins show cryoprotective activities for lactate dehydrogenase depending on

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