

2 **Dehydrin, alcohol dehydrogenase, and central metabolite**
3 **levels are associated with cold tolerance in diploid strawberry**
4 **(*Fragaria* spp.)**

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10 **Abstract** The use of artificial freezing tests, identification
11 of biomarkers linked to or directly involved in the low-
12 temperature tolerance processes, could prove useful in
13 applied strawberry breeding. This study was conducted to
14 identify genotypes of diploid strawberry that differ in their
15 tolerance to low-temperature stress and to investigate

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whether a set of candidate proteins and metabolites corre- 16
late with the level of tolerance. 17 *Fragaria vesca*, 2 17
F. nilgerrensis, 2 *F. nubicola*, and 1 *F. pentaphylla* genotypes 18
were evaluated for low-temperature tolerance. Estimates of 19
temperatures where 50 % of the plants survived (LT₅₀) 20
ranged from -4.7 to -12.0 °C between the genotypes. 21
Among the *F. vesca* genotypes, the LT₅₀ varied from 22
-7.7 °C to -12.0 °C. Among the most tolerant were three 23
F. vesca ssp. *bracteata* genotypes (FDP821, NCGR424, 24
and NCGR502), while a *F. vesca* ssp. *californica* genotype 25
(FDP817) was the least tolerant (LT₅₀ -7.7 °C). Alcohol 26
dehydrogenase (ADH), total dehydrin expression, and 27
content of central metabolism constituents were assayed in 28
select plants acclimated at 2 °C. The LT₅₀ estimates and 29
the expression of ADH and total dehydrins were highly 30
negatively correlated ($r_{adh} = -0.87$, $r_{dehyd} = -0.82$). 31
Compounds related to the citric acid cycle were quantified 32
in the leaves during acclimation. While several sugars and 33
acids were significantly correlated to the LT₅₀ estimates 34
early in the acclimation period, only galactinol proved to 35
be a good LT₅₀ predictor after 28 days of acclimation 36
($r_{galact} = 0.79$). It is concluded that ADH, dehydrins, and 37
galactinol show great potential to serve as biomarkers for 38
cold tolerance in diploid strawberry. 39

Keywords Galactinol · Hierarchical clustering · Lethal 41
temperature 50 · Metabolite profiling · Raffinose pathway · 42
Survival analysis 43

Abbreviations 44
ABA Abscisic acid 45
ADH Alcohol dehydrogenase 46
CBF C-repeat/dehydration responsive element binding 47
factor
FDP *Fragaria* diploid project 48

49 GC–MS Gas chromatography and mass spectrometry
 50 LT₅₀ Temperature where 50 % of the plants are killed
 51 NCGR National Clonal Germplasm Repository
 52 PCA Principal component analyses
 53 PPFDF Photosynthetic photon flux density
 54
 55

56 **Introduction**

57 In areas where strawberry is grown in a perennial growing
 58 system, the plants have to survive through the winter. In
 59 Scandinavia, the majority of the strawberry produce comes
 60 from a perennial production system, and winter survival is
 61 a major limiting factor for the strawberry industry. A
 62 typical annual yield reduction is 20 %, with an occasional
 63 total loss (Davik et al. 2000).

64 The survival of strawberry plants in areas with low
 65 temperatures is affected by several physiological respon-
 66 ses, e.g., growth cessation, effective cold hardening in
 67 autumn, and the response to growth stimulation in periods
 68 of temporarily increased temperatures. Abiotic stresses like
 69 ice encasement, desiccation, and soil heaving add to the
 70 problem. With a range of biotic and abiotic factors con-
 71 tributing and interacting, disentangling the full story of
 72 winter survival has proven difficult. However, low-tem-
 73 perature stress per se is one important aspect of winter
 74 survival, and in particular during periods with little or no
 75 snow cover. The cycles of freezing and thawing during
 76 wintertime have been shown to be particularly harmful to
 77 strawberry plants. In such cases, the use of insulating
 78 cover, either snow or ice, has a significant impact on both
 79 the yield and the quality of the yield (Nestby et al. 2000).

80 Differences among cultivars in winter survival have
 81 been known to the industry and also experimentally con-
 82 firmed both for octoploid *Fragaria* × *ananassa* (Nestby
 83 and Bjørgum 1999) and diploid *F. vesca* genotypes (Søn-
 84 stebly and Heide 2011). Hence, selecting for winter survival
 85 is a prime objective for strawberry breeding programs.
 86 Given the complexity of the trait and the often fluctuating
 87 winter weather, extensive field testing over many years
 88 would be required to gain reliable results. More rapid
 89 laboratory tests are therefore required. Testing for frost
 90 tolerance under controlled conditions to grade genotypes
 91 has been used in several plant breeding programs to iden-
 92 tify superior genotypes, e.g., in wheat (Gusta et al. 1997)
 93 and oilseed rape (Teutonico et al. 1993). This approach
 94 could also be a valuable alternative for the strawberry
 95 breeder.

96 In addition to the use of artificial freezing tests, identi-
 97 fication of biomarkers linked to or directly involved in low-
 98 temperature tolerance processes could prove useful in

applied strawberry breeding. During acclimation, plants
 from temperate and cold climates develop increased tol-
 erance to subsequent low-temperature exposure, and
 changes in expression of hundreds of genes have been
 demonstrated in *Arabidopsis thaliana* (Kaplan et al. 2007).
 In strawberry vegetative tissue, metabolite profiles are
 totally reconfigured as a result of the low-temperature
 impact (e.g., Rohloff et al. 2012). The metabolic cold
 response results in increased levels of compatible solutes
 such as free amino acids, amines, polyols, and mono-, di-,
 and trisaccharides as described for the model *Arabidopsis*
thaliana (Korn et al. 2010). The molecules’ osmo-protect-
 ive role is based on their properties to stabilize and pre-
 vent proteins, membranes (as reviewed by Kaplan et al.
 2007), and nucleic acids (Kurz 2008) from the damaging
 effects of freezing temperatures. Moreover, secondary
 metabolism is also strongly affected leading to the up-
 regulation of photoprotective flavonoids (Hannah et al.
 2006). The prominent role of the raffinose pathway
 (Rohloff et al. 2009) and central carbohydrate metabolism
 is documented in several studies (Guy et al. 2008), and a
 significant correlation between freezing tolerance and
 carbohydrate content and accumulation during acclimation
 has been demonstrated in *A. thaliana* (Hannah et al. 2006).

In other species, expression of alcohol dehydrogenase
 (ADH) is known to increase under various stresses,
 including low temperature, drought, abscisic acid (ABA),
 and salinity (Christie et al. 1991; Jarillo et al. 1993;
 Dolferus et al. 1994; de Bruxelles et al. 1996; Lindlöf et al.
 2007; Diab et al. 2008). In particular, ADH genes are
 among the most commonly found cold-induced genes in
 cereal crops and *Arabidopsis* (Lindlöf et al. 2007). Our
 own preliminary observations have shown a high correla-
 tion between ADH levels and cold tolerance in the octo-
 ploid strawberry.

Another group of candidate marker proteins are the
 dehydrins. Dehydrins comprise a family of proteins that are
 produced in response to low temperatures and drought
 stress. Dehydrins are often regulated by the CBF cold-
 responsive pathway and are among the most commonly
 reported proteins accumulating in plants in response to cold
 stress (Close 1996). Dehydrins are well conserved in the
 plant genera, and homologs are readily identified by
 sequence similarity and occurrence of the dehydrin con-
 sensus sequence (Close 1997). In Rosaceous species, de-
 hydrins have been identified that have high similarity to
Arabidopsis dehydrins (Artlip et al. 1997; Bassett et al.
 2009; Garcia-Bañuelos et al. 2009) and in strawberry
 (*F. × ananassa*) Koehler et al. (2012) identified two de-
 hydrin-like proteins (COR47-like, XERO2-like) that were
 regulated by cold exposure.

Indirect selection using a marker-assisted approach
 could enhance the efficiency of cultivar development.

Author Proof

152 Although rudimentary linkage maps have been emerging
 153 (Rousseau-Gueutin et al. 2008; Sargent et al. 2009), com-
 154 mercially grown strawberry cultivars are difficult to dis-
 155 entangle in genetic studies due to their octoploid genome.
 156 To understand the molecular basis for low-temperature
 157 stress and develop molecular markers linked to stress tol-
 158 erance, we chose a model system using diploid *Fragaria*
 159 species in a screening for diverging genotypes. The octo-
 160 ploid strawberry progenitors *F. virginiana* and *F. chilo-*
 161 *ensis* are believed to be diploidized allopolyploids, each
 162 descending from four diploid ancestors. The ancestry of
 163 *F. virginiana* and *F. chiloensis* is not fully known, but
 164 *F. vesca*, *F. iinumae*, *F. nubicola*, and *F. orientalis* have
 165 been suggested by some authors (Potter et al. 2000; Folta
 166 and Davis 2006), while Rousseau-Gueutin et al. (2009)
 167 have found evidence for *F. vesca*, *F. mandshurica*, and
 168 *F. iinumae* being strong candidates. So, there appears to be
 169 a consensus among the authors that at least *F. vesca* is one
 170 of the early ancestors.

171 Diploid strawberry species have several features that make
 172 them attractive as model species. The plants are easily grown
 173 and propagated both through seeds and runners, and they are
 174 relatively easy to transform genetically (Oosumi et al. 2006).
 175 Moreover, the *F. vesca* genome is relatively small
 176 (~240 Mb) and has recently been sequenced (Shulaev et al.
 177 2011). Finally, a high degree of macrosynteny and collin-
 178 earity between diploid and octoploid strawberry exist, and no
 179 major chromosomal rearrangements seem to have occurred
 180 (Rousseau-Gueutin et al. 2008). This conserved organization
 181 within the *Fragaria* genus supports the use of diploid *Fra-*
 182 *garia* as a model system to gain genetic knowledge that
 183 subsequently can be transferred to the more complex and
 184 economically important octoploid *F. × ananassa* (Davis and
 185 Yu 1997; Sargent et al. 2004).

186 This study was conducted to identify genotypes of dip-
 187 loid strawberry that diverge in their tolerance to low-tem-
 188 perature stress and investigate whether a set of candidate
 189 proteins and metabolites show correlation with the level of
 190 tolerance. The work presented here is part of a project
 191 where the main goal is to gain basic knowledge about the
 192 genetic variation of winter survival of strawberry. The
 193 development of molecular markers useful in the amelio-
 194 ration of strawberry cultivars with improved winter sur-
 195 vival rate is our long-term goal.

196 Materials and methods

197 Plant material and multiplication

198 The plants were either collected as runners in Norway (Alta,
 199 Bukammen, and Haugastøl) or obtained as seeds from the
 200 National Clonal Germplasm Repository (NCGR-accessions)

in Corvallis, OR, USA, and East Malling Research (FDP-
 201 accessions), UK. Seeds were propagated and one single plant
 202 was collected from each of the accessions mentioned in
 203 Table 1, hereafter called 'genotype' or 'genotypes', even
 204 though we retain the original label. Multiplication of each of
 205 the genotypes was subsequently done by runnering, aiming
 206 for uniform test plants. The plants were then raised in a heated
 207 greenhouse for 5 weeks maintained at 20 ± 2 °C and 20-h
 208 photoperiod. Throughout the experiments, the plants were
 209 grown in 10 cm plastic pots containing a peat-based potting
 210 compost (90 % peat, 10 % clay), with the addition of 1:5
 211 (v/v) of granulated perlite. The plants were watered twice a
 212 week (and 1 day immediately before harvesting for freezing
 213 treatments), sufficient to keep the soil moist at all times. A
 214 balanced nutrient solution containing 7.8 mmol N, 1 mmol P,
 215 and 4.6 mmol K per liter (used in 1:100 ratio) was applied
 216 twice a week.
 217

Freezing experiments 218

219 For the LT₅₀ determinations, the plants were subsequently
 220 acclimated for 6 weeks at 2 °C and 10-h photoperiod. Sup-
 221 plemental light was provided by high-pressure sodium lamps
 222 (SON-T) at a PPFD of about 90 μmol quanta m⁻² s⁻¹. After
 223 hardening, the plants were exposed to freezing temperatures
 224 ranging from 0 to -27 °C (0, -8, -9, -10, -12, -14, -15,
 225 -18, -21, -24, and -27 °C). The freezing was performed in
 226 darkness in freeze cabinets initially set at 2 °C. The temper-
 227 ature was immediately lowered to -2 °C, and kept at this
 228 temperature for 12 h to ensure that the soil in the pots was
 229 frozen. The temperature was then lowered by 2 °C/h until the
 230 target temperature was reached where it was held for 4 h,
 231 before raising the temperature by 2 °C/h to 2 °C and holding
 232 for 10 h. Control plants were exposed to 0 °C in darkness for
 233 12 h. After completion of the freezing exposure, the plants
 234 were moved into a greenhouse maintained at 18 ± 2 °C and
 235 20-h photoperiod for 5 weeks before survival was scored
 236 (dead or alive).

Setup and statistical analysis of the freezing experiments 237 238

239 Six freezing experiments were performed under identical
 240 conditions with the 22 genotypes presented in Table 1. In
 241 each experiment, we used 12 clonally propagated plants
 242 from each genotype in each of the temperature treatments.
 243 Occasionally, and for some genotypes, only nine plants
 244 were used due to the great variation in stolon formation
 245 between the genotypes. For the same reason, some geno-
 246 types were represented in four experiments, while one was
 247 represented only once. On average, each genotype was
 248 represented 2.5 times in one of the six experiments.
 249 However, statistical connectivity between the experiments

Table 1 Strawberry genotypes included in this study

Accession ID/genotype	Species subspecies	Origin	Altitude (m a.s.l.)	LT ₅₀ (°C) ± SE
FDP821/NCGR546	<i>F. vesca</i> ssp. <i>bracteata</i>	Wyoming, USA	1,200	-12.0 ± 1.2
NCGR1428	<i>F. vesca</i>	Bolivia	n/a	-12.0 ± 1.7
Alta	<i>F. vesca</i> ssp. <i>vesca</i>	Alta, Norway	50	-11.6 ± 1.2
NCGR1603	<i>F. vesca</i>	Rakitovo, Bulgaria	1,070	-11.1 ± 1.3
NCGR424	<i>F. vesca</i> ssp. <i>bracteata</i>	Oregon, USA	1,300	-11.1 ± 1.5
NCGR1309	<i>F. vesca</i>	Italy	1,200	-11.0 ± 1.5
NCGR1364	<i>F. vesca</i>	Epinel, Italy	1,300	-11.0 ± 1.5
Haugstøl	<i>F. vesca</i> ssp. <i>vesca</i>	Haugstøl, Norway	1,080	-10.4 ± 2.0
NCGR198	<i>F. vesca</i>	Hawaii, USA	2,135	-10.4 ± 2.0
FDP815	<i>F. vesca</i> ssp. <i>vesca</i>	Inbred from Baron Solemacher	n/a	-10.3 ± 1.7
NCGR502	<i>F. vesca</i> ssp. <i>bracteata</i>	New Mexico, USA	2,500	-10.3 ± 1.7
Bukammen	<i>F. vesca</i> ssp. <i>vesca</i>	Stjørdal, Norway	250	-9.8 ± 1.5
NCGR1780	<i>F. vesca</i>	Ukraine	n/a	-9.6 ± 1.3
NCGR1001	<i>F. vesca</i>	Ecuador	2,460	-9.2 ± 1.5
NCGR1848	<i>F. vesca</i>	Hokkaido, Japan	180	-8.9 ± 1.3
NCGR522	<i>F. nubicola</i>	Kohistan, Pakistan	2,400	-8.4 ± n/a
FDP701	<i>F. pentaphylla</i>	Wolong Preserve, Sichuan, China	2,400	-8.3 ± 1.6
NCGR1363	<i>F. vesca</i>	Bolivia	n/a	-8.2 ± 1.2
FDP301	<i>F. nubicola</i>	Uttar Pradesh, Pakistan	n/a	-7.7 ± 1.7
FDP817/NCGR371	<i>F. vesca</i> ssp. <i>californica</i>	California, USA	28	-7.7 ± 0.5
NCGR1825	<i>F. nilgerrensis</i>	Yunnan, China	2,100	-6.1 ± 1.9
NCGR1188	<i>F. nilgerrensis</i>	Guizhou, China	1,550	-4.7 ± 3.2

Origin and altitude of collection site, the estimated temperatures for 50 % survival (LT₅₀), and the corresponding standard errors 5 weeks after low-temperature exposure are presented

n/a not available

250 was ensured by replicating some genotypes across experi- 268
 251 ments. To analyze the unbalanced survival data (dead/ 269
 252 alive), the following logistic model was used.

$$\pi_{ijkl} = P(y_{ijkl} = 1 | E_j, (E\alpha)_{ij}) = P(y = 1) \\ = P(\text{a plant survives}) = \frac{e^{\beta_0 + \alpha_i + \beta_1 \cdot t + E_j + (E\alpha)_{ij}}}{1 + e^{\beta_0 + \alpha_i + \beta_1 \cdot t + E_j + (E\alpha)_{ij}}}$$

254 where β_0 is an unknown constant, α_i is the main effect of 272
 255 the genotype i ($i = 1, \dots, 22$), β_1 is the coefficient that 273
 256 estimates the effect temperature (t) has on plant survival, E_j 274
 257 is the effect of experiment or run j ($j = 1, \dots, 6$), k denotes 275
 258 a clonal plant from each genotype in a given experiment, 276
 259 $k = 1, \dots, 12$, t is the temperature plant k is exposed to 277
 260 ($t = -15$ °C to 0 °C), $(E\alpha)_{ij}$ is the interaction between 278
 261 genotype i in experiment j , and π_{ijkl} is the observation 279
 262 [alive (1)/dead (0)] made on plant k from genotype i , in 280
 263 experiment j , exposed to temperature t . 281

264 The LT₅₀ for genotype i was estimated as

$$\hat{E}(\text{LT}_{50}) = -\frac{\hat{\beta}_0 + \hat{\alpha}_i}{\hat{\beta}_1}$$

266 The Glimmix procedure in SAS[®] was used to implement 282
 267 this model. The standard errors for the estimated LT₅₀ 283

values were computed from the covariance matrix using 268
 the delta method (Coles 2001). 269

Protein extraction, Western blot, and quantitative 270
 analysis 271

For the SDS-PAGE and subsequent blot analyses, a subset 272
 of ten genotypes from Table 1 was used (Alta, Bukammen, 273
 FDP817, FDP821, NCGR424, NCGR522, NCGR1363, 274
 NCGR1603, NCGR1780, and NCGR1848). Plant cultiva- 275
 tion was carried out as described previously. Cold treat- 276
 ment was performed at 2 °C for a 10-h photoperiod at 277
 90 μmol m⁻² s⁻¹ for 0, 1, 2, 14 and 42 days. Tissue from 278
 crowns was harvested, immediately shock frozen in liquid 279
 nitrogen and stored at -80 °C until processing. Control 280
 samples (0 day) were harvested prior to the transfer to the 281
 cold room. Each time point was represented by tissue 282
 samples from 12 crowns (4 crowns per replicate). 283

Total protein extracts were isolated from cold-treated 284
 and control crown tissues. Tissue samples (200 mg FW) 285
 were ground to a fine powder in liquid nitrogen and then 286
 extracted with homogenizing buffer composed of 1.5 M 287
 Tris (pH 8.8), 2 % glycerol, 2 % SDS w/v, 2 % 288

289 mercaptoethanol, and 1× Complete Roche Protease inhibitors. The homogenates were then centrifuged for 5 min at
290 10,000g and the supernatants were stored at −80 °C.
291 Loading of the SDS-PAGE was normalized by adding
292 equivalent amounts of protein in each lane. Protein con-
293 centration was estimated using the Amido Black method
294 (Kaplan and Pedersen 1985).
295

296 Proteins extracted from crown tissue (5 µg) were sepa-
297 rated by 12 % SDS-PAGE and transferred to nitrocellulose
298 membranes overnight at 0.2 constant Amps at 4 °C. Mem-
299 branes were blocked and then probed in PBS/5 % non-fat
300 milk (pH 7.4) with either anti-dehydrin antibody (1:2000
301 supplied by Tim Close, UC Riverside, CA, USA) or anti-
302 alcohol dehydrogenase (ADH) (Agrisera, Vannas, Sweden)
303 followed by peroxidase-labeled goat anti-rabbit (1:4,000
304 Sigma®, St Louis, MO, USA). SuperSignal® West Dura
305 (Thermo Scientific, Rockford, IL, USA) was used to visualize
306 chemiluminescence on a ChemiDoc™ XRS Molecular
307 Imager (Bio-Rad). Image analysis and densitometry were
308 performed with ImageJ (NIH IMAGE, <http://rsbweb.nih.gov/ij/>).
309

310 Since the anti-dehydrin antibody had not been used
311 previously in strawberry, experiments confirming speci-
312 ficity, using K-peptide competition, were performed
313 (Suppl. Fig. S1).

314 These data were analyzed statistically and plotted using
315 the *Reg* and the *Sgscatter* procedures in SAS®.

316 Metabolite experiment

317 Since strawberries are propagated by stolons from the crown,
318 a most efficient breeding strategy would be to screen non-
319 essential tissues rather than to destroy the propagule. Thus,
320 we wanted to investigate the potential of using metabolite
321 profiles from leaf tissues to predict low-temperature toler-
322 ance. For this experiment, a subset of ten *F. vesca* genotypes
323 (Table 1) with contrasting freezing tolerance was selected
324 (Alta, Bukammen, FDP817, FDP821, Haugastøl, NCGR13
325 63, NCGR1428, NCGR1603, NCGR1780, and NCGR1848).
326 Twelve-week-old runner-propagated *Fragaria* plants, raised
327 on fertilized soil in plug trays (3 × 6 cells) in a greenhouse at
328 18 ± 2 °C under natural light and long-day conditions, were
329 transferred to a cold storage room at 2 °C under artificial light
330 (fluorescent tubes, 90 µmol m⁻² s⁻¹) for a period of
331 4 weeks. Three mature leaves were sampled from individual
332 plants (*n* = 3 per genotype and time point) at the following
333 time points: 0, 1, 2, 14, and 28 days. The control samples
334 (0 day) were harvested before cold exposure. Samples were
335 immediately shock frozen in liquid nitrogen and stored at
336 −80 °C prior to further processing. A modified extraction
337 and derivatization protocol (Roessner et al. 2001) was uti-
338 lized, based on mechanical sample crushing using a handheld
339 high-speed mixer (300 mg FW). A lyophilized aliquot

(300 µl) was further processed using methoxyamine and
340 trimethylsilyl derivatization. Samples were transferred to
341 1.5 ml autosampler vials with glass inserts and stored at
342 −20 °C prior to analysis by gas chromatography and mass
343 spectrometry (GC–MS).
344

345 An Agilent 6890/5975 GC–MS was used for all analy-
346 ses. Sample volumes of 1 µl were injected with a split ratio
347 of 25:1. GC separations were carried out on an HP-5MS
348 capillary column (30 m × 0.25 mm i.d., film thickness
349 0.25 µm). The injection temperature was 230 °C and the
350 interface was set to 250 °C. The carrier gas was helium at a
351 constant flow rate of 1 ml/min. The GC temperature pro-
352 gram was held isothermally at 70 °C for 5 min, ramped
353 from 70 to 310 °C at a rate of 5 °C/min, and finally held at
354 310 °C for 7 min (analysis time: 60 min). The MS source
355 was adjusted to 230 °C and a mass range of *m/z* 50–700
356 was recorded. All mass spectra were acquired in EI mode.
357 Chromatogram visualization and peak area integration
358 were carried out using the Agilent ChemStation software.
359 For mass spectra evaluation and peak identification, the
360 AMDIS software (v. 2.64) was used in combination with
361 the following mass spectral libraries: NIST05 database and
362 a target library containing MS spectra of trimethylsilylated
363 (TMS) metabolites (Hummel et al. 2010). Numerical
364 analysis was based on peak area integration being corrected
365 for FW variation, using the internal standard ribitol
366 (normalized response). For the statistical analyses, the
367 ribitol-corrected peak areas within each time point were
368 standardized to zero mean and a standard deviation of one
369 for each metabolite.

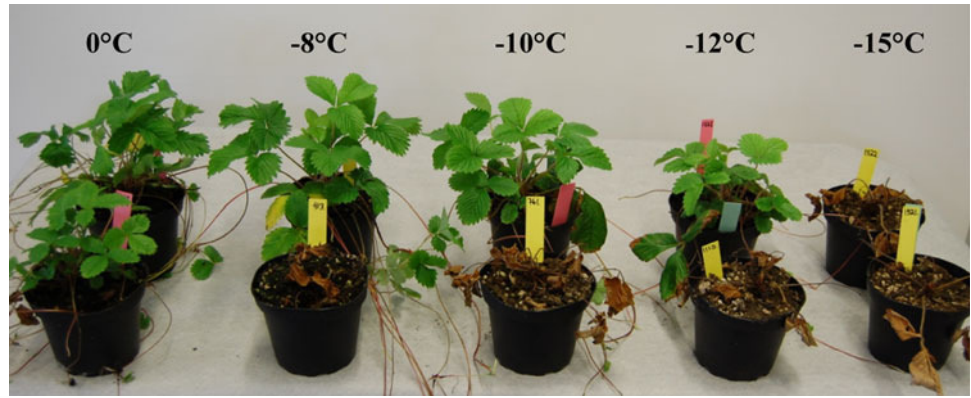
370 A multivariate regression approach was taken to model the
371 LT₅₀ estimates using the metabolite data at 28 days of
372 acclimation. *Proc Reg* (SAS Institute Inc. 2008) with the
373 *stepwise* option was used for this. In order to reveal structures
374 in the metabolite data that could be associated with the impact
375 of the acclimation period or with the specific genotype, we
376 used principal component analyses (PCA) including all the 13
377 compounds observed at time points 0 and 28. The SAS®
378 *Princomp* procedure was used for the PCA, and the *Sgplot*
379 procedure was used for generating the PC loading plot (SAS
380 Institute Inc. 2008). Finally, heat maps were made to visu-
381 alize structures and metabolic responses to cold acclimation.
382 For this, the *heatmap.2* function in *R* (<http://www.r-project.org>) was used.
383

384 Results

385 Freezing tests of 22 genotypes

386 Typical results of the freezing tests are shown in Fig. 1 where
387 one of the low-temperature-tolerant genotypes (*F. vesca* ssp.
388 *bracteata*, NCGR424) and one low-temperature-sensitive

Fig. 1 Typical result from low-temperature stress experiments. A frost-tolerant accession (NCGR424, rear, *F. vesca* ssp. *bracteata*) and a frost-susceptible (FDP 817, front, *F. vesca* ssp. *californica*) accession of *F. vesca* exposed to five levels of freezing stress. The plants had been grown at 18 °C in the greenhouse for 5 weeks after low-temperature exposure when the picture was taken



389 genotype (*F. vesca* ssp. *californica*, FDP817) are presented. 425
 390 The estimated LT_{50} values (temperature at which 50 % of 426
 391 plants survived) and their corresponding standard errors are 427
 392 presented in Table 1. 428

393 In general, there was a negative ($r = -0.47$) and sig- 429
 394 nificant ($P = 0.04$) correlation between LT_{50} estimates and 430
 395 geographical latitude. The correlation to altitude was, 431
 396 however, not significant. 432

397 The LT_{50} estimates have a range from -4.7 (NCGR1188) 433
 398 to -12.0 °C (FDP821 and NCGR1428). NCGR1188 is an 434
 399 *F. nilgerrensis*, while the two most tolerant are *F. vesca* 435
 400 species. In general, the *F. vesca* genotypes seem to be more 436
 401 low-temperature tolerant than the other species tested. In 437
 402 particular, the three *F. vesca* ssp. *bracteata* genotypes were 438
 403 all in the low-temperature-tolerant side of the distribution 439
 404 (Table 1), while the two *F. nilgerrensis* genotypes appeared 440
 405 on the susceptible side of the same distribution, to some 441
 406 extent together with the *F. nubicola* genotypes. Regarding 442
 407 their average LT_{50} values, *F. vesca* ssp. *bracteata* (Avg LT_{50} 443
 408 -11.1) differ significantly from the value of the one *F. vesca* 444
 409 ssp. *californica* genotype ($LT_{50} -7.7$, $P = 0.003$), the 445
 410 average of the two *F. nilgerrensis* genotypes (Avg $LT_{50} -5.4$, 446
 411 $P < 0.0001$), and from the average of the two *F. nubicola* 447
 412 genotypes (Avg $LT_{50} -8.1$, $P = 0.0002$). Finally, the 448
 413 *F. nilgerrensis* average also differs significantly ($P = 0.0003$) 449
 414 from the *F. vesca* ssp. *vesca* average ($LT_{50} -10.5$). 450

415 The NCGR1363 is another low-temperature susceptible 451
 416 *F. vesca* genotype ($LT_{50} -8.2$). Pair-wise tests showed that 452
 417 Alta ($LT_{50} -11.6$) was significantly different from both 453
 418 FDP817 ($LT_{50} -7.7$) and NCGR1363. Hence, these are 454
 419 excellent candidates for parent mapping populations. 455

420 Alcohol dehydrogenase and dehydrin levels

421 Western blotting and probing with anti-dehydrin (K-seg- 456
 422 ment specific) for the full time course sample series 457
 423 (noncold-acclimated treated control, 1, 2, 14, and 42 days 458
 424 cold) was carried out for eight *F. vesca* genotypes with 459

425 three biological replicates each (Fig. 2). Dehydrins were 426
 427 not detected in the untreated control or in the 1-day and 428
 429 2-day cold-treated crowns. Interestingly, dehydrins in the 429
 430 leaves could not be detected at any time points (data not 430
 431 shown). However, three bands were first observed at 431
 432 14 days, which accumulated to much higher levels at 432
 433 42 days (Fig. 2). This should be considered a relatively 433
 434 slow cold response, particularly relative to *Arabidopsis* 434
 435 where dehydrin levels are readily detected by 2 days and 435
 436 are at near maximum at 4–6 days after initiation of cold 436
 437 treatment. Competition experiments (Suppl. Fig. S1) 437
 438 showed that all bands represented true dehydrins as they 438
 439 were competed by the K-peptide. Four distinct patterns of 439
 440 dehydrin expression were observed in the genotypes and 440
 441 were exemplified by FDP821, Alta, NCGR522, and 441
 442 NCGR1603 (Suppl. Fig. S2). The dehydrin masses were 442
 443 extrapolated from the competition experiment (Suppl. Fig. S1). 443
 444 Bioinformatic analysis identified seven distinct dehydrins 444
 445 (Suppl. Fig. S3). Application of antibodies specific to 445
 446 *Arabidopsis* dehydrins revealed multiple polypeptides, 446
 447 confounding identification of specific *Fragaria* orthologs 447
 448 (Suppl. Fig. S2). The total dehydrin content (obtained by 448
 449 summing all K-peptide antibody-reactive bands) after 449
 450 14 days of cold acclimation was not correlated to the LT_{50} 450
 451 values (data not shown); however, a strong correlation was 451
 452 evident at 42 days ($r = -0.81$, $P < 0.0001$; Fig. 3a). 452

453 Our own preliminary observations in the octoploid 453
 454 *F. × ananassa* indicated that there was a high correlation 454
 455 between alcohol dehydrogenase (ADH) levels and cold 455
 456 tolerance as indicated by LT_{50} values (Koehler et al. 2012). 456
 457 In the present experiments with *F. vesca*, ADH levels were 457
 458 very low in control crowns (not shown), but strongly 458
 459 induced in the cold-treated crowns. For some genotypes 459
 460 and in particular the ones that turned out to possess the 460
 461 highest tolerance to cold, a 200-fold increase in ADH 461
 462 protein levels was observed after 42 days of cold treatment, 462
 463 relative to the controls (Figs. 3b, 4). The correlation 463
 464 between the estimated LT_{50} values and the ADH 464

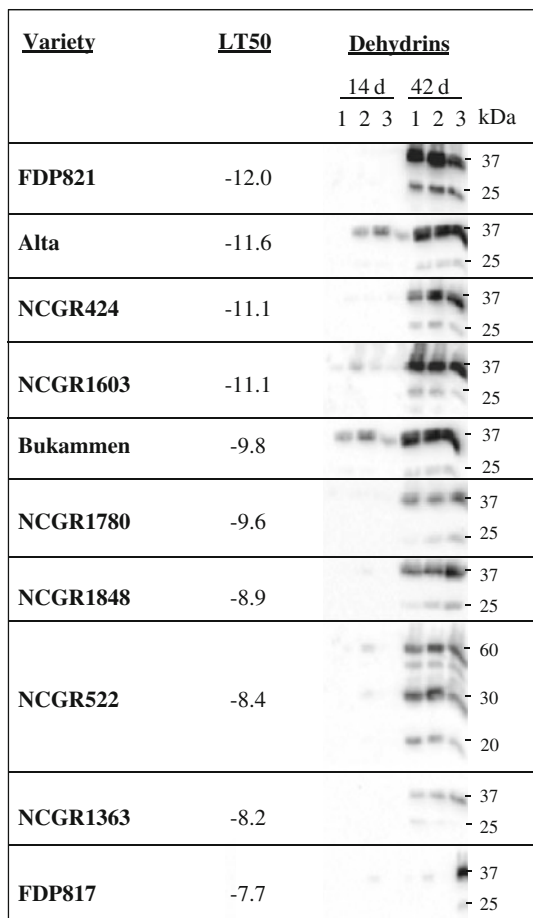


Fig. 2 Dehydrin levels in ten *Fragaria* genotypes. Extracts of crowns in three biological replicates from plants treated for 0 h, and 1, 2, 14, and 42 days at 2 °C were separated on 12 % SDS-PAGE and probed with anti-K peptide. Probing and visualization were done in two groups. Antibody-reactive bands appeared only for the 14 and 42 days cold-treated samples, and only blot sections with these samples are shown

463 expression levels after 6 weeks of cold acclimation was
464 significant with an $r = -0.86$ ($P < 0.0001$; Fig. 3b).

465 The relationship between dehydrins or ADH levels and
466 cold tolerance in *F. vesca* genotypes revealed by linear
467 regression (Fig. 3) did not include two of the genotypes tested
468 in the Western blots (Figs. 2, 4). *F. nubicola*, while of interest
469 for its low cold tolerance and dehydrin expression, repre-
470 sented a distinct species from *F. vesca*. FDP821, a distinct
471 subspecies which did not produce any fertile hybrids when
472 used as a parent in hybridization experiments with *F. vesca*
473 (not shown), suggesting significant chromosomal differences
474 or an efficient incompatibility system between FDP821 and all
475 the other *F. vesca* genotypes, was also not included in the
476 regression analysis. It was interesting; however, that FDP821,
477 the most cold tolerant of the tested genotypes, showed the
478 highest levels of dehydrin accumulation, but relatively low
479 expression of ADH, though not as low as the least cold-tol-
480 erant genotypes. *F. nubicola* (NCGR522), a cold-susceptible

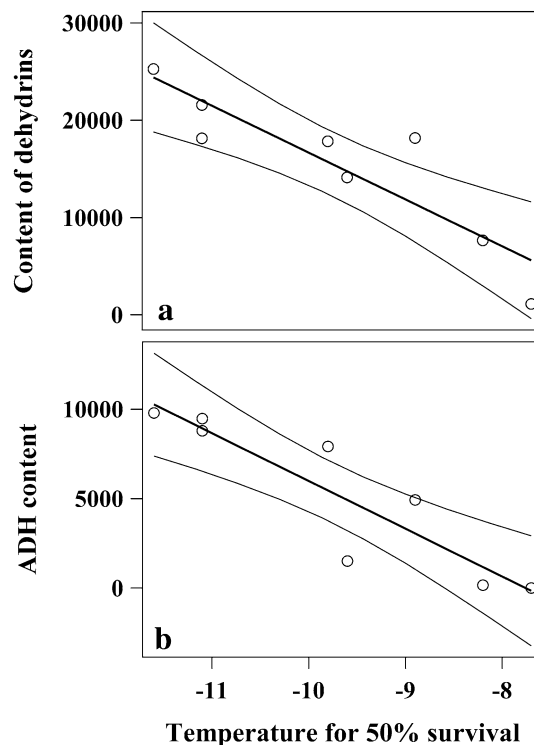


Fig. 3 Linear regression line fit between dehydrin content (a, $R^2 = 0.67$, $r_{\text{dehyd}} = -0.82$, $P < 0.0001$), alcohol dehydrogenase content (b, $R^2 = 0.74$, $r_{\text{adh}} = -0.87$, $P < 0.0001$), and the LT₅₀ estimates. The 95 % confidence intervals are indicated. Protein levels are expressed as dimensionless arbitrary values. Each data point is the average of three measurements. Only *F. vesca* genotypes were used for these correlations. When all the genotypes were included in the regression analyses, the R^2 values fell to $R^2_{\text{dehyd}} = 0.24$ ($r = -0.49$, $P = 0.0034$) and $R^2_{\text{adh}} = 0.47$ ($r = -0.69$, $P < 0.0001$)

genotype, had a moderate but distinctive dehydrin expression 481
482 pattern and no detectable ADH. However, when these geno-
483 types were included in the regression analyses, the R^2 obtained
484 were $R^2_{\text{dehyd}} = 0.24$ ($r = 0.49$, $P = 0.0034$), and $R^2_{\text{adh}} = 0.47$
485 ($r = 0.69$, $P < 0.0001$).

Metabolite profiling 486

487 From the table of means (Table 2), there seem to be dif-
488 ferent patterns of leaf metabolic responses across the time
489 points. Metabolites like fumaric acid, aspartic acid, glu-
490 tamic acid, asparagine, citric acid, galactose, sucrose, and
491 raffinose by and large show an increase in content during
492 the whole acclimation period. Others in general decrease
493 toward the last time point (succinic acid, malic acid,
494 fructose, and glucose), and finally there are metabolites that
495 do not seem to change much as the acclimation proceeds
496 (galactinol). These general patterns are, however, fre-
497 quently broken by local peaks or troughs, e.g., the galact-
498 ose content at day 14 (Table 2).

499 Some of the metabolites show significant positive cor-
 500 relations to the LT₅₀ estimates (Table 3). A positive cor-
 501 relation would indicate that at the particular time point, the
 502 content of the metabolite is lower for the more cold-tol-
 503 erant genotypes. The significant correlations observed for
 504 succinic acid at four of the five time points is to a large
 505 extent caused by the relatively high content of succinic acid
 506 in the low-tolerant genotype, FDP817 (Suppl. Table S1).

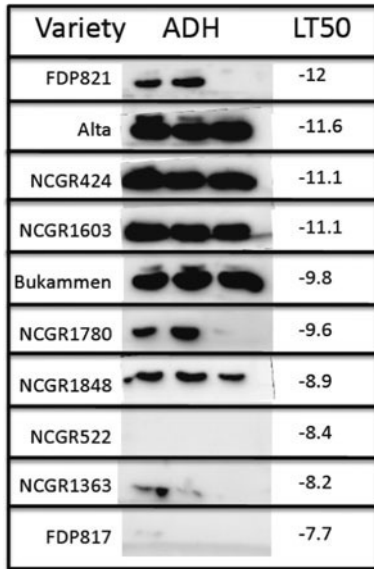


Fig. 4 Alcohol dehydrogenase (ADH) protein levels in *F. vesca* genotypes. Extracts of crowns from plants treated for 6 weeks at 2 °C were separated on 12 % SDS-PAGE and then probed with anti-ADH. Since bands were not visible for controls (0 h at 2 °C) at this exposure, they are not shown. Gels were all blotted onto the same nitrocellulose paper and thus probed simultaneously with antibodies. For each variety, triplicates are shown

The lack of correlation between the raffinose content and LT₅₀ estimates is notable, but consistent across all time points (Table 3). On the other side, both sucrose and galactinol correlate well with LT₅₀, at least at some of the early time points.

One of our goals in the current work was to identify compounds that could be correlated to low-temperature tolerance (LT₅₀ estimates). We expected plants to be fully acclimated after 28 days of low-temperature exposure and choosing this time point for our multivariate data analyses seemed natural. In the multiple regression analysis approach using data from day 28, only the galactinol content was retained as the only significant ($R^2 = 0.63$, $P < 0.0001$) explanatory variable for the variation in low-temperature tolerance (LT₅₀). The content of raffinose showed no such co-variation with the LT₅₀ estimates. The linear regression lines for the raffinose and the galactinol contents are presented in Fig. 5.

The PCA of the metabolite data from before the acclimation started (day 0) and at the end of the acclimation period (day 28) showed that five components were required to account for 90 % of the total variation in the metabolite data matrix. The first principal component had contributions from most of the metabolites, but not from succinic acid, malic acid, fructose, and glucose. The component loadings varied between 0.36 and 0.42. The second principal component was dominated by glucose, fructose, and galactinol.

The plot of the two first PC axes is given in Fig. 6. At the starting point (day 0), all the genotypes cluster relatively well together. After 28 days of cold acclimation, their metabolite profiles had become much more heterogeneous and spread in various directions. This response

Table 2 Leaf metabolite changes averaged across ten diploid *Fragaria* genotypes during acclimation at 2 °C

Metabolite	Abbrev.	Days in cold acclimation					
		0	Content of metabolite relative to day 0				
			0	1	2	14	28
		$\mu\text{g g}^{-1}$ FW					
Succinic acid	SucA	44.7 ± 5.0	100	46	58	33	36
Fumaric acid	FumA	23.7 ± 3.9	100	73	84	100	224
Malic acid	MalA	1,913 ± 171	100	150	86	77	55
Aspartic acid	AspA	19.5 ± 5.4	100	204	344	590	806
Glutamic Acid	GluA	74.7 ± 16.7	100	111	180	475	576
Asparagine	Asp	6.5 ± 3.5	100	52	866	2,616	4,526
Citric acid	CitA	1,824 ± 93	100	164	109	144	247
Fructose	FruS	1,130 ± 75	100	159	119	83	63
Galactose	GalS	7.9 ± 3.5	100	1,329	4,000	7,089	3,177
Glucose	GluS	574 ± 24	100	170	113	87	79
Sucrose	SucS	16,202 ± 490	100	452	121	126	184
Galactinol	Galact	141 ± 10	100	257	112	140	116
Raffinose	RafS	309 ± 24	100	249	128	389	520

Metabolite content at initiation was set to 100 % and percent increases/decreases are relative to these initial values. Actual contents in $\mu\text{g g}^{-1}$ FW and the corresponding standard errors at day 0 are also presented. An extended table of the metabolite contents is given in Suppl. Table S1

Table 3 Pearson correlation coefficients (r) of selected metabolites versus LT_{50} values for ten diploid *F. vesca* genotypes from cold acclimation trials at 2 °C over a period of 4 weeks (sample material: leaf)

Metabolite	0	1 day	2 days	14 days	28 days
Succinic acid	0.63***	0.63***	0.62***	0.52**	0.15
Fumaric acid	0.46*	0.12	0.23	0.40*	0.02
Malic acid	0.05	0.25	0.17	0.29	0.31
Aspartic acid	0.04	-0.23	-0.17	-0.11	-0.14
Glutamic acid	-0.01	-0.14	-0.14	-0.11	-0.13
Asparagine	-0.01	-0.30	0.19	-0.02	-0.25
Citric acid	0.51**	0.35	-0.15	0.27	-0.13
Fructose	-0.09	-0.14	-0.03	-0.01	0.31
Galactose	-0.28	-0.14	-0.09	0.08	0.21
Glucose	-0.03	0.10	-0.03	0.03	0.53**
Sucrose	0.51**	0.68***	0.42*	0.29	0.29
Galactinol	0.52**	0.49**	0.30	0.27	0.79***
Raffinose	-0.04	-0.16	-0.22	0.29	0.24

* $P < 0.05$
 ** $P < 0.01$
 *** $P < 0.001$

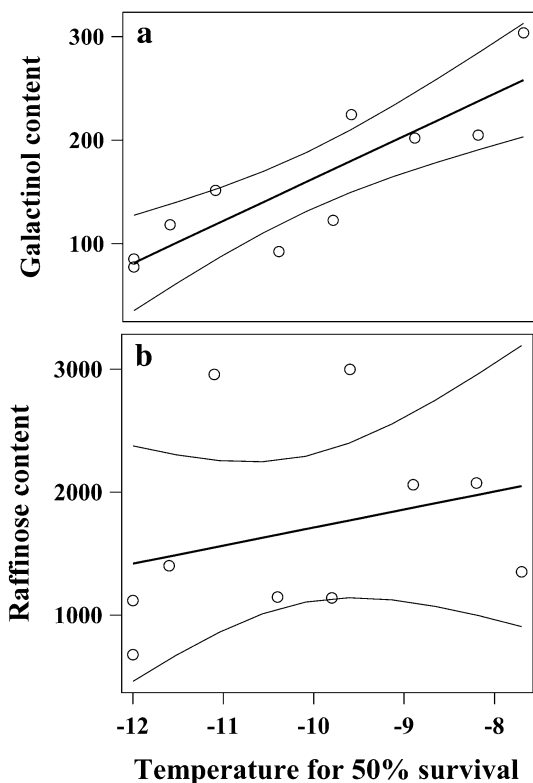


Fig. 5 Linear regression line fit between the LT_{50} estimates and galactinol (a, $R^2 = 0.63$, $r_{galact} = 0.79$, $P < 0.0001$) and raffinose content (b, $R^2 = 0.06$, $r_{raff} = 0.24$, $P = 0.24$) after 28 days of cold acclimation at 2 °C. The 95 % confidence intervals are indicated. Each data point is an average of three measurements and the sugar content unit is $\mu\text{g/g FW}$. Only *F. vesca* genotypes were used for these regression plots

540 due to acclimation is also illustrated in the heat maps of the
 541 two time points (Suppl. Fig. S4). FDP817 and NCGR1780
 542 appear to respond in a similar way, while the remaining
 543 genotypes form a more or less diffuse cluster (Fig. 6). The

one exception is possibly the Alta genotype. It seems to 544
 545 behave differently from the other genotypes by showing
 546 little movement or regrouping caused by acclimation
 547 (Fig. 6).

Since Fig. 6 only depicts two of the five axes necessary to 548
 549 account for the bulk variation (>90 %) in the metabolite data,
 550 a more nuanced illustration is provided by the heat map in
 551 Fig. 7. Here, the simultaneous hierarchical clustering of the
 552 two time points is presented as a heat map. There appear to be
 553 two genotype clusters that mainly consist of genotypes before
 554 acclimation (0 days) and those after acclimation (28 days).
 555 However, a couple of exceptions are notable. The Alta
 556 genotype appears in the same cluster both before and after
 557 acclimation, in agreement with Fig. 6, indicating that the
 558 acclimation results in only subtle changes in this genotype's
 559 metabolite composition. Alta originates from the very
 560 northern parts of Norway and has been shown to respond
 561 differently also in other traits, e.g., in response to flowering
 562 stimuli (Heide and Sønsteby 2007). Moreover, at time point
 563 28, the NCGR1848 and FDP817 are clustered together with
 564 the bulk of the time point 0 entries. Figure 7 indicates that
 565 these entries do not respond typically to the acclimation. For
 566 instance, they lack the accumulation of aspartic acid, glu-
 567 tamic acid, and asparagine observed after the acclimation
 568 period in the majority of the entries (Fig. 7). Finally,
 569 NCGR1780 also responds atypically (Figs. 6, 7). The
 570 implication is that there are varying responses to acclimation
 571 at the metabolic level, and given the diverse origin of the
 572 accessions, this may not be surprising.

As for the metabolites, there seem to be two or three 573
 574 structures in their responses to the cold acclimation
 575 (Fig. 7). The first cluster consists of fructose, glucose,
 576 succinic acid, malic acid, and galactinol, and their content
 577 is reduced toward the end of the acclimation period. A
 578 second cluster consists of aspartic acid, glutamic acid,
 579 citric acid, and asparagine, and these metabolites are in

621 deep cold hardiness. The 6 weeks we used as acclimation
622 may thus be too short to attain the full level of hardiness.
623 We considered the ranking of the genotypes to be our
624 major goal and, although we could not dismiss the possi-
625 bility of genotypes interacting with other environmental
626 factors resulting in shifts in ranking, we expect that such
627 possible shifts would be of minor importance.

628 The range in the LT_{50} values we found for *F. vesca* is in
629 some agreement with observations in *F. × ananassa*
630 (Marini and Boyce 1977, 1979) where normally hardened
631 plants are slightly wounded at -4 °C. Severe injuries
632 occurred at -12 °C, but survival was observed with crown
633 temperatures reaching -20 °C. These authors used a
634 slightly different testing regime, i.e., they only allowed the
635 plants to reach the target temperature and then removed
636 them immediately to thaw in the greenhouse, while we kept
637 the target temperature for 4 h and then slowly, with con-
638 trolled temperatures, returned plants to above freezing
639 temperatures, and finally to 18 °C growing conditions.

640 *Fragaria* dehydrins and alcohol dehydrogenase
641 are associated with low-temperature tolerance

642 Dehydrins are known to be involved in response to a wide
643 range of abiotic stresses, such as cold, drought, and salt
644 stress (Campbell and Close 1997). Dehydrins are well
645 conserved between the plant genera and homologs are
646 readily identified based on sequence similarity and in
647 particular by the presence of the K-segment, a signature
648 dehydrin consensus sequence. However, Koehler et al.
649 (2012) identified changes in dehydrin transcript levels
650 during cold acclimation in two Norwegian *F. × ananassa*
651 cultivars, Frida and Jonsok. These two cultivars differ in
652 their low-temperature tolerance—Jonsok being the most
653 tolerant one. The XERO2-like dehydrin increased in both
654 cultivars during the acclimation period, but to a much
655 larger extent in Jonsok. While the COR47-like dehydrin
656 transcript decreased with acclimation, the level in Jonsok
657 before the acclimation period was much higher than the
658 less cold-tolerant cultivar (Koehler et al. 2012).

659 In other species within Rosaceae, dehydrins have been
660 identified to have a high similarity to *Arabidopsis* dehyd-
661 rins. In peach, COR47-like (Bassett et al. 2009) and
662 XERO2-like dehydrins have been found (Artlip et al.
663 1997), and in apple, an ERD10-like dehydrin (Garcia-
664 Bañuelos et al. 2009). To obtain a better understanding of
665 the dehydrin family of proteins in strawberry, we based our
666 bioinformatic analyses (Suppl. Fig. S3) on the recently
667 published *F. vesca* genomic sequence (Shulaev et al. 2011).
668 We predicted that the *Arabidopsis*-derived dehydrin anti-
669 bodies used in our present experiments were likely to react
670 with the predicted *Fragaria* dehydrin proteins (Suppl.
671 Figs. S2 and S3).

All dehydrin bands correlate with the LT_{50} ; however, for
the eight *F. vesca* genotypes the total level of dehydrin at 6
weeks was highly correlated ($r = -0.81$) with LT_{50} . This
makes the overall dehydrin content a very good candidate
for a freezing tolerance protein marker. That increased
dehydrin expression is sufficient to increase frost tolerance
was previously shown by transforming a *F. × ananassa*
cultivar with the wheat dehydrin gene WCOR410. Freezing
tolerance, as measured by the electrolyte leakage test,
increased by -5 °C compared to the wild type (Houde
et al. 2004). In blueberry stem and leaf tissue, two varieties
of differing cold hardiness were compared (Danyluk et al.
1994), and in agreement with our results, the most winter
hardy variety showed the strongest induction of dehydrin,
both at the protein and mRNA levels. A positive correla-
tion between a dehydrin and freezing tolerance was also
found in a segregating F_2 population of *Rhododendron*
(Lim et al. 1999).

We examined alcohol dehydrogenase as it can enhance
stress survival by ameliorating hypoxic conditions brought
on by melting snow or ice encasement. Thus by increasing
the glycolytic fermentation pathways and shifting the end
point away from lactate and toward ethanol (Drew 1997),
elevated levels of ADH can prevent accumulation of toxic
end products of anaerobic metabolism, preventing injury
and thus increasing winter survival. Based on the high
correlation of ADH levels with LT_{50} ($r = -0.86$), it is
likely that ADH contributes to cold hardiness in *F. vesca*.
This protein is thus a very good candidate as a molecular
marker for cold stress tolerance.

Central metabolites in the leaf showed correlation
to LT_{50} -based freezing tolerance

Sucrose accumulation in response to cold exposure is a
common observation and is a result of the increased
activity of sucrose phosphate synthase and sucrose syn-
thase (Sasaki et al. 2001). Recently, Schulze et al. (2011)
observed significant increases in leaf content of glucose,
fructose, and sucrose during cold acclimation of *A. thali-*
ana, in agreement with other authors (Cook et al. 2004;
Kaplan et al. 2007; Guy et al. 2008) and also in accordance
with our overall response observations. But a closer look
showed that the genotypes responded differently as also
reported in our earlier study (Rohloff et al. 2012). If we
look at the correlations between these sugars and the sur-
vival rate of the plants after cold exposure, the LT_{50} esti-
mates, there is a positive correlation to the sucrose.
So, even though on an overall basis there is a significant
accumulation of the mono- and disaccharides (i.e., galact-
ose, sucrose, and raffinose), the positive correlation
between sucrose levels and LT_{50} at the beginning of the
acclimation period indicates that the genotypes with the

723 lowest sucrose content are the most low-temperature tol-
724 erant ones.

725 Raffinose is often found up-regulated in other plant
726 species, e.g., *Arabidopsis thaliana*, during cold acclimation
727 (e.g., Korn et al. 2010), and this was also the case with our
728 material (Table 2). However, we did not observe a signif-
729 icant correlation to the LT₅₀ estimates at any time point
730 during acclimation (Table 3). It has been shown, however,
731 that raffinose accumulation is neither necessary nor suffi-
732 cient for the induction of freezing tolerance in *A. thaliana*
733 (Zuther et al. 2004). While our present results (Table 2)
734 and previous ones (e.g., Saito and Yoshida 2011; Rohloff
735 et al. 2012) show that both raffinose and galactinol contents
736 are enhanced during acclimation (i.e., the raffinose path-
737 way), only galactinol content showed a significant correla-
738 tion to cold stress tolerance in our study (Fig. 5).
739 Moreover, this correlation was positive, implying a rela-
740 tively lower level of metabolite in the hardiest genotypes.

741 The majority of studies on low-temperature tolerance
742 have been conducted with *Arabidopsis thaliana*, which
743 survives winter either as a small plantlet (winter annual) or
744 as seed. The strawberry, however, prepares for winter by
745 senescence and translocation of the majority of assimilates
746 to the crown. Could this explain why we, for instance,
747 observe that the most cold-tolerant genotypes exhibit the
748 lowest levels of galactinol after acclimation? Is it because
749 these are the genotypes that most efficiently transport the
750 solutes to the crown in preparation for winter? Our ongoing
751 research addresses these issues.

752 To examine the natural variation in cold/freezing toler-
753 ance, 22 diploid *Fragaria* genotypes were acclimated and
754 then tested to obtain plant survival estimates (LT₅₀). Cor-
755 relation of plant survival with leaf metabolite profiles and
756 with the expression of dehydrin and alcohol dehydrogenase
757 proteins in the crown during acclimation indicated that the
758 proteins and the sugar alcohol galactinol showed the
759 clearest association with cold tolerance and thus the
760 greatest potential to be developed into biomarkers.

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773 **Conflict of interest** The authors declare that they have no conflict
774 of interest.

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