Vernalization Requirement and the Chromosomal VRN1-Region can Affect Freezing Tolerance and Expression of Cold-Regulated Genes in Festuca pratensis

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Plants adapted to cold winters go through annual cycles of gain followed by loss of freezing tolerance (cold acclimation and deacclimation). Warm spells during winter and early spring can cause deacclimation, and if temperatures drop, freezing damage may occur. Many plants are vernalized during winter, a process making them competent to flower in the following summer. In winter cereals, a coincidence in the timing of vernalization saturation, deacclimation, downregulation of cold-induced genes, and reduced ability to reacclimate, occurs under long photoperiods and is under control of the main regulator of vernalization requirement in cereals, VRN1, and/or closely linked gene(s). Thus, the probability of freezing damage after a warm spell may depend on both vernalization saturation and photoperiod. We investigated the role of vernalization and the VRN1-region on freezing tolerance of meadow fescue (Festuca pratensis Huds.), a perennial grass species. Two F2 populations, divergently selected for high and low vernalization requirement, were studied. Each genotype was characterized for the copy number of one of the four parental haplotypes of the VRN1-region. Clonal plants were cold acclimated for 2 weeks or vernalized/cold acclimated for a total of 9 weeks, after which the F2 populations reached different levels of vernalization saturation. Vernalized and cold acclimated plants were deacclimated for 1 week and then reacclimated for 2 weeks. Flowering response, freezing tolerance and expression of the cold-induced genes VRN1, MADS3, CBF6, COR14B, CR7 (BLT14), LOS2, and IRI1 was measured. We found that some genotypes can lose some freezing tolerance after vernalization and a deacclimation–reacclimation cycle. The relationship between vernalization and freezing tolerance was complex. We found effects of the VRN1-region on freezing tolerance in plants cold acclimated for 2 weeks, timing of heading after 9 weeks of vernalization, expression of COR14B, CBF6, and LOS2 in vernalized and/or deacclimated treatments, and restoration of freezing tolerance during reacclimation. While expression of VRN1, COR14B, CBF6, LOS2, and IRI1 was correlated, CR7 was associated with vernalization requirement by other mechanisms, and appeared to play a role in freezing tolerance in reacclimated plants.

Keywords: CBF6, COR14B, CR7, deacclimation, meadow fescue, IRI1, LOS2, photoperiod
INTRODUCTION

Overwintering temperate plants cold acclimate in the autumn and develop resistance to freezing damage. Upon exposure to warmer temperatures in spring plants deacclimate and gradually lose this resistance (reviewed by Kalberer et al., 2006; Rapacz et al., 2014). Warm spells in mid-winter or early spring can cause deacclimation when there is still a risk of freezing temperatures. To some extent, depending on circumstances, plants have the ability to reacclimate if temperatures drop again. The annual variation in freezing tolerance is one of many developmental processes that are regulated largely by temperature (Penfield, 2008). Deacclimation and reacclimation processes are highly complex, and although temperature is a main driving force, other environmental and physiological conditions have strong influence. Resistance to deacclimation and/or the ability to reacclimate is thought to be crucial for plant winter survival in areas with a variable winter climate and temperatures fluctuating around the freezing point. In the face of global warming, where a higher frequency of warm spells during winter can be expected (Shabbar and Bonsal, 2003; Johansson et al., 2011), understanding these processes in plants is important both in an agricultural context (Gu et al., 2008; Bokhorst et al., 2009; Rapacz et al., 2014).

The relationships between vernalization (the process of becoming competent to flower after a prolonged period of cold) and freezing tolerance, and to some extent photoperiod, have been particularly studied in cereals. In these species, it has been shown that freezing tolerance and expression of genes involved in freezing tolerance are down-regulated in leaf and stem base tissue when the vernalization requirement is saturated, but before any development of the apex is visible in the microscope (Fowler et al., 1996; Limin and Fowler, 2006; Laudencia-Chingcuanco et al., 2011). There is an interaction between vernalization and photoperiod on this deacclimation and also on the ability to reacclimate. In cultivars with a long day requirement for flowering, the negative effect of vernalization on freezing tolerance is stronger when plants are vernalized under long days than under short days, whereas vernalization- and photoperiod-insensitive cultivars are not able to develop much freezing tolerance at all (Mahfoozi et al., 2001a, 2005, 2006; Dhillon et al., 2010). Also, plants vernalized and deacclimated under long days are less able to reacclimate (Mahfoozi et al., 2001b). VRN1 is an inducer of the transition to generative development in cereals and other temperate grass species (reviewed by Trevaskis, 2010; Fjellheim et al., 2014). It is gradually upregulated during vernalization and appears to act in the down-regulation of freezing tolerance genes in vernalized plants under long days (Fowler et al., 1996; Limin and Fowler, 2006; Dhillon et al., 2010; Laudencia-Chingcuanco et al., 2011). It is not entirely clear, however, whether it is VRN1 itself, or a very closely linked gene, that is responsible. It is also not known how long days interact with the VRN1 locus in down-regulation of freezing tolerance genes. Mahfoozi et al. (2005, 2006) suggested that in regions with long, mild winters, mechanisms extending the vegetative phase (through vernalization and/or photoperiod requirements) might actually be more important for winter survival than a high maximum attainable freezing tolerance. This is increasingly relevant for winter cereals and perennial grasses in the context of climate change in Northern areas, where winters will become milder, but remain dark.

There are several reports describing deacclimation and reacclimation in response to various treatments in perennial grass species (Tronsmo, 1985; Gay and Eagles, 1991; Tompkins et al., 2000; Jørgensen et al., 2010; Espevig et al., 2014; Hoffmann et al., 2014). These have, however, not specifically tested the effect of vernalization or vernalization requirement on deacclimation and reacclimation, or characterized accompanying changes in gene expression. Here, we addressed these aspects by studying freezing tolerance during a cold acclimation (CA)/vernalization – deacclimation cycle in genetic material of the perennial forage grass species Festuca pratensis Huds. (meadow fescue), divergently selected for high or low vernalization requirement. We coupled this with measurements of flowering response and expression of VRN1 and MADS3 (a VRN1-like gene), and genes known to play a role in CA in F. pratensis and other temperate grass species (Table 1).

MATERIALS AND METHODS

Plant Material and Growth Conditions

Genotypes of two F1-populations from the “HF2/7 × BF14/16” F1 mapping population of Festuca pratensis Huds. (Alm et al., 2003), VRmin and VRmax, were studied. These two populations were produced by crossing F1 individuals selected for either high or low vernalization requirement in two separate groups (Ergon et al., 2013). VRMin segregates for the ability to flower without vernalization, while VRMax requires 9 or more weeks of vernalization in order to flower. Seeds were sown in November 2011 and the plants were grown in the greenhouse under non-vernalizing temperatures and 12 h photoperiod. Over the summer the pots were kept outdoors with natural light conditions. In late August 2012, twenty-one and six genotypes of VRmin and VRmax, respectively, were clonally propagated and pregrown in the greenhouse. A higher number of VRmin genotypes were included due to the segregation for the ability to flower without vernalization in this population. The plants were first grown at 15/18°C (day/night), 12 h photoperiod and approximately 100 μmol m−2 s−1 PAR for 2 months and cut and fertilized at 1 or 2 weeks intervals. After this the plants were grown for another 2 months with approximately 200 μmol m−2 s−1 PAR, and not cut, but fertilized weekly. After pregrowth, the plants were exposed to four different temperature treatments (Figure 1): CA at 2°C for 2 weeks, V-CA: vernalization and CA at 6°C for 7 weeks followed by 2 weeks at 2°C, DA: V-CA followed by deacclimation at 12°C for 1 week, RA: V-CA and DA followed by reacclimation for 2 weeks at 2°C. All treatments had a 8 h photoperiod and a light intensity of approximately 250 μmol m−2 s−1.
TABLE 1 | Genes included in the gene expression analysis.

| Gene  | Protein                          | Expression                                      | Function                                                                 | Reference                                                                 |
|-------|----------------------------------|-------------------------------------------------|--------------------------------------------------------------------------|----------------------------------------------------------------------------|
| VRN1  | MADS-box transcription factor    | Induced by prolonged cold.                      | Induces transition to generative development                             | Yan et al., 2003; Jensen et al., 2005; Ergon et al., 2008; Trevaskis, 2010 |
|       |                                  | Induced by long photoperiod under certain        |                                                                          |                                                                            |
|       |                                  | circumstances.                                  |                                                                          |                                                                            |
| MADS3 | VRN1-like MADS-box transcription factor | Associated with transition to generative         | Unknown                                                                  | Schmitz et al., 2000; Petersen et al., 2004; Preston and Kellogg, 2008;  |
|       |                                  | development                                     |                                                                          | Ergon et al., 2013                                                        |
| CBF6  | Member of family of AP2/EREBP     | Induced rapidly by cold                         | Induces cold-regulated genes with CRT/DRE promoter element.              | Xiong and Fei, 2006; Tamura and Yamada, 2007; Galiba et al., 2009; Alm et al., 2011; Rudi et al., 2011; Sandve et al., 2011; Jurczyk et al., 2013a |
|       | transcription factors            |                                                 |                                                                          |                                                                            |
| COR14B| Soluble protein localized in the stroma compartment of the chloroplast | Induced by CBFs. Regulated by light. Some expression also in etiolated tissue and stem base. | Unknown                                                                  | Crosatti et al., 1995, 1999, 2003; Dal Bosco et al., 2003; Galiba et al., 2009; Rudi et al., 2011; Jurczyk et al., 2013a |
| CR7 (BLT14) | Member of a family of proteins predicted to be secreted into the apoplastic | Cold-induced                                   | Unknown                                                                  | Phillips et al., 1997; Pearce et al., 1998; Rudi et al., 2011 |
| LOS2  | Bifunctional enolase and         | Cold-induced                                    | Positive regulator of cold-induced genes                                 | Lee et al., 2002; Rudi et al., 2011; Jurczyk et al., 2013a |
|       | transcription factor             |                                                 |                                                                          |                                                                            |
| IRI1  | Member of a family of ice         | Cold-induced                                    | Protects against freezing damage                                        | Tremblay et al., 2005; Sandve et al., 2008; Zhang et al., 2010; Rudi et al., 2011 |
|       | recrystalization inhibitor proteins |                                                |                                                                          |                                                                            |

FIGURE 1 | Overview of the four different temperature treatments.

Clonal plants of the VRmin and VRmax F2 populations of Festuca pratensis, divergently selected for low and high vernalization requirement, respectively, were pregrown at 12 h photoperiod and 15/18°C and then exposed to four different temperature treatments: CA: cold acclimation at 2°C for 2 weeks, V-CA: vernalization and cold acclimation at 6°C for 7 weeks + 2°C for 2 weeks, DA: V-CA followed by deacclimation at 12°C for 1 week, RA: V-CA and DA followed by reacclimation for 2 weeks at 2°C, all at 8 h photoperiod. After each of the four temperature treatments plants were sampled for determination of flowering response, freezing tolerance, and gene expression.

PAR. Throughout the experiment plants were organized into trays with one plant of each genotype, and random trays of plants were used for testing of freezing tolerance, flowering response and tissue sampling after each of the four temperature treatments.

Determination of Flowering Response and Freezing Tolerance

For determination of the heading phenotype, three plants per genotype and temperature treatment were placed in the greenhouse at approximately 18°C and 16 h photoperiod. The flowering response was recorded as DTH (number of days from transfer to the greenhouse until the tip of the first panicle was visible), and as PHP (percent heading plants) and PHS (percent heading shoots per plant), both recorded when heading ceased (no new plants heading for a week).

For determination of freezing tolerance (LT50), four plants of each genotype and temperature treatment were divided into individual tillers. The shoots were cut at 5 cm and the roots at 2 cm and 4–9 random tillers from each genotype were placed in loose bundles in each of 13 boxes of moist sand in programmed freezing chambers initially set at 2°C. One control box was placed in a chamber with a constant temperature of 2°C, while the other 12 boxes were distributed among three chambers where the temperature was first lowered from 2°C to −3°C at 1°C h−1 and kept at this level for 12 h, after which the temperature was lowered again by 1°C h−1. Four test temperatures were used: −5, −10, −15, and −20°C for the CA treatment, and −13, −17, −21, and −25°C for the V-CA, DA, and RA treatments. When
the temperature reached one of the four test temperatures, one box from each of the three chambers was removed and placed at 2°C for thawing. After thawing tillers were planted in soil. Survival of individual plants, rated dead or alive, was determined after 3 weeks of growth in a greenhouse at approximately 18°C and 16 h photoperiod. Freezing tolerance (LT50; temperature required to kill 50% of the tillers) was calculated by probit analysis using PROC PROBIT in SAS 9.2 (SAS Institute, Inc., Cary, NC, USA).

**Gene Expression Analysis**

The shoot bases (1 cm, outer leaves peeled off) of all tillers from one plant per genotype and temperature treatment were excised, immediately frozen in liquid nitrogen and then kept at −80°C. Sampling was done during a 3 h period from 1 to 4 h after dawn, and the genotypes were sampled in a random order. Total RNA was extracted with the RNeasy Plant Mini kit (Qiagen). Ten microgram RNA of each sample was DNase-treated with TURBO DNase (Ambion, Life Technologies) and 1 µg of DNase-treated RNA was used as a template for cDNA-synthesis using the SuperScript VILO cDNA synthesis kit (Invitrogen, Life Technologies). A 10 µl control reaction without reverse transcriptase was included for all samples in order to confirm the absence of genomic DNA contamination. All cDNA samples were diluted 5x and 2 µl was used as template in real-time PCR reactions with SYBRGreen PCR Master Mix (Applied Biosystems, Life Technologies) in order to quantify transcript levels of VRN1, MADS3, CBF6, COR14B, CR7, LOS2, and IRI1 (see Supplementary Table S1 for primer sequences). The house-keeping gene ACTIN was used as a reference gene. PCR products were quantified in a 7500 Fast Real-Time PCR system (Applied Biosystems, Life Technologies). Relative quantity (RQ) of transcripts in each sample was determined by the ΔΔCt-method, where \( RQ = 2^{-ΔΔCt} \) and \( ΔCt = C_{gene\ of\ interest} - C_{ACTIN} \). Twenty-seven of the 756 sample-gene combinations, for which the qRT-PCR were not successful, were regarded as missing values in the analyses. In addition, four sample-gene combinations with more than 3x higher expression than the other samples in the same temperature treatment were considered as outliers and also regarded as missing values.

**Classification into Phenotypic and Genotypic Classes**

Based on the ability to head or not without vernalization (after CA only), the genotypes of VRmin were divided into two phenotypic classes: VRmin- (able to head without vernalization, seven genotypes) and VRmin+ (unable to head without vernalization, 14 genotypes). The VRN1-locus of all individuals was genotyped using the CAPS-marker described by Ergon et al. (2006), which recognizes one of the four haplotypes (b) of the VRN1-locus in this population. This haplotype is one of the two maternal haplotypes of a region of chromosome four containing QTLs controlling vernalization requirement (Ergon et al., 2006) and freezing tolerance (Alm et al., 2011) in the F1 mapping population. The b-haplotype is associated with low vernalization requirement and high freezing tolerance. Based on the b-allele, individuals in the VRmin population were divided into three genotypic classes: homozygous for the b-allele (four genotypes), heterozygous (nine genotypes), or no b-allele (eight genotypes). VRmax was not divided into genotypic classes due to the limited number of genotypes included from this population (bb1, −b;3, −2).

**Statistical Analysis**

To test for differences between F2 populations and phenotypic and genotypic classes of VRmin, the flowering response, freezing tolerance and gene expression data were subjected to analysis of variance using PROC GLM procedure in SAS 9.2. Pairwise differences between treatments were identified with the LSD-test of GLM. PROC CORR procedure was used to test for correlation among all traits of VRmin genotypes within temperature treatments.

### RESULTS

#### Flowering Response

The heading response differed significantly between the two F2 populations and between the four temperature treatments (Table 2). Some VRmin genotypes (defined as phenotypic class VRmin-) headed sparsely after only 2 weeks of CA followed by greenhouse conditions (Figure 2). VRmin- genotypes did not differ significantly from VRmin+ in the other three temperature treatments. After a total of 9 weeks at low temperatures (V-CA), VRmin genotypes had an average PHP of 94, PHS of 23 and DTH of 46 (Table 3). VRmax genotypes headed significantly less and later with PHP, PHS and DTH values of 61, 11, and 55, respectively. When V-CA plants were exposed to 1 week of de-acclimation (DA) the PHP and PHS did not change significantly, and both populations headed approximately 10 days earlier, indicating that the process of heading started during the

| Variable | Temperature treatment (T) | Population (P) | T × P |
|----------|--------------------------|----------------|-------|
| PHP      | 114.5***                 | 12.0***        | NS    |
| PHS      | 74.6***                  | 8.6***         | NS    |
| DTH      | 399.7***                 | 10.7***        | NS    |
| LT50     | 221.5***                 | NS             | NS    |
| CBF6     | 3.71*                    | NS             | NS    |
| COR14B   | 7.11***                  | NS             | NS    |
| CR7      | 19.0***                  | 13.8***        | NS    |
| LOS2     | 100.1***                 | NS             | NS    |
| IRI1     | 9.0***                   | NS             | NS    |
| VRN1     | 50.6***                  | NS             | NS    |
| MADS3    | 19.0***                  | NS             | NS    |

*df., degrees of freedom; PHP, percent of heading plants; PHS, percent of heading shoots per plant; DTH, days to heading; LT50, freezing temperature at which 50% of the plants are estimated to die according to results from a freezing test. Remaining variables are expression levels (RQ) of various genes in the shoot base tissue. F-values are given when the effect was significant. ***P < 0.001; **0.01 < P < 0.001; *0.01 < P < 0.05; NS, 0.05 < P.*
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FIGURE 2 | Heading response (measured as percent heading plants, percent heading shoots per plant and days to heading) and freezing tolerance of clonal plants of Festuca pratensis genotypes exposed to four different temperature treatments, CA, V-CA, DA, and RA (see Figure 1). Genotypes of two F2 populations selected for either high (VRmax, six genotypes) or low (VRmin) vernalisation requirement were tested; VRmin were further divided into two phenotypic classes (VRmin−, seven genotypes, and VRmin+, 14 genotypes), based on the heading response in CA. Plants that did not head during the experiment were given the value 149 for days to heading. LT50: freezing temperature at which 50% of the plants were estimated to die. Significant differences between groups within temperature treatments ($P < 0.05$) are indicated with different letters above bars.

DA treatment. When DA plants were exposed to 2 weeks of reacclimation (RA) all the plants of both populations headed (PHP = 100%). Both populations obtained higher PHS than V-CA and DA plants and headed earlier than V-CA plants. This response was stronger in VRmax genotypes and a significant difference between the populations remained for DTH only. After RA, the PHP and PHS values had changed by 6 and 64% relative to the values after the V-CA treatment in VRmin, while in VRmax, they changed by 64 and 209%, respectively. Thus, the vernalization requirement was not saturated after 9 weeks of low temperature in any of the populations, but VRmin was closer to saturation than VRmax. After V-CA, DA and RA, VRmin genotypes homozygous for the $b$-haplotype of the VRN1-region tended to head later and have lower PHS than other genotypes, although this was not significant for all treatments (Figure 3).

Freezing Tolerance

As expected, freezing tolerance differed between temperature treatments (Tables 2 and 3). After 2 weeks of CA LT50 was on average $-19.5^\circ C$, while after a total of 9 weeks at low temperature it was $-24.2^\circ C$. One week of de-acclimation lowered it to $-12.8^\circ C$, and 2 weeks of re-acclimation raised it to $-18.4^\circ C$, which was a significantly lower freezing tolerance than after the first 2 weeks of CA. There were no significant differences between populations within each temperature treatment. However, while VRmax was able to obtain the same level of freezing tolerance after RA as it had after CA, VRmin was not. This means that, on average, VRmin genotypes lost some ability to cold acclimate during vernalization and deacclimation, and this was caused by the VRmin- phenotypic class of VRmin (Figure 2). VRmin- was significantly more freezing tolerant than VRmin+ and VRmax in the CA treatment (LT50 $-21.1$, $-19.1$, and $-18.4$, respectively, $P = 0.05$). In the other treatments, there were no significant differences between populations or phenotypic classes. When grouping the VRmin genotypes into genotypic classes, those that were homozygous for the $b$-allele of the VRN1-region were more freezing tolerant than the other genotypes in the CA treatment (LT50 $-22.0$ and $-19.2$, respectively, $P = 0.05$, Figure 3).
Gene Expression

There was a significant effect of temperature treatment on expression of all genes (Table 2). The expression of CBF6, COR14B, and CR7 were not significantly different after the V-CA treatment as compared to only CA treatment, while the expression of VRN1 and MADS3 was more than 20x higher, and LOS2 and IRII more than 5x higher, in the V-CA treatment than in the CA treatment (Figure 4). All genes except CBF6 were down-regulated by deacclimation. In DA, LOS2, IRII, and MADS3 were down-regulated to only 0.06x or less of the RQ in V-CA, while VRN1, COR14B, and CR7 were down-regulated to around 0.25x of the RQ in V-CA (and CA in the case of COR14B and CR7). Only VRN1 and CR7 were significantly up-regulated by re-acclimation (4x higher RQ in RA than in DA). The expression of VRN1 became 16x higher than in CA plants but only 0.7x as high as in V-CA, whereas for CR7 the expression level was similar in CA, V-CA, and RA. The two F2-populations had similar gene expression patterns, except for CR7, which was significantly different, with VRmax having almost twice as high expression of CR7 as VRmin in the CA and RA treatments (Figure 5). There were no significant differences in gene expression between the two phenotypic classes of VRmin, but there were some differences between the genotypic classes based on the haplotype defined by the b-allele of VRN1. The b-haplotype was not significantly associated with expression of COR14B in the V-CA and DA treatments and of CBF6 and LOS2 in the DA treatment (Figure 6). There were also some weak, but significant, correlations between expression levels of different genes, and between expression of specific genes and PHS and LT50 in VRmin (Table 4). The strongest correlations (positive) were found between expression of VRN1, CBF6, COR14B, and LOS2 in the CA, DA, and RA temperature treatments, and there were also significant positive correlations between expression of these genes and IRII.

DISCUSSION

Flowering Response

The different vernalization response in the two F2-populations, and the segregation within VRmin for the ability to head without vernalization reported previously (Ergon et al., 2013), was confirmed in the experiment reported here. The ability to head without vernalization and the heading responses after 9 weeks of vernalization were not associated, and hence appear to be controlled by different genetic factors. As expected, the two F2 populations differed in the level of vernalization saturation after 9 weeks of cold treatment (V-CA). Although the b-haplotype was associated with the ability to flower without vernalization in the F1 generation (Ergon et al., 2006), it was not associated with this trait in the VRmin F2 population, which segregated for this trait, and after vernalization it was instead associated with later heading. This difference between generations could be due to epistatic effects being masked in the F1 generation or due to an effect of the other maternal haplotype, which could not be identified with the CAPS-marker.

Interestingly, plants from the DA treatment headed 10 days earlier than plants exposed to the V-CA treatment only (counted from the time of transfer to the greenhouse after the temperature treatment was ended). This shows that the reproductive development progressed at the same, or at a slightly faster, rate during DA (12°C, 8 h photoperiod), as compared to greenhouse conditions (~18°C, 16 h photoperiod), in spite of a lower temperature. Short photoperiods stimulate

| Trait | Population | Temperature treatment |
|-------|------------|-----------------------|
| PHP   | VRmin      | CA: 19 ± 7 (0–100)² | V-CA: 94 ± 3 (67-100)³ Ab | RA: 100a |
|       | VRmax      | 0c                    | 61 ± 20 (0-100)³ | 89 ± 11 (33-100)³ab | 100a |
| PHS   | VRmin      | 2 ± 1 (0–9)³c         | 23 ± 2 (6-37)³b | 22 ± 1 (11-31)³b | 37 ± 3 (16-55)³a |
|       | VRmax      | 0c                    | 11 ± 5 (0-30)³b  | 15 ± 4 (4-29)³b   | 34 ± 6 (18-54)³a |
| DTH   | VRmin      | 133 ± 6 (61–max²)    | 46 ± 1 (41–66)³b | 35 ± 1 (32-42)³c | 33 ± 1 (29-38)³c |
|       | Max²       | 55 ± 4 (41-66)³b     | 44 ± 3 (36-51)³c | 37 ± 2 (32-42)³c |
| LT50  | VRmin      | –19.8 ± 0.5 (~15.2 to –26.1)² | –24.1 ± 0.4 (~19.5 to –28.0)³ | –12.9 ± 0.04 (~12.6 to –13.2)³ | –18.5 ± 0.3 (~14.5 to –21.0)³ |
|       | VRmax      | –18.4 ± 0.7 (~15.4 to –19.4)² | –24.6 ± 0.8 (~21.8 to –26.6)³ | –12.8 ± 0.07 (~12.6 to –13.0)³ | –18.1 ± 0.4 (~17.1 to –19.7)³ |

Clonal plants were exposed to four different temperature treatments (see Figure 1) before being tested for the heading response and freezing tolerance. For the heading characteristics there were three replicate plants per genotype and temperature treatment. For the LT50 test tillers were exposed to four test temperatures (4–10 tillers per test temperature in each of three replicate chambers). The values given are the averages of the genotypes ±SE, the genotype ranges are given in brackets. Values not followed by the same letter are significantly different according to one-way analyses of variance within the temperature treatments (capital letters) or within populations (small letters; P < 0.05). ¹DTH was measured from the time of transfer to long days and growing temperatures in a greenhouse. ²For plants that did not head during the course of the experiment the DTH was set to 149 days (the maximum DTH observed + 1). ³In treatment CA, there were only two replicate LT50 tests for 10 and 2 genotypes in VRmin and VRmax, respectively, due to a limited number of tillers available for testing. In treatment DA, LT50 could only be estimated for 17 and 4 genotypes in VRmin and VRmax, respectively, because there was 100% mortality in the other genotypes. The LT50 was for these were set at –12.6°C (the highest observed LT50 + 0.1) and the reported LT50 may therefore be an overestimate.
induction of flowering and to some extent replace vernalization in perennial grasses, while long photoperiods accelerate flowering in vernalized plants and in plants without a vernalization requirement (Heide, 1988, 1994). The effect of photoperiod during the different stages of meristem and inflorescence development has not been described in detail in grasses; the effect may vary between species and genotypes. In the partially vernalized *F. pratensis* in our experiment, short photoperiods appeared to be more efficient in promoting floral development than long photoperiods as it was able to compensate for the difference in temperature.

**Effects of Temperature Treatments on Gene Expression**

The ability of some VRmin genotypes to head after only 2 weeks of cold (CA) was not significantly associated with higher VRN1 or MADS3 expression levels at the shoot basis in this temperature treatment. In previous experiments (Ergon et al., 2013), the ability to head without vernalization was associated with a higher expression of these two genes, particularly MADS3, in non-vernalized plants. In those experiments non-vernalized plants were grown under an 18 h photoperiod in the greenhouse prior to sampling and not 12 h followed by 8 h at 2°C as in the experiment reported here. The expression of VRN1 and MADS3 in non-vernalized plants of the VRmin-phenotypic class may depend on long days. There is little information available on expression patterns of MADS3 in other species. VRN1 is known to be induced or enhanced by short days in leaves and shoot apices of temperate grasses if certain conditions are met, i.e., after vernalization (Sasani et al., 2009, *Hordeum vulgare*), after a long period of short days (Dubcovsky et al., 2006, *Triticum aestivum*), and in genotypes that are not restricted by a requirement for vernalization and/or a short day period in order to flower (Gocal et al., 2001, *Lolium temulentum*). It is not clear whether VRN1 expression is actually required for transition to reproductive development in these cases. Instead, it may be upregulated in the apex after transition, where it appears to have a role in
FIGURE 4 | Gene expression of CBF6, LOS2, VRN1, COR14B, IRI1, MADS3, and CR7 in shoot bases of Festuca pratensis exposed to four different temperature treatments, CA, V-CA, DA, and RA (see Figure 1). Average values of all 27 genotypes are shown. Gene expression was measured by qRT-PCR as the relative quantity (RQ) of transcripts using ACTIN as the reference gene. Different letters indicate significant differences among temperature treatments (P < 0.05). In a few cases 1–4 genotypes are not represented due to failed or outlier qPCR reactions.

the development of the inflorescence and flowers (Gocal et al., 2001; Preston and Kellogg, 2007). This could also be the case for MADS3.

The genes that we studied differed in the way their expression was affected by the prolonged cold treatment. There were no significant differences between the CA and V-CA treatments in the expression of CBF6, COR14B, and CR7, but VRN1, MADS3, LOS2, and IRI1 were all present at higher transcript levels after the V-CA than after the CA treatment. The level of VRN1 expression increases during prolonged cold in winter cereals with a vernalization response (reviewed by Trevaskis, 2010), and this appears to also be the case in perennial grasses, such as L. perenne (Petersen et al., 2004) and Phleum pratense (Seppänen et al., 2010). MADS3 was more strongly expressed in L. perenne after 12 weeks at 5°C and 8 h photoperiod than after 6 weeks (Petersen et al., 2004). LOS2 tended to have a higher expression in F. pratensis after 18 and 21 days at 4/2°C (day/night) and 10 h photoperiod than after 1 day (Jurczyk et al., 2013a,b). IRI1 genes have been shown to be cold induced in leaves of L. perenne (Zhang et al., 2010), Brachypodium distachyon (Li et al., 2012; Colton-Gagnon et al., 2014) and Deschampsia antarctica (Chew et al., 2012). In these studies, expression was only tested for up to 1–2 weeks of cold exposure, except for Colton-Gagnon et al. (2014), who found a decline in expression from 1 to 2 weeks up to 5 weeks.

CBF6 did not change significantly in expression levels from 2 weeks (CA) to 9 weeks (V-CA) of cold, it was not down-regulated by deacclimation, and it was expressed at a higher level after reacclimation than before deacclimation. This is in contrast to many CBF-genes, which are induced rapidly by cold, and then revert to a basic level. CBF genes vary in their expression patterns, however. Among the wheat and barley CBF genes (Skinner et al., 2005), FpCBF6 has the highest identity with HvCBF6 and TaCBF6. HvCBF6 was found to have a delayed cold response with a peak in expression at 24 h after transfer to cold (2°C, 16 h photoperiod), and then maintained at that level (Skinner et al., 2005).

Across all genotypes COR14B and CR7 expression was also maintained during prolonged cold in our experiment, without any significant change in expression levels between plants...
exposed to 7 weeks of vernalization temperatures +2 weeks of cold (V-CA) compared with those exposed only to 2 weeks of cold (CA). Studies of cereals have reported both up- and downregulation of COR14B in response to prolonged cold. The outcome appears to be related to photoperiod. Dhillon et al. (2010) found that in *Triticum monococcum*, COR14B was downregulated during prolonged cold (lower expression level after six compared to 2 weeks), and more so under long days (16 h photoperiod) than under short days (8 h photoperiod). In line with this, in several studies of wheat a decline in the expression of COR14B was found during prolonged cold in long days (16 h photoperiod, Ganeshan et al., 2008; Laudencia-Chingcuano et al., 2011). Barley seedlings germinated in darkness, however, had higher expression of COR14B after exposure to 7 weeks of cold than after only 4 days of cold (Greenup et al., 2011). Similarly, Gana et al. (1997) found an increase in the expression of CR7 in crown tissue of wheat seedlings exposed to cold in darkness for up to 4 weeks. COR14B is known to be regulated by light-dependent factors (Crosatti et al., 1995, 1999), but this has not been described for CR7. The long day-induced gene FT1/VRN3 (Turner et al., 2005) may play a role in this as the locus harboring this gene is found to affect COR14B expression in barley (Cuesta-Marcos et al., 2015).

We found that all the studied genes except CBF6 were significantly downregulated in the shoot basis after 1 week of deacclimation at 12 h photoperiod. VRN1 was also downregulated in above-ground parts of *B. distachyon* when vernalized plants were placed at growth temperatures (16 h photoperiod, Colton-Gagnon et al., 2014). Similarly, Greenup et al. (2011) found a down-regulation of VRN1 in etiolated and vernalized barley seedlings when exposed to growth temperatures. The expression in deacclimated plants in both these studies were still higher than that in non-vernalyzed plants, and when etiolated and vernalized barley seedlings were placed in the greenhouse with a 16 h photoperiod, VRN1 expression in leaf blades remained high (Greenup et al., 2011). Sasani et al. (2009) had previously found that this expression in leaves was lower under short days (8 h photoperiod) than under long days (16 h photoperiod), while the expression in the shoot apices was not sensitive to photoperiod. Taken together, these and our results suggest that VRN1 is downregulated in shoot apices by deacclimation, but not to the level of non-vernalyzed plants, and that in leaf blades, but not apices/stem bases, the down-regulation is limited to short day conditions. Inclusion of other tissues than leaf blades, or species differences, may explain the down-regulation observed in shoots of *B. distachyon* by Colton-Gagnon et al. (2014).

It was only VRN1 and CR7 that were significantly upregulated by reacclimation relative to the deacclimated plants. Two weeks of reacclimation after deacclimation resulted in different expression levels than the two first weeks of CA for CBF6 (discussed above), VRN1 and COR14B. VRN1 expression was higher after RA than after CA, while the opposite occurred for COR14B. CR7, LOS2, and IRI1 were expressed at similar levels after CA and RA. Thus, it appears that VRN1 expression is somehow primed after prolonged cold; this may be related to changes in chromatin structure showed to occur at the VRN1 locus (Oliver et al., 2009, 2013). The lower expression of COR14B, but not CR7, after RA compared with CA, is interesting. Laudencia-Chingcuano et al. (2011) studied the effect of the VRN1 locus on the expression of COR14B and BLT14.1 in wheat during prolonged cold treatment under 16 h photoperiods (BLT14.1 is the wheat/barley CR7/BLT-gene with the highest similarity to FpCR7). Expression of both genes was affected by the VRN1 locus; their expression levels increased or was maintained for a longer time when the spring allele of VRN1 was absent and the expression of VRN1 was delayed. Expression of COR14B, however, was more strongly reduced after 10 weeks of cold than BLT14.1 (CR7). Our results are in agreement with this, and shows that vernalization may also have a down-regulating effect on COR14B, but not CR7, under short photoperiods.

### Freezing Tolerance as Affected by Vernalization and the VRN1 Chromosomal Region

The two phenotypic classes of VRmin differed in freezing tolerance in the CA treatment, the only treatment for which they also displayed different heading phenotypes. Here, the tendency to head was associated with better freezing tolerance. This effect may be attributable to genetic linkage. The genetic control of the ability to head without vernalization in the *F*1 generation of this plant material is mainly controlled by loci on chromosome 4F, of which some, but not the strongest ones, are closely linked to VRN1 (Ergon et al., 2006). There is also a QTL for freezing tolerance after 2 weeks of CA located close to VRN1 (Alm et al., 2011), which may account for the difference in freezing tolerance between the two phenotypic classes. Indeed, a re-examination of the marker data of Alm et al. (2003) showed that, in the *F*1...
FIGURE 6 | Gene expression of CBF6, LOS2, COR14B, and VRN1 in shoot bases of the Festuca pratensis VRmin population exposed to four different temperature treatments, CA, V-CA, DA, and RA (see Figure 1). Plants were grouped according to the genotype identified by the b-haplotype of the VRN1-region. Gene expression was measured by qRT-PCR as the RQ of transcripts using ACTIN as the reference gene. Averages of 8 (–), 9 (-b), and 4 (bb) genotypes are shown. Different letters indicate significant differences within temperature treatments ($P < 0.05$). There were no differences between genotypic classes in the expression of VRN1, MADS3, CR7, or LOS2 (only VRN1 shown).

population, the maternal marker haplotype of the VRN1-region associated with the ability to head without vernalization was also the one that was associated with better freezing tolerance after 2 weeks of CA (the paternal haplotypes did not have different effects on these traits in F$_1$). Šimkūnas et al. (2013) also found a positive correlation among Festulolium cultivars between heading prior to vernalization and survival in the field during the following winter. They speculated that this might be due to more young tillers in the plants heading prior to vernalization, but had no data supporting this. An alternative explanation could be a linkage between alleles conferring the ability to head without vernalization and freezing tolerance. The effect of the b-haplotype on freezing tolerance observed in the F$_1$ generation was retained in VRmin, and appeared to be recessive. No effect of the b-haplotype on freezing tolerance was seen after prolonged cold or after a deacclimation–reacclimation cycle, thus it appears to be involved in the relatively early stages of CA. The b-haplotype was also associated with later and less heading after the vernalization treatment, i.e., a lower responsiveness to vernalization, also in a recessive manner.

The ability to flower without vernalization on one hand, and vernalization response on the other hand, may be controlled by different genetic factors. Indeed, in our previous QTL analysis using the F$_1$ generation, the trait “vernalization requirement” included both these traits but was dominated by the variation in the ability to flower without vernalization, and the QTLs of largest magnitude were located 10 cM proximal to VRN1 (Ergon et al., 2006). In cereals, a QTL conferring freezing tolerance (Fr-1) is closely linked to VRN1 (Sutka and Snape, 1989), and VRN1 alleles conferring a vernalization requirement/slower vernalization response in cereals is associated with better freezing tolerance, similarly to the b-haplotype in our material. Based on phenotypic characterization (freezing tolerance, gene expression) of Triticum monococcum mutants where a small region encompassing VRN1 and a few more genes had been deleted, it was suggested that VRN1 is actually responsible for both traits (Dhillon et al., 2010). We found that freezing tolerance increased substantially by the vernalization treatment (7 weeks at 6$^\circ$C) prior to the 2 weeks of CA at 2$^\circ$C. In the V-CA treatment there was no difference in freezing tolerance.
between the two $F_2$ populations or between the phenotypic classes of $VR_{min}$. Reacclimation after deacclimation was efficient, but $VR_{min}$ had a significantly lower freezing tolerance after RA than after CA, while $VR_{max}$ did not, indicating some effect of the degree of vernalization on the de-acclimation/re-acclimation process, even under short photoperiods. The $VRN1$ $b$-haplotype was associated with better freezing tolerance in the CA treatment and less restoration of freezing tolerance in the RA treatment. The $b$-haplotype was also associated with later heading and a lower percent of heading shoots per plant in the vernalized treatments, and with a lower level of $CBF6$, $COR14B$, and $LOS2$ expression in vernalized and/or deacclimated treatments. This suggests that the $VRN1$-region somehow regulates the expression of these genes in vernalized plants or during deacclimation, and that less saturation of the vernalization requirement is associated with lower expression of cold-induced genes. In cereals, the opposite relationship has been found; vernalization saturation is associated with a down-regulation of cold-induced genes through mechanisms controlled by the $VRN1$-region (Fowler et al., 1996; Limin and Fowler, 2006; Laudencia-Chingcuanco et al., 2011). However, as shown by Dhillon et al. (2010), the effect that the $VRN1$ locus has on down-regulation of cold-induced genes, when the vernalization requirement is saturated, is dependent on long photoperiods. The short photoperiod used in our cold treatments, representing Norwegian mid-winter/early spring conditions, is a possible

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**TABLE 4** | Correlation coefficients (R) between variables measured for 21 genotypes in the *Festuca pratensis* $VR_{min}$ population given four different temperature treatments (see Figure 1).

| Variables/temperature treatments | PHS | DTH | LT50 | $VRN1$ | $COR14B$ | CR7 | $CBF6$ | $IRI1$ |
|----------------------------------|-----|-----|------|--------|---------|-----|--------|--------|
| DTH                             |     |     |      |        |         |     |        |        |
| CA                              | -0.89*** |     |      |        |         |     |        |        |
| V-CA                            | -0.75*** |     |      |        |         |     |        |        |
| DA                              | -0.49* |     |      |        |         |     |        |        |
| RA                              | NS   |     |      |        |         |     |        |        |
| LT50                            |     |     |      |        |         |     |        |        |
| CA                              | NS   | NS  |      |        |         |     |        |        |
| V-CA                            | NS   | NS  |      |        |         |     |        |        |
| DA                              | NS   | NS  |      |        |         |     |        |        |
| RA                              | NS   | NS  |      |        |         |     |        |        |
| $VRN1$                          |     |     |      |        |         |     |        |        |
| CA                              | NS   | NS  | NS   |        |         |     |        |        |
| V-CA                            | NS   | NS  | NS   |        |         |     |        |        |
| DA                              | NS   | NS  | NS   |        |         |     |        |        |
| RA                              | NS   | NS  | NS   |        |         |     |        |        |
| $COR14B$                        |     |     |      |        |         |     |        |        |
| CA                              | NS   | NS  | NS   | 0.63** |         |     |        |        |
| V-CA                            | NS   | NS  | NS   | NS     |         |     |        |        |
| DA                              | NS   | NS  | NS   | -0.47* | 0.55**  |     |        |        |
| RA                              | NS   | NS  | NS   | 0.59** |         |     |        |        |
| CR7                             |     |     |      |        |         |     |        |        |
| CA                              | NS   | NS  | NS   | NS     | NS      |     |        |        |
| V-CA                            | NS   | NS  | NS   | NS     | NS      |     |        |        |
| DA                              | NS   | NS  | NS   | NS     | NS      |     |        |        |
| RA                              | NS   | NS  | NS   | -0.46* | NS      |     |        |        |
| $CBF6$                          |     |     |      |        |         |     |        |        |
| CA                              | NS   | NS  | NS   | 0.58** | 0.75*** | NS  |        |        |
| V-CA                            | NS   | NS  | NS   | NS     | NS      | NS  |        |        |
| DA                              | NS   | NS  | NS   | NS     | NS      | NS  |        |        |
| RA                              | NS   | NS  | NS   | 0.66** | 0.61*   | NS  | NS      |        |
| $IRI1$                          |     |     |      |        |         |     |        |        |
| CA                              | NS   | NS  | NS   | NS     | 0.59**  | NS  | NS      |        |
| V-CA                            | NS   | NS  | NS   | 0.49*  | NS      | NS  | NS      |        |
| DA                              | NS   | NS  | NS   | NS     | NS      | NS  | NS      |        |
| RA                              | 0.56* | NS  | NS   | 0.55*  | NS      | NS  | NS      |        |
| $LOS2$                          |     |     |      |        |         |     |        |        |
| CA                              | NS   | NS  | NS   | 0.58** | 0.51*   | NS  | 0.48*   | NS     |
| V-CA                            | NS   | NS  | NS   | 0.43*  | NS      | NS  | 0.49*   | NS     |
| DA                              | NS   | NS  | NS   | 0.62** | NS      | 0.44* |        | NS     |
| RA                              | NS   | NS  | NS   | NS     | NS      | NS  | 0.55*   |        |

PHS, percent of heading shoots; DTH, days to heading; LT50, freezing temperature at which 50% of the plants are estimated to die. Remaining variables are expression levels of various genes in shoot basis tissue. ***$P < 0.001$; **$0.01 < P < 0.001$; *$0.01 < P < 0.05$; NS, $0.05 < P$. Note that a negative correlation with LT50 indicates a positive correlation with freezing tolerance. Expression of MADS3 was not significantly correlated with any of the other variables (not shown).
some correlation between VRN1 and down-regulation of cold-induced genes. We observed between vernalization saturation or vernalization requirement explanation for why we did not observe a similar association. 

Ergon et al. Vernalization and Freezing Tolerance mechanisms in early CA, possibly through the action of CBF and suggested that these genes may be regulated by similar patterns of VRN1 always positive. Oliver et al. (2013) observed similar expression SD, on gene expression, generative development, and freezing tolerance in Festuca pratensis. A pointed arrow indicates a positive, and a blunt arrow a negative, effect on, or association with, a process. Green and red lines indicate segregating effects observed here. Gray boxes indicate sets of co-regulated genes. LD, long days.

**CONCLUSION**

Some genotypes of *F. pratensis* are able to head to a limited extent without vernalization. This ability appears to be controlled by other genetic factors than the VRN1-region and is not associated with the responsiveness to vernalization or timing of heading. Timing of heading is associated with the VRN1-region but also with other genetic factors. Under short day conditions VRN1, CBF6, COR14B, LOS2, and IRI1 appear to be largely co-regulated, while CR7 and MAD3 are regulated by other mechanisms. Our results indicate that the relationship between vernalization and freezing tolerance in *F. pratensis* is complex. After 2 weeks of CA (Figure 7A), the genotypes that are able to head to a limited extent also has a better freezing tolerance, and the VRN1-region also has some effect on freezing tolerance. CR7 is more strongly expressed in genotypes with a higher vernalization requirement, but this does not result in better freezing tolerance at this stage. During prolonged cold (Figure 7B), VRN1, LOS2, IRI1, and MAD3 continue to be upregulated, while the expression of CBF6, COR14B, and CR7 are maintained at a constant level. The VRN1-region has some effect on the expression of COR14B. During deacclimation (Figure 7C), VRN1, LOS2, IRI1, MAD3, COR14B, and CR7 are down-regulated, and the expression of LOS2, CBF6, and COR14B is affected by the VRN1-region. At this stage expression of COR14B is associated with better freezing tolerance. During reacclimation (Figure 7D), VRN1 and CR7 are upregulated, while the expression of other cold-induced genes remains relatively stable. At this stage expression of CR7, which is associated with a lower vernalization requirement, is associated with better freezing tolerance.

Our results indicate that in *F. pratensis*, some genotypes can more easily lose some freezing tolerance after a deacclimation–reacclimation cycle, even under short photoperiods. The effect appears to be limited, but may increase significantly as photoperiods rapidly become longer than 12 h after the spring equinox. The role of the photoperiod in deacclimation and reacclimation processes in *F. pratensis* and other perennial grasses cultivated or growing naturally in the north, where photoperiods become long while there is still a risk of freezing, deserves further investigation.

**AUTHOR CONTRIBUTIONS**

ÅE, MH, and OR designed the study. ÅE conducted the experiment, analyzed the data, and wrote the manuscript. TM
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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fpls.2016.00207

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