Vernalization response of *Phleum pratense* and its relationships to stem lignification and floral transition

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- **Background** Timothy is a long-day grass species well adapted for cultivation in northern latitudes. It produces elongating tillers not only in spring growth but also later in summer. As the quantity and quality of harvested biomass is dictated by canopy architecture and the proportion of stem-forming flowering tillers, the regulation of flowering is of great interest in forage grass production.
- **Methods** Canopy architecture, stem morphology and freezing tolerance of vernalized timothy were investigated in greenhouse and field experiments. The molecular control of development was examined by analysing the relationship between apex development and expression of timothy homologues of the floral inducer VRN1 and repressor VRN2.
- **Key Results** True stem formation and lignification of the sclerenchyma ring occur in both vernalized and regrowing stems irrespective of the developmental stage of the apex. The stems had, however, divergent morphology. Vernalization enhanced flowering, and the expression of the VRN1 homologue was elevated when the apex had passed into the reproductive stage. High VRN1 homologue expression was not associated with reduction in freezing tolerance and the expression coincided with increased levels of the floral repressor VRN2 homologue. Field experiments supported the observed linkage between the upregulation of the VRN1 homologue and the transition to the reproductive stage in vernalized tillers. The upregulation of putative VRN1* or VRN2* genes was restricted to vernalized tillers in the spring yield and, thus, not detected in non-vernalized tillers of the second yield; so-called regrowth.
- **Conclusions** The formation of a lignified sclerenchyma ring that efficiently reduces the digestibility of the stem was not related to apex development but rather to a requirement for mechanical support. The observed good freezing tolerance of reproductive timothy tillers could be one important adaptation mechanism ensuring high yields in northern conditions. Both VRN1 and VRN2 homologues required a vernalization signal for expression so the development of yield-forming tillers in regrowth was regulated independently of the studied genes.

**Key words:** Apex development, canopy structure, elongation, flowering, freezing tolerance, lignification, *Phleum pratense*, regrowth, sclerenchyma ring, spring growth, timothy, vernalization, VRN1, VRN2.

**INTRODUCTION**

The canopy architecture of grasses is controlled genetically, although it is sensitive to environmental stimuli (Doust, 2007). In perennial grasses and winter cereals, vernalization has a great impact on the canopy architecture during spring growth, since the inhibition of flowering is released in vernalized shoots and the canopy consists of elongating and flowering tillers. Some species, such as *Lolium perenne* and *Festuca arundinacea*, require vernalization for the formation of true stems and flowering tillers (Heide, 1994). The digestibility of grasses, and especially that of stems, declines during maturation (Akin, 1989). Accumulation of lignin and the formation of a highly lignified sclerenchyma ring are thought to be responsible for the reduced microbial degradation and digestibility of stem cell walls (Grabber et al., 1992; Wilson and Hatfield, 1997; Chen et al., 2002). The transition of the apex from the vegetative to the generative stage is considered to be necessary for the formation and gradual accumulation of true stem biomass (Bélanger and McQueen, 1997), so there is a negative correlation between the quality and quantity of grass yield. The spring growth of vernalized grass tillers and the development of the yield and its quality can be predicted accurately by the accumulation of temperature sum in Nordic conditions (Rinne et al., 2001). Forage grasses are harvested several times during the year, and the growth and quality development of non-vernalized tillers of second and thirds yields – so-called regrowth – cannot be predicted accurately, leading to challenges in optimizing harvesting times (Kuoppala et al., 2008). The yield of regrowth tends to be quantitatively important but lower than that of spring growth, and consists of living and senesced leaves and tillers with vegetative structures (Pakarinen et al., 2008). The formation of flowering stems in regrowth is inhibited, most probably due to the lack of vernalization and to the presence of signals of shortening daylength.

Timothy (*Phleum pratense*) is well adapted for cultivation in Nordic conditions, since it has good overwintering capacity...
and the harvested forage grass is palatable to livestock. Compared with other cultivated forage grass species such as *L. perenne*, it has a superior freezing tolerance (Moriyama et al., 1995). The flowering does not require vernalization, but it is induced under long-day conditions (Heide, 1982). Knievel and Smith (1970) showed, however, that vernalization does affect the canopy structure in timothy, and the non-vernalized summer tillers produce more secondary tillers than the vernalized winter tillers. The release of flowering during vernalization and expression of floral inducers have also been connected with decreased freezing tolerance in winter cereals (Mahfoozi et al., 2001; Danyuk et al., 2003). Thus, the fulfillment of the vernalization requirement may increase the risk of winter deaths of tillers shifted to a reproductive stage. In grasses, such deaths of vernalized tillers may have an effect on the canopy structure and number of flowering tillers in spring growth. However, the compensation capacity of grasses to replace lost tillers is high, which may explain the lack of reports on the relationship between vernalization and freezing tolerance in forage grasses.

The amount of flowering and stem-forming tillers is important for the quantity and quality of forage biomass. The floral transition and formation of stem-forming tillers is induced by signals from both environmental and endogenous (or autonomous) pathways. According to current understanding, the floral inductive signal is perceived in vegetative tissues such as leaves, and then transported to the shoot apical meristem (Colasanti and Coneva, 2009). Flowering is known to be induced by several pathways such as vernalization, photoperiod, gibberellic acid (GA) and an autonomous pathway (He and Amasino, 2005). The autonomous and GA pathways respond to endogenous factors such as plant age, leaf number and energy status, whereas the photoperiod and vernalization pathways respond to environmental factors. In both dicots and monocots, vernalization promotes flowering by repressing the transcription of controlling gene(s) with simultaneous upregulation of flowering inducer gene(s) (Greenup et al., 2009). The molecular control of vernalization has been studied in *L. perenne* and *L. tenuifolium* among the forage grasses, and orthologues of the floral inducers VRN1 (VERNALIZATION 1) and VRN3 (VERNALIZATION 3) have been identified (Gocal et al., 2001; Andersen et al., 2006; Ciannamea et al., 2006; Studer et al., 2009). Moreover, floral repressors *LpMADS10* (*L. perenne* SHORT VEGETATIVE PHASE like gene) (Ciannamea et al., 2006) and *LpTFL1* (*L. perenne* TERMINAL FLOWER 1) (Jensen et al., 2001), and other genes, *LpCO* (*L. perenne* CONSTANS) (Martin et al., 2004) and *LpLIR1* (*L. perenne* LIGHT INDUCED RICE) (Ciannamea et al., 2007), related to the vernalization and daylength response of *L. perenne*, have been cloned from subtractive libraries. The identification of genes controlling flowering and manipulation of their expression can have great potential in attempts to increase the quantity as well as the quality of economically important forage grasses. Therefore, delayed or suppressed flowering has been one of the targets in breeding forage grasses for improved digestibility (Jensen et al., 2004).

The objective of this study was to reveal the physiological and molecular mechanisms of the vernalization response in timothy. Vernalization promotes both stem elongation and flowering in forage grasses and, therefore, the relationships of stem elongation and apex development to stem morphology and lignin localization was also studied. A homology-based identification of vernalization genes was used to design primers for VRN1 and VRN2 homologues in timothy, and the expression was examined under controlled vernalization and daylength conditions in greenhouse as well as in field experiments. While having an important role for yield formation in second and third harvests, the molecular control of development in regrowing tillers of grasses has not been previously studied. This study provides the first insights into how the apex development is related to stem lignification and how the two vernalization genes VRN1 and VRN2 are regulated in spring growth and regrowing tillers of timothy.

**MATERIALS AND METHODS**

**Plant material, growth conditions and determination of apex development**

For controlled vernalization conditions, timothy plants (*Phleum pratense* L. ’Iki’) were grown in a greenhouse at 20 °C under a 16 h day for 2 weeks prior to the transfer to vernalization conditions. The light intensity was 200–500 μmol m⁻² s⁻¹ and natural daylight was supplemented with 400 W high-pressure sodium lamps (Lucalox, LU 400/ HO/T/40 NG, Hungary). The plants were grown in 5 L pots containing fertilized and limed peat (Kekkilä B2, Finland).

The timothy plants (’Tammiisto II’) grown in field conditions were located at the MTT Agrifood Research, Maaninka Research Station, Finland (63°10’N, 27°18’E). The experimental field was established in 2005 in three replicates of experimental plots of 12 m² using barley as a cover crop, which was harvested after heading. During three consecutive years, the experimental field was fertilized for primary growth with 90, 13.5 and 22.5 kg ha⁻¹ and for regrowth with 90, 0 and 31.5 kg ha⁻¹ of N, P and K, respectively, and cut twice per growing season according to the typical cultivation practice of the area. The samples for anatomical and RNA analysis were harvested during the development of the sward in the second and third harvesting years, namely seasons 2007 and 2008.

The developmental stage of the shoot apex of tillers was recorded using a scale adapted from the 11-stage scale developed for *L. perenne* by Sweet et al. (1991). In this determination, the shoot apices at stages A1 (fully vegetative) and A2 (apex vegetative but elongated) were considered as vegetative, at stages A3 (apex elongated with visible leaf and spikelet primordia) and A4 (double ridge stage) as being at the transition from the vegetative to reproductive stage, and of stages A5 (development of the apex beyond the double ridge stage) and further as reproductive shoot apex. In addition to the original scale, three further stages of development were defined: at stage A12, lammas were more extended and had reached the height of the floret initials; at stage A13, glumes were well above the floret initials, which were more hidden by the lemmas; and at stage A14, some hairy structures could be seen on the external surface of the glumes. The apex height was measured from ground level to the base of the apex. The dissection of the apex was done if...
needed. The total length was measured from ground level to the top of the plants as the plant was stretched to its full length.

**Vernalization treatments and freezing tests**

Timothy plants grown in a greenhouse were subjected to vernalization treatment at 6 °C/4 °C (day/night), 8 h daylength in a growth chamber (Weiss Technik, Germany) for 0, 2, 10, 18 or 20 weeks. The pots were arranged in a completely randomized design with three replications in the growth chamber, and rotated once a week in order to minimize the effects of possible temperature or light difference within the chamber. The development of the shoot apex was followed during vernalization and the developmental stage was recorded. For RNA samples, the leaves and apices of ten plants were harvested immediately after the indicated vernalization time points and after 7–28 d of growth in the greenhouse.

The development of freezing tolerance during vernalization was evaluated by regrowth tests. After vernalization, plants were carefully removed from the soil, and the roots were rinsed with water. The plants had one to two tillers which was not dependent on the length of the vernalization period. Plants were placed in a controlled-temperature cooling bath (−1 °C) in covered culture tubes, four plants per tube, with a piece of wet tissue paper in the bottom. Extracellular ice was initiated by adding a small piece of ice to each tube, and the temperature was held at −1 °C for 1 h, then lowered at 2.5 °C h⁻¹ to the pre-determined minimum temperature (−15 to −25 °C). Samples were removed from the cooling bath at 2.5 °C intervals and thawed at 4 °C overnight. Meanwhile, the control samples were kept at 4 °C. After thawing, the plants were planted in soil in the greenhouse and the regrowth was evaluated weekly until the development of new tillers had ceased (5–6 weeks). Four plants per replication were subjected to each test temperature. The development of plants was monitored after vernalization treatments and freezing tests by calculating the number of leaves, stems and inflorescences weekly.

**Anatomical studies**

Tillers for anatomical studies were collected from the field experiment during spring growth and regrowth in 2007. At each collection time, 12–20 random tillers containing true stems were cut at the base. Tillers were kept in 4 °C, and total height, ligule and apex height, leaf and node number and the developmental stage of the shoot apex were all determined. Tillers were separated by the developmental stage of the apex into classes A1, A2, A3, A4 and A5 or further. Pieces of true stem were fixed immediately after collection and stored at room temperature in formalin--acetic acid--alcohol (FAA) before sample preparation. After measurement of ligule height, the internode located halfway up the true stem was used for anatomical studies. Stem samples were collected from the middle part of the selected internodes. Simultaneously samples from two or three of the youngest fully developed leaves were collected in RNA later solution (Applied Biosystems, USA) and kept at 4 °C until frozen at −70 °C before analysis. The stem samples were dehydrated in a graded ethanol series and embedded in paraffin at 60 ± 2 °C. Cross-sections of 8–12 μm were cut with a microtome (Leica RM2125, McBain Instruments). Paraffin was removed with Histo-Clear (National Diagnostic, UK), and samples were rehydrated in a graded ethanol series. Sections were stained with conventional safranin–alcan blue stain for lignin and polysaccharides (1 % safranin in 50 % ethanol and 0.1 % alcan blue in 3 % acetic acid), and finally mounted in Histomount (National Diagnostic, UK). Three replicate stems of each apex developmental stage were prepared and examined, and photographed with an Olympus DP 50 digital camera connected to a Leitz Laborlux light microscope or a Leica MZFL III stereomicroscope (Nilmok).
RESULTS

Interaction of the developmental stage of the apex with stem morphology

Under field conditions, a higher proportion of tillers are reproductive in spring growth than in summer or autumn regrowth. When both the main and side tillers were taken into account, the percentage of flowering tillers in primary growth was 30–40 % whereas the percentage in regrowth was only 1–2 % of all tillers. In this study, it became clear that the transition of the apex to the reproductive stage is not necessary for the initiation of stem elongation, and thus elongating tillers, in which the apical meristem is vegetative, can be found from both primary growth and regrowth (Table 1). In spring growth, the stem elongation occurs with increasing temperatures and lengthening photoperiods. Prior to this stage, all nodes are developed but they are still packed tightly together. As the internodes elongate, the nodes become visible. In field conditions in 2007, vegetative shoot apices were clearly above the ground level and true stems had been formed: the stem height of elongating tillers at the vegetative or transition stages (A1–A4) in spring growth ranged from 320 to 426 mm and in regrowth from 95 to 270 mm (Table 1). There were 2–4 visible nodes above the basal node in the elongating tillers of spring growth and 1–3 visible nodes in the regrowth (data not shown), and the lengths of the internodes were shorter in the regrowth than in the spring growth (Table 1). Elongating tillers with a reproductive apex (A12) in spring were considerably taller than those in regrowth. Although the height of the studied stems varied significantly between different apex classes, the height of the stem was not determined by the developmental stage of the apex.

During booting in spring, the internodes of vernalized tillers became hollow at an earlier developmental stage than internodes of regrowing tillers in July and August (Fig. 1). In spring growth, all analysed stem samples at all developmental stages of the apex already had hollow internodes. In contrast, most of the stem samples harvested in the autumn had solid internodes, although some hollow internodes were also observed. The total height of elongating tillers in spring and regrowth was similar from apex stage A2 onward, but the internodes in regrowing stems were shorter and this may explain the observed differences in stem morphology (Table 1, Fig. 1). A lignified schlenchyma ring formed at early stages of apex development, especially in the vernalized tillers during spring growth (Fig. 1). The formation seemed not to be associated with the developmental stage of the apex, but was rather related to stem height during spring growth (Fig. 1, Table 1) and sampling time during regrowth (Fig. 1, July vs. August). Although the anatomical analysis does not allow precise quantification of lignin content, the sclerenchyma ring in regrowing tillers was apparently less lignified in July than in August.

Vernalization, freezing tolerance and canopy architecture

The transition of apices to the reproductive stage was first observed in plants vernalized for 10 weeks. The percentage of flowering plants was also the highest (67 %) in plants vernalized for 10 weeks (Fig. 2) and it decreased gradually from 60 to 33 % in plants vernalized for 12–20 weeks. The decrease was not, however, statistically significant. After 12 weeks of vernalization, the number of developing tillers started to decrease, and after 20 weeks of vernalization, the canopy was thin, consisting of only a few lateral tillers, some with inflorescences. The leaf number increased until 10 weeks of vernalization and then was unaffected by prolonged vernalization time.

The plants vernalized for 10 weeks had the best freezing tolerance as they could recover from freezing down to –20 °C (Fig. 3A). Prolonged vernalization reduced the number of developing tillers gradually, so that at each tested freezing temperature there were fewer tillers than in plants vernalized for 10 weeks. The reduced number of tillers per plant was observed simultaneously with a higher percentage of flowering plants in plants vernalized for 18–20 weeks after freezing (Fig. 3B). The freezing test procedure seemed to recover the flowering induction in plants vernalized for 18–20 weeks.
since the percentage of flowering plants was 50–100 % after freezing tests in plants vernalized for 18–20 weeks (Fig. 3B) but only 33–53 % if the plants were directly transferred to the greenhouse (Fig. 2).

Expression of putative timothy VRN1 and VRN2 orthologues

PCR primers, designed to amplify putative VRN1 and VRN2 orthologues in timothy, produced fragment of 1103 and 192 bp, respectively. The first 75 bp of the putative genomic VRN1 sequence had a high homology (83 %) with the second exon of the VRN1 gene of T. monococcum (EU875079-2, 2012–2090), and with L. perenne (96 %) (AY198326, 361–468) and L. temulentum (96 %) MADS1 (AF035387, 338–445) VRN1 genes (Fig. 4A). The position of the T. monococcum intron at 2091–3225 and, thus, the potential start site of an intron at 66 bp also in the 1103 bp genomic sequence in timothy, led to primer design on a sequence with high homology with VRN1 genes in other species. The VRN2 candidate had 82 % homology to H. vulgare and T. monococcum VRN2 genes, also named ZCCT-Hb (DQ492696, 1853–2042) and ZCCT1 (AY485969-1, 542–731), respectively (Fig. 4B).

The expression of the timothy VRN1 candidate gene was upregulated in leaves and apex by vernalization (Fig. 5A, B). Significantly increased expression of VRN1 was first detected in the leaves of plants vernalized for 10 weeks and not in non-vernalyzed controls, and the level of expression remained high during prolonged vernalization. In the apex, VRN1 transcripts were also first detected after 10 weeks of vernalization but higher transcript accumulation was found as the vernalization time increased. VRN1 transcripts accumulated under short days during vernalization (10 + 0, 18 + 0, 20 + 0), and the transfer to long days in the greenhouse did not have a consistent effect on transcript levels. The VRN2 candidate transcript accumulated in long days after vernalization in leaves of plants vernalized for 2–10 weeks (2 + 1.5, 10 + 1, 10 + 2), but not in those vernalized for longer. During vernalization, altered expression of VRN2 was not observed in the apex. The VRN1 expression in the leaves of plants vernalized for 10 weeks coincided with the observed transition of apical meristem from the vegetative to the reproductive phase, as well as with the highest percentage of flowering plants.

Results from the field experiments supported the observations obtained from controlled vernalization conditions. The expression of the VRN1 candidate gene was elevated in elongating tillers during spring growth, and the upregulation coincided with the transition of the shoot apex from the vegetative (June A2–A3) to the reproductive stage (June A5) (Fig. 6). In the regrowing sward in July and August, elevated transcript levels of VRN1 were not detected in elongating tillers at the same apex developmental stage (July A3, August A3). The highest level of VRN2 expression was observed at the transition from the vegetative to the reproductive stage of the apex (June A3) and coincided with slightly reduced levels of VRN1 expression.

**DISCUSSION**

The quantity and quality of harvested forage biomass are defined by the canopy structure of the grass sward. The spring growth is mainly regulated by vernalization and photoperiod, leading to a high number of flowering tillers and gradually decreasing digestibility of the yield, whereas the regrowth is probably affected by photoperiod and plant endogenous signals. A significant reduction in the digestibility of developing forage yield is thought to be the result of extensive lignification of flowering stems (Akin, 1989; Grabber, 2005), and the transition of the apex from vegetative to reproductive is often thought to be critical for the digestibility of the forage biomass. Our observations with timothy under field conditions proved that the differentiation of the stem apex to the reproductive stage is not needed for the formation of true stem in this species, and a lignified sclerenchyma ring, which reduces the digestibility of the true stem (Wilson and Hatfield, 1997), can already be found in vegetative tillers. In F. arundinacea, a lignified sclerenchyma ring was observed in elongating tiller stems when the second node was palpable, and a significant increase in cell wall lignin content as well as
FIG. 1. The morphology of stems in spring growth (11–18 June) and regrowth (16 July and 6–13 August) at different developmental stages of the apex. A1 to A2 refer to the vegetative stage, A3 to A4 to the transition stage and A5 and more to reproductive stages of apexes. Stem samples of three individual tillers of the indicated apex stage were used for anatomical studies; \( n = 3 \) replicate stem sections per apex stage and sampling time. Scale bars = 200 μm in each photomicrograph.
accumulation of S-lignin monomers was observed later, when tillers were reproductive (Chen et al., 2002), although the developmental stage of the apex was not analysed in those studies. It seems that in timothy, the lignification process may be related to stem elongation and to the requirement for mechanical support of the stem rather than to the developmental stage of the apex. The sclerenchyma cell walls increase in thickness during maturation, leading to reduced lumen volume (Wilson and Hatfield, 1997). Such thick-walled sclerenchyma cells were observed only in regrowing stems in late summer (stage A9). This indicates that the development of the apex may have been arrested during regrowth, due to environmental and endogenous signals inhibiting flower induction, but the stem internodes had continued maturing processes without further development of the apex. Thus, the developmental stage of the apex does not necessarily correlate with the maturation stage of the stem. In timothy, arrested or backwards development of the apex was reported by Heide (1994), who found that flowering induction is not necessarily unidirectional but in some conditions a reproductive apex can revert back to a vegetative, leaf-forming apex.

The morphology of regrowing tillers differed notably from that of vernalized tillers; hollow stems were seldom observed in regrowing tillers but were already seen at the apex developmental stage A1 in vernalized tillers. The observed difference in morphology could not be explained by total height or by ligule height of the stems. Rather, the regrowing tillers also had shorter internodes and the height of the apex was lower, especially at early developmental stages of the apex. The maturation of the vernalized tillers seemed to be relatively synchronized, leading to high number of flowering tillers with hollow stems and most probably high cell wall content in the spring harvest. Taken together, the results show that the yield-forming stems in spring and regrowth have different morphology and the regulatory processes for stem formation and flowering are probably different. Our results support the observations by Knievel and Smith (1970) who also reported that non-vernalized and vernalized timothy tillers can have the same height. In their experiments vernalization did not increase the number of inflorescences but rather enhanced the rate of development. Those field experiments were, however, conducted during spring when the photoperiod is not repressing the transition to the reproductive stage as is probably the case in the regrowing tillers in this study.
Vernalization time seemed to have an optimum, after which longer times under vernalization conditions did not increase, but rather decreased, the number of flowering stems. **Langer** (1955) made similar observations on the reduction of the number of ears in timothy plants as the vernalization time of seeds was extended from 4 to 8–10 weeks. The induction of flowering in forage grass species that require double induction, e.g. both low temperature and long-day treatment, is shown to be sensitive to temperature and daylength conditions after low temperature treatment (Heide, 1994). Vernalization causes chromatin modifications that allow flowering induction to proceed (reviewed in Greenup et al., 2009). In this study, the formation of flowering stems after long vernalization was more intensive in plants which had been transferred to freezing test conditions for 1 d prior to planting in the greenhouse. The result indicates that the freezing test treatment after prolonged vernalization may have released some epigenetic regulation of flowering. Alternatively, the temperature and light conditions during freezing treatment were more optimal than greenhouse conditions for flowering induction in those plants.

In cereals, the winter or spring growth habit is defined by the vernalization requirement. Perennial grass species are harvested several times during the growing season, and the vernalization requirement for stem elongation and flowering most probably affect the yield and canopy structure of the first yield but not the second and third yields during summer. The molecular control of flowering in regrowing tillers of grasses has not been studied previously, although they have an important role for yield formation in the second and third yields. There is great variation in the vernalization requirement between species of forage grasses (Heide, 1994). Some species have an obligatory vernalization requirement (**L. perenne**, **Festuca** spp.) whereas in others vernalization enhances flowering as shown in this study on timothy. The spring growth of forage grasses is probably mainly regulated via vernalization and photoperiod signals, whereas the endogenous pathway may have a significant role in developmental processes during summer. However, in several temperate grasses (**Bromus inermis**, **F. pratensis**, **Dactylis glomerata** and **L. perenne**), it has been shown that flowering induction can be transferred from vernalized tillers to their offspring, probably by molecular messenger exchange through vascular tissue at the base of the tiller (Havstad et al., 2004). In *Arabidopsis* and rice, **FT** protein and its orthologue **Hd3a** have been shown to act as long-distance flowering signals (Jaeger and Wigge, 2007). Orthologues of **FT** have been identified in cereals and *Lolium* (**VRN3**) (Yan et al., 2006), and it is likely that the function is conserved across grasses. Thus, orthologues of **FT** and **VRN3** may have a role in transferring signals from vernalized timothy tillers to their offspring in regrowing swards.

### A VRN1

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**Fig. 4.** Sequence alignment of **VRN1** and **VRN2** candidate genes in timothy (*Phleum pratense, P. p.*) with corresponding orthologues in *T. monococcum* (*T.m.*, EU875079-2), *L. perenne* MADS1 (*L.p.*, AY198326) and *L. temulentum* MADS1 (*L.t.*, AF035387) for **VRN1**, and *T. monococcum* (*T.m.*, AY485969) and *H. vulgare* (*H.v* DQ492696) for **VRN2**. Primer sequences used for qPCR analysis are underlined.
Fig. 5. Quantitative real-time PCR expression analysis of putative VRN1 and VRN2 orthologues, as compared with Actin transcript levels, in (A) leaves and (B) apex of timothy plants during vernalization for 0, 2, 10, 18 and 20 weeks in a growth chamber and after 1–4 weeks of growth in a greenhouse. Data are the means ± s.e of three replicate samples.

Fig. 6. Relative amounts of putative VRN1 and VRN2 orthologue transcripts as compared with Actin transcript levels in the leaves of field-grown timothy plants at different developmental stages. Samples were harvested from vernalized plants of spring growth during June and regrowth during July–August. The leaves of individual plants were classified according to the developmental stage of the apex. Data are the mean ± s.e. of 3–4 plants. The transition from vegetative to reproductive stage occurs at apex stage A3–A4.
and VRN2 peaked in June when the daylength was the longest, 20 h. Also, the relative expression level was higher in field conditions compared with plants grown in the greenhouse. These observations indicate that the expression level of VRN1 and VRN2 was dependent on daylength and light intensity. The studied genes were also active only in vernalized tillers, which may indicate a low-temperature requirement of VRN1 induction in timothy similar to that which has been reported in barley (Treviskas et al., 2006; Sasani et al., 2009). An increased level of VRN1 expression and transition from the vegetative to the reproductive stage is associated with lowered freezing tolerance in winter cereals, and in spring-habit genotypes VRN1 was constitutively upregulated (Danylik et al., 2003). Elevated VRN1 expression did not seem to decrease the freezing tolerance of timothy, indicating that its reproductive tillers are not very sensitive to freezing. This may be one of the crop’s adaptation mechanisms ensuring high yields in northern conditions.

In cereals, vernalization releases VRN1 expression and induces flowering through repression of VRN2 and induction of VRN3 by long days (Hemming et al., 2008). VRN2 expression is repressed at low temperatures and it integrates the vernalization and photoperiod pathways in winter cereals. VRN2 is not strictly a vernalization gene but it is also thought to repress floral induction under long-day conditions during summer to avoid flowering just prior to winter (Dubcovsky et al., 2006; Treviskas et al., 2007). Although rice does not require vernalization for flowering, it has a gene closely related to VRN2, Ghd1, that represses flowering under non-inductive long days (Xue et al., 2008). In contrast to cereals, elevated levels of the VRN2 transcript were absent in regrowing timothy tillers. The hexaploid nature of timothy and the possibility for allelic variation in the VRN2 gene may have affected the results and, thus, definite conclusions on the role of VRN2 in regrowing timothy tillers cannot be drawn yet. A full understanding of how the development of elongating and flowering stems is regulated in vernalized and non-vernallized timothy tillers also requires detailed studies on the expression of other vernalization genes, such as VRN3, and regulatory genes of the photoperiod, GA and autonomic pathways.

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LITERATURE CITED


