Review

COR/LEA proteins as indicators of frost tolerance in Triticeae:
A comparison of controlled versus field conditions

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Abstract: Low temperatures in the autumn induce enhanced expression/relative accumulation of several cold-inducible transcripts/proteins with protective functions from Late-embryogenesis-abundant (LEA) superfamily including dehydrins. Several studies dealing with plants grown under controlled conditions revealed a correlation (significant quantitative relationship) between dehydrin transcript/protein relative accumulation and plant frost tolerance. However, to apply these results in breeding, field experiments are necessary. The aim of the review is to provide a summary of the studies dealing with the relationships between plant acquired frost tolerance and COR/LEA transcripts/proteins relative accumulation in cereals grown in controlled and in the field conditions. The impacts of cold acclimation and vernalization processes on the ability of winter-type Triticeae to accumulate COR/LEA proteins are discussed. The factors determining dehydrin relative accumulation under controlled cold acclimation treatments versus field trials during winter seasons are discussed. In conclusion, it can be stated that dehydrins could be used as suitable indicators of winter survival in field-grown winter cereals but only in plant prior to the fulfillment of vernalization requirement.

Keywords: cold acclimation; vernalization; frost tolerance; dehydrins; COR14b; growth chambers; field trials

1. Introduction

Low temperatures represent the dominant environmental factor determining plant growth and development during autumn/winter/spring seasons in temperate climates. Plants have evolved complex mechanisms aimed at an enhancement of their low-temperature tolerance in the process of cold acclimation [1]. In addition, plants growing in temperate climates where regular long periods of cold temperatures occur during the winter season have evolved developmental adaptation preventing a premature transition from more tolerant vegetative stage to cold-susceptible reproductive stage, i.e., complex mechanisms known as „vernalization” [2]. In Triticeae, two major loci determining the resulting frost tolerance were identified on the long arm of group 5 chromosomes including Vrn1/Fr1 locus encoding the major vernalization gene VRN1 induced by vernalization fulfilment [3, 4], and Fr2 locus encoding a cluster of cold-inducible CBF transcription regulators upstream of COR/LEA genes [5, 6]. In the field, winter seasons in temperate zones are characterised not only by low temperatures but by multiple environmental stresses including freeze-thaw cycles, temporal floods during snow melting or winter drought. All these environmental stresses induce
plant stress responses aimed at an enhancement of stress tolerance. Since low temperatures pose an enhanced risk of cellular dehydration and oxidative stress resulting from imbalances in photosynthesis, induction of enhanced low-temperature tolerance in plants is underpinned by enhanced accumulation of several stress-protective proteins including several COR/LEA proteins including several COR/LEA proteins and dehydrins.

Late embryogenesis abundant (LEA) proteins, originally identified in cotton seeds during embryo maturation and dessication, represent a superfamily of hydrophilic intracellular proteins which accumulate in plant cells under cellular dehydration [7]. Several LEA proteins belonging to either LEA-II (D-11; dehydrins) group characterized by the presence of at least one copy of 15-mer conserved lysine-rich sequence, the K-segment, or LEA-III (D-7/D-29) group characterized by the presence of at least one copy of conserved 11-mer motif, are induced by low temperatures thus are collectively termed as cold-regulated (COR) proteins [8]. Dehydrins are intracellular hydrophilic proteins which are classified as group II of LEA proteins and are defined by the presence of at least one copy of lysine-rich sequence known as K-segment near to C-terminus of their molecule [9, 10, 11]. In addition, they can also contain a tyrosin-rich sequence (Y-segment) near to N-terminus and a stretch of serine residues (S-segment) which could be phosphorylated and function as a nuclear localization signal [12]. Dehydrins are generally characterised by high content of glycine and hydrophilic amino acids and the absence of cysteine and tryptophan. They belong to intrinsically disordered proteins (IUPs) since they bind relatively large amounts of water molecules and acquire random coils in well-hydrated environments [13, 14, 15, 16]. Under reduced hydration, they form hydrogen bonds between side amino acid residues which results in the formation of defined secondary structures such as α-helices and β-sheets. For example, K-segments form amphipathic α-helices with acidic, basic and hydrophobic sides thus enabling interactions with both hydrophilic and hydrophobic surfaces such as membranes [17]. Under stress conditions, stress-responsive COR/LEA proteins accumulate to very high levels in plant cells to prevent cellular dehydration; it is estimated that they can reach up to 1-2% of total cellular proteins [18]. Due to versatility in their structure responding to cell hydration status, dehydrins can adopt multiple molecular functions corresponding to cell hydration status and interaction partners which include not only proteins but also membrane surfaces [15, 16]. Studies dealing with dehydrin molecular functions under cold reported their chaperone, cryoprotective, antifreeze, ion-binding and radical scavenging functions for some cold-responsive dehydrins [14, 19]; for more details on dehydrin molecular structure and functions, see Kosová et al. (2019) [20]. In winter wheat and barley plants, the major cold-inducible dehydrins belong to Kn structural group harboring multiple K-segments in their molecules (K6 in wheat WCS120 while K9 in barley DHN5) and revealing significant cryoprotective activities [18, 19]. Cold-regulated LEA-III proteins include several chloroplast-targeted proteins regulated by both light and low temperatures such as WCS19, WCOR14a,b, WCOR15, etc. [21, 22].

Several studies in controlled conditions (growth chambers) revealed a significant positive relationship (a positive correlation) between COR/LEA (dehydrin) transcript/protein relative accumulation and acquired frost tolerance for some cold-inducible proteins [23, 24, 25] not only under cold but also under a broad range of growing temperatures from cold to optimum ones [26, 27] due to higher threshold induction temperatures of acquired frost tolerance determined as lethal temperature for 50% samples (LT50) for highly tolerant genotypes of winter cereals when compared to the less tolerant ones [28]. Some cold-inducible dehydrins, e.g., WCS120 in wheat or DHN5 in barley, can be thus considered as markers of plant acquired frost tolerance for young plants in vegetative stage grown under controlled conditions. However, to be utilized in breeding, field trials are necessary. The results of field experiments indicated that dehydrins could be used as indicators of plant winter survival but only under certain conditions [29]. The aim of the present review is to provide a summary of the recent studies dealing with cold-inducible COR/LEA proteins and their relationships with plant acquired frost tolerance both in controlled and the field conditions. Environmental and developmental factors affecting COR/LEA protein relative accumulation are discussed.
2. Cold acclimation studies in controlled conditions

Plants as poikilothermic organisms have to adjust their metabolism to ambient temperatures. Adjustment of chilling-tolerant and freezing-tolerant plants such as Triticaceae cereals or winter oilseed rape to low above zero temperatures in +15 to 0°C range resulting in an acquisition of enhanced frost tolerance (FT) is known as cold acclimation (CA) [1, 30]. Acquired FT is usually expressed as lethal temperature of 50% of the sample (LT50) which can be determined either by a direct frost test including regulated freezing and thawing of the samples in laboratory freezers [31] or by determination of tissue damage via electrolyte leakage [32]. The key inducing factors in plant cold acclimation process represent both low temperature (chilling) and photoperiod (short days) which determine chloroplast redox status and the activity of photosynthetic apparatus which provides energy for biosynthesis of novel molecules of transcripts, proteins and metabolites necessary for plant adaptation to altered environment [33]. CA appears to represent a modular response associated with alterations in nucleosomal histone H2A isoforms indicating chromatin remodelling resulting in transcriptome and proteome reprogramming [34, 35]. In cold acclimation studies performed in controlled conditions such as growth chambers, plants are grown under defined photoperiod and temperature regimes which usually represent a continuous cold treatment resulting in an acquisition of maximum frost tolerance. Controlled conditions of growth chambers thus represent an ideal model for the study of the effects of cold on cold-inducible CBF-COR/LEA genes (pathways) expressions. Under cold, dehydrin expression is regulated by both ABA-dependent and ABA-independent pathways such as CBFs as indicated by the presence of multiple copies of ABRE and CRT/DRE motifs in their promoters [36, 37]. Moreover, both ABA-dependent and ABA-independent pathways reveal crosstalks in their regulation since ABA-induced kinase SnRK2 phosphorylates ICE1 which in turn activates CBF pathway [38, 39]. Continuous cold treatments lead to significantly enhanced expression of CBF/COR pathway indicated by COR/LEA transcript peaks at 3-4 days of cold treatment while a slower rise of corresponding proteins resulting in a maximum at 2-3 weeks of cold treatment [40, 41]. Moreover, organ-related differences in dehydrin protein relative accumulation were found by a comparison of leaf and crown tissues with crowns representing the crucial organ for a whole plant survival [41]. Thus, determination of COR/LEA proteins in crown tissues in winter cereals seems to represent a useful approach for assessment of plant winter survival. Due to the high level of cold-induced dehydrin protein accumulation, their direct relationship to plant acquired frost tolerance was repeatedly found [23]. A correlation between dehydrin protein relative accumulation and plant acquired frost tolerance determined as LT50 was found not only for winter-type cereals such as winter wheats [24] and barleys [25] but also for other winter-type crops such as winter oilseed rape [42] when grown in controlled environment.

In addition to temperature, CA process including the expression of some CBF-COR/LEA pathway genes and the resulting FT seems to be affected by light intensity and light quality, i.e., the red to far-red (R:FR) light ratio [21]. Ahres et al. [43] studied barley response to low temperatures (5 and 15°C) and different light conditions including three light intensities and FR light supplementation on acquired FT and CBF14, COR14b and DHN5 transcript levels and found a significant response to both cold and light intensity and FR supplementation for CBF14 and chloroplast-located COR14b genes while DHN5 responded only to cold.

The differences between cold-tolerant (winter) and cold-susceptible (spring) genotypes in cereals (wheat, barley) related to CBF-COR/LEA pathway induction:

1/ Genetic differences: cold-tolerant winter genotypes encode higher gene copy number of cold-inducible CBF genes at Fr2 locus when compared to cold-susceptible spring ones.

The differential induction level of CBF/COR pathways between frost-tolerant and frost-susceptible genotypes can be determined genetically; for example, a comparative study by Tondelli et al. [44] revealed that frost-tolerant winter
barley Nure has a higher CBF gene copy number in cold-inducible Fr2 locus than frost-susceptible spring barley Tremois. The differences in gene copy number of CBFs and other cold-inducible genes can thus underlie the differences in COR/LEA/dehydrin protein accumulation and the resulting frost tolerance between contrasting genotypes (e.g., spring vs winter ones). Similarly, enhanced levels of WCBF2 TF and downstream Cor/Lea transcripts Wdhn13, Wcor14 and Wcor15 were found in frost-tolerant winter wheat Mironovskaya 808 in comparison to frost-susceptible spring wheat Chinese Spring; moreover, the transcript levels of WCBF2 as well as Wdhn13, Wcor14 and Wcor15 peaked later (at 42 days of cold treatment) in Mironovskaya 808 than in Chinese Spring (at 21 days of cold treatment) [45].

2/ Threshold induction temperatures: Fowler [28] had shown that highly frost-tolerant winter cereals such as rye start inducing enhanced acquired frost tolerance determined as LT50 (lethal temperature for 50% of the samples) at higher growth temperatures in comparison with the less tolerant ones. Analogous patterns to LT50 were found also for cold-inducible CBFs and COR/LEA proteins, i.e., cold-tolerant winter cultivars start inducing cold-inducible genes such as CBFs and downstream COR/LEA proteins at higher temperatures in comparison with cold-susceptible ones. Vágújfalvi et al. [46] detected Cor14b transcripts in frost-tolerant winter line G3116 of einkorn wheat (T. monococcum) at higher temperature (up to 20 °C) than in frost-susceptible spring line DV92. Campoli et al. [47] detected different threshold induction temperatures for different CBF structural groups based on their phylogenetic analysis in winter barley, winter wheat, two winter rye and one spring rye cultivar. Similarly, Badawi et al. [48] distinguished ten CBF phylogenetic groups in two Triticum species out of which five Pooidae-specific groups revealed higher constitutive and low temperature inducible expression in winter wheat Norstar. Our previous studies [26, 27] demonstrated that cold-inducible proteins such as wheat WCS120 or barley DHN5 can be detected in the highly frost-tolerant cultivars such as Mironovskaya 808 or Odesskij 31 at higher temperatures (17-20 °C) than in the less tolerant winter wheats or barleys (around 10 °C).

3/ Differential phytohormonal regulation of CA process: CA leads to repression of plant growth and development. In A. thaliana, Achard et al. [49] observed a positive effect of CBF1 on accumulation of DELLA proteins known as growth repressors due to stimulation of GA-2 oxidase resulting in reduction of active gibberellins (GA). Our comparative studies on winter wheat Samanta and spring wheat Sandra, winter wheat Cheyenne - Chinese Spring 5A substitution lines as well as einkorn wheats G3116 (facultative) and DV92 (spring) revealed similar patterns of FT (LT50) and dehydrins (COR14b, WCS120 family) induction during the first days of CA treatment; however, at later stages (21 to 42 days CA), spring genotypes revealed significantly lower FT and WCS120 proteins levels. Phytohormone analyses also revealed a two-phase CA response with an alarm and early acclimation phase (1-3 days CA) with increased ABA in all growth habits inducing stress acclimation response while differential responses in later CA phases (7-42 days) when winter types maintained high levels of stress acclimation related phytohormones (ABA, JA, SA) while spring types revealed induction of phytohormones involved in vegetative-to-reproductive phase transition such as auxin, bioactive CKs and GAs probably due to VRN1 gene expression and floral meristem development [50, 51, 52]. Differential phytohormone dynamics may thus be the reason of lower FT and COR/LEA transcript/protein levels found in spring genotypes compared to winter ones at full CA (2-3 weeks CA treatment).

3. Effect of vernalization on cold-inducible pathways including dehydrins

Vernalization, i.e., requirement of a sufficiently long period of low temperatures in order to gain potential to flowering represents an evolutionary adaptation to temperate climates with harsh winter seasons in winter annuals. At molecular level, fulfillment of vernalization requirement puts an imprint to plant stress memory due to epigenetic modifications of specific vernalization genes resulting in irreversible downregulation of flowering repressors such as
FLC in *A. thaliana* or an activation of flowering inducers such as VRN1 in Triticeae [53]. In Triticeae, fulfillment of vernalization requirement in cereals winter growth habits leads to a downregulation of the major flowering repressor VRN2 leading to an induction of VRN1 which is a MADS-box gene revealing both morphogenetic and transcription regulation functions and acting as a flowering integrator indicating plant preparedness to flowering [54, 55, 56]. In Triticeae, up-regulation of VRN1 gene expression is also accompanied by changes in histone methylation status: the level of H3K27me3 (histone 3 lysine 27 trimethylation) decreases and the level of H3K4me3 increases resulting in an active chromatin state for transcription [57]. Vernalization studies in winter-type cereals repeatedly confirmed a dominant negative effect of VRN-1 gene expression on cold-inducible CBF/COR pathway expression [4, 58] while study of vernalization in model plant *Arabidopsis thaliana* revealed a crucial role of vernalization-induced *SOC1* gene as a direct inhibitor of CBF/COR pathway [59].

Studies on VRN1/Fr1 substitution lines between winter wheat Norstar and spring wheat Manitou revealed that an induction of VRN1 gene expression following vernalization fulfillment correlates with a downregulation of CBFs and COR/LEA gene expression [4]; however, the studies on Norstar-Manitou substitution lines indicate that the type of Vrn-1A locus does not have a significant effect on the initial rates of CBF-COR/LEA pathway gene expression which is rather governed by the genetic background [60]. A negative correlation between the expression of VRN1 gene expression and CBF-COR/LEA cold-inducible pathway. However, the molecular mechanisms underlying VRN1 and CBF-COR/LEA interactions still remain to be elucidated since no direct link between VRN1 and CBFs was identified [61]. Some VRN1 downstream regulated genes were identified in barley by Deng et al. [62] including a flowering promoter VRN3 gene also known as FLOWERING LOCUS T-1 (*FT1*) gene whose expression is activated by VRN1, flowering repressors VRN2 and ODDSOC2 genes whose expressions are downregulated by VRN1, and some genes involved in abscisic acid, jasmonic acid, and gibberellin biosynthesis pathways. In addition, VRN1 also seems to positively regulate some morphogenetic genes involved in floral meristem development such as *AP1* [63].

Fulfillment of vernalization requirement in winter cereals is indicated by induction of VRN1 gene expression. In *Arabidopsis thaliana*, the link between vernalization fulfillment and a downregulation of CBF-COR/LEA pathway has already been elucidated. Vernalization-induced downregulation of a major flowering repressor FLC results in an upregulation of *SOC1*, a MADS-box flowering integrator which was reported to inhibit CBF gene expression [59]. The study by Li et al. [63] on Vrn1/Fr1 substitution lines between winter wheat Norstar and spring wheat Manitou („winter Manitou“ and „spring Norstar“) revealed that Vrn1/Fr1 locus is a dominant but not a sole locus determining acquired frost tolerance expressed as LT50 and COR/LEA gene levels since „winter Manitou“ has ca 5 °C higher LT50 when compared to „Norstar“ (and even „spring Norstar“ under short days) indicating a joint effect of both Vrn1/Fr1 locus and the genetic background on the resulting LT50 and COR/LEA gene expression. Under short-day conditions which delay vegetative-to-reproductive transition, spring Norstar revealed nearly the same acquired frost tolerance as Norstar due to the same genetic background including predominantly *Fr2* loci harbouring CBF cluster on the long arms of chromosome 5 group and loci on the long arms of chromosome 6 group encoding WCS120 genes from dehydrin group [64]. Li et al. [63] detected 21 cold-responsive differentially expressed transcripts including *CBF3*, *WCO-R410b*, *CS120*, three *CS66* genes highly expressed in the lines with Norstar background while 14 cold-responsive transcripts including *LOS2* and two *CS66* genes highly expressed in lines with Manitou background. The effect of vernalization fulfillment on the ability of winter wheat Mironovskaya 808 to reacclimate to cold following a period of warmer temperatures inducing deacclimation was studied by Vitámvás and Prášil [65] in controlled conditions. They found out that fulfillment of vernalization significantly reduced winter wheat ability to accumulate WCS120 proteins after reacclimation.
In a summary, cold acclimation leads to cessation of plant growth and development aimed to induce enhanced stress tolerance while vernalization results in an acquisition of the competence to flowering. A comparison of the effects of cold acclimation and vernalization on winter cereals is provided in Table 1.

Table 1. A comparison of cold acclimation and vernalization processes and their impacts on winter Triticeae.

| Characteristics | Cold acclimation                                      | Vernalization                                           |
|-----------------|-------------------------------------------------------|--------------------------------------------------------|
| Inducing conditions | Short-term cold (days to weeks), Short-day photoperiods | Long-term cold (weeks to months), Long-day photoperiods (VRN3/FT1 pathway) |
| Plant response  | Conservation of vegetative stage; shoot apex: single-ridge (new leaves) | Transition to reproductive stage; shoot apex: double-ridge (floral meristem) reduced ability to induce FT under LT | Phytohormones: auxin, active cytokinins and gibberellins |
| Phytohormones: ABA, JA, SA (stress tolerance induction), DELLA (growth repressors) | Phytohormones: auxin, active cytokinins and gibberellins | |
| Gene expression | Upregulation of genes associated with enhanced FT (CBF-COR/LEA) | Downregulation of genes associated with FT acquisition (CBF-COR/LEA) |
| High levels of flowering repressors (VRN2 in winter cereals) are associated with high FT | Upregulation of VRN1 and floral meristem identity genes (API, AGL19, AGL24) | |

4. Field studies: COR/LEA proteins and winter hardiness

Winter seasons in temperate climates can be characterized by a complex of environmental stresses including not only low temperatures (both chilling and freezing temperatures) but also other stress factors such as water regime related stresses including both winter drought as well as temporal flooding events following snow melting, mechanical stress caused by soil heaving, and others. Studies aimed at the effects of multiple stress treatments on plants indicated that the resulting impacts of multiple stresses do not simply equal to the sum of the individual stress treatments thus highlighting the urgent need to study multiple stress impacts in controlled experiments as well as in the field [66]. All these environmental factor determine the final winter hardiness of winter crops. Winter hardiness thus represents a complex phenotypic trait which is usually determined as percentage of plant winter survival. Since different dehydrin proteins reveal a differential response to various environmental stresses it is thus evident that a sum of various dehydrin proteins detected in the plant samples reveals a better correlation with plant winter survival than a single dehydrin protein [29].
Basically, two major types of temperate climate can be distinguished: a continental climate which is characterized by continuous freezing temperatures throughout the whole winter, and a maritime /transitional type of climate which is characterized by fluctuations between freezing and above-zero temperatures that is associated with freezing/thawing cycles of water as well as mechanical stress (damage) caused by ice crystals, soil heaving, etc. Periods of low temperatures are thus interspersed by periods of relatively warm temperatures inducing deacclimation in overwintering plants. It is known that vernalization fulfillment adversely affects plant ability to reacclimate to low temperatures and newly induce several cold-responsive genes including CBF-COR pathway. Therefore, the level of relative accumulation of several COR/LEA proteins including dehydrins (LEA-II) but also chloroplast-located COR14b (LEA-III) proteins reveals a correlation with plant winter survival only in plants prior to vernalization fulfillment, i.e., sampled in early winter (till Christmas) [29]. Samplings carried out in later stages (January, February) revealed no significant correlation between plant winter survival and accumulation of COR/LEA proteins [29, 67] due to periods of relatively warm temperatures inducing partial deacclimation.

Continental climates are characterized by continuous freezing throughout the whole winter season thus enhanced levels of COR/LEA/dehydrin proteins were found during the whole winter season up to March in the crown tissues of winter wheat, triticale, and rye [68, 69]. Pomortsev et al. [68] found high quantitative levels but differences in qualitative profiles in dehydrins in winter wheat cv. Irkutskaya at different sampling dates (November, February, March) during winter season in Central Siberia, Irkutsk, Russia. The authors detected the presence of 29 kDa dehydrin band only in autumn samplings (November) while no such band was detected in winter and spring samplings (February, March). Ganeshan et al [69] presented a 3-year study on cold acclimation and COR/LEA gene expression in Norstar-Manitou set of substitution lines. The studied COR/LEA (WCS120, WCOR410, WCOR14) transcript levels decreased throughout winter except for WCOR14 transcript in winter Manitou; however, significant alterations among the different seasons were observed which corresponded with fluctuations in soil temperature at crown depth. In contrast, in maritime and transitional zones such as Italy or Czechia where freezing periods are interspersed by relatively long periods of above-zero temperatures during winter (freeze-thaw cycles) when partial cold deacclimation occurs, relatively low levels of COR/LEA/dehydrin proteins were found in plant tissues in later phases of winter (January, February) and these levels did not correspond with the plant freezing tolerance [29, 67] since plants are already vernalized at this time and cannot efficiently re-acclimate after a period of deacclimation.

In Italian studies, a positive relationship between COR14b transcript accumulation and plant winter survival was found in samplings from November and December [70] while no significant relationship was found in samplings from January and February [67]. Rizza et al. [71] found no significant relationship between COR14b transcript accumulation and acquired frost tolerance in a set of European barley cultivars; in contrast, they proposed Fv/Fm chlorophyll fluorescence parameter as a reliable indicator of acquired FT in barley.

Vítámvás et al. (2019) [29] found a positive relationship between cold-inducible dehydrin (sum of WCS120 proteins in winter wheats; DHN5 protein in barleys) accumulation in crown tissues and winter survival in field grown winter wheat and barley plants at two locations (Prague-Ruzyně, Lužany) and two winter seasons in Czechia in December samplings prior to the fullfilment of vernalization while the positive relationship between plant winter survival and dehydrin proteins was lost in later samplings (January, February) when plants were already vernalized and exposed to partial deacclimation during warmer periods throughout the winter season.

5. Conclusion

Cold acclimation process leading to enhanced low temperature tolerance is associated with profound alterations in gene expression including activation of CBF-COR/LEA pathway. There are significant differences between
frost-tolerant and frost-susceptible genotypes at both genotypic and phenotypic levels with frost-tolerant genotypes encoding higher copy numbers of cold-inducible CBF genes as well as revealing higher threshold induction temperatures for cold-inducible genes. Vernalization fulfillment associated with an induction of VRN1 gene leads to a downregulation of CBF-COR/LEA pathway thus decreasing the possibility of cold reacclimation following a period of warmer temperatures. For the study of cold acclimation process in controlled conditions (growth chambers), defined environmental conditions such as temperature, photoperiod, irradiance, substrate (soil), fertilization and plant protection (no pathogens) are used in most experiments. These conditions lead to exact and repeatable results which are surely valuable for plant science research; however, they are very different from those the plants have to face in the field (Table 2).

Table 2. A comparison of the impacts of controlled and field conditions on dehydrin relative accumulation and plant tolerance to environment.

| Characteristics       | Controlled (growth chamber)                                                                 | Field experiments                                                                 |
|-----------------------|---------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------|
| Growth conditions     | Controlled (growth chamber): a very few variables (usually temperature, or photoperiod) - distinct and contrasting values (e.g., optimum, e.g., +20 °C vs cold, e.g., +4 °C; long-day 16 h/8 h vs short-day 8 h/16 h day/night); constant irradiance; defined watering | Very variable, continuously changing conditions with significant fluctuations (temperature) or continuously changing values (photoperiod - day shortening in autumn, day prolongation in spring); several additional stress factors including water-related stress (transient drought or wet - waterlogging and flooding), nutrient-related stress, mechanical wounding, biotic stress (pathogens) |
| Plant growth stage    | Defined: usually early vegetative stage (e.g., 3-leaf stage) or (less often) after vernalization fulfillment | Continuous development from vegetative to reproductive transition (vernalization fulfillment) |
| Plant stress tolerance| Frost tolerance expressed as lethal temperature for 50% samples (LT50) determined by laboratory methods (direct frost test) under defined freezing, thawing and recovery conditions | Winter hardiness expressed as percentage of survived plants (winter survival) as a result of joint effects of all stress factors during winter |
| COR/LEA proteins      | A correlation between relative abundance of a single cold-induced COR/LEA protein (a single band on the immunoblots) and LT50 both before and after vernalization fulfillment under continuous cold | A correlation between relative abundance of the sum of cold-inducible COR/LEA proteins (all COR/LEA bands on the immunoblots) and plant winter survival only at early sampling dates prior to vernalization fulfillment |

Winter seasons in temperate climates are characterised not only by low temperatures but a complex of environmental stresses related to plant water regime, mechanical wounding, and pathogens. Plant winter hardiness thus represents a
complex phenotypic trait affected by various environmental factors and determined by expression of wide arrays of genes. Therefore, rather a sum of all dehydrin proteins than a single protein identified in the plant materials reveals a correlation with winter survival in winter cereals. In temperate climate zone, winter seasons differ between maritime/transitional climatic regions such as western/central Europe and continental climates regions such as central Canada and Siberia. Continental climates are characterised by continuous freezing temperatures thus overwintering cereals reveal enhanced levels of COR/LEA proteins throughout the whole winter season. In contrast, in maritime/transitional climates characterised by freeze/thaw cycles, high levels of COR/LEA proteins correlating with plant winter survival can be found only prior to vernalization fulfillment whereas no significant correlation between COR/LEA protein accumulation and winter survival was found in later phases of winter (January, February) when vernalized plants were exposed to periods of relatively warm temperatures inducing partial deacclimation.

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