Physiological Research on Winter-hardiness: Deacclimation Resistance, Reacclimation Ability, Photoprotection Strategies, and a Cold Acclimation Protocol Design

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\textit{Abstract.} Freezing is a major environmental stress during an annual cycle of overwintering, temperate-zone perennials. The timing and extent of seasonal cold acclimation (development of freezing tolerance in the fall) and deacclimation (loss of acquired freezing tolerance in response to warm temperatures) are of critical importance for winter survival, particularly in view of the climate change, i.e., unpredictable extreme weather occurrences. For example, plants may acclimate inadequately if exposed to a milder fall climate and may be damaged by sudden frosts. Alternatively, they may deacclimate prematurely as a result of unseasonable, midwinter warm spells and be injured by the cold that follows. Efficient cold acclimation ability, high deacclimation resistance, and efficient reacclimation capacity are, therefore, important components of winter survival in overwintering perennials. These components should be evaluated separately for a successful breeding program focused on improving winter-hardiness. Another layer of complexity that should be carefully considered is that endodormant status (shallow versus deep) of the reproductive/vegetative apices can significantly impact these components of winter-hardiness. Winter survival, especially by woody evergreens, requires tolerance of light stress, which can result in photo-oxidative damage at cold temperatures when biochemistry of photosynthesis is somewhat compromised but light harvesting is unaffected. Accumulation of \textit{Elips} (early light-induced proteins) in overwintering evergreens during winter represents a relatively novel strategy to cope with such light stress, and investigations on the precise cellular mechanism and genetic control of this strategy deserve research in the future. Investigations into the mechanisms for cold acclimation use laboratory-based, artificial acclimation protocols that often do not closely approximate conditions that plants are typically exposed to in nature. To draw meaningful conclusions about the biology of cold acclimation and ultimately improve freeze resistance under field conditions, one should also include in cold acclimation regimens parameters such as exposure to subfreezing temperatures and realistic diurnal temperature fluctuations and light levels to simulate natural conditions. One of the main objectives of this article is to highlight two areas of research that we believe are important in the context of plant cold-hardiness but, so far, have not received much attention. These are: 1) to understand the biology of deacclimation resistance and reacclimation capacity, two important components of freeze-stress resistance (winter-hardiness) in woody perennials; and 2) to investigate the cellular basis for various strategies used by broad-leaved evergreens for photoprotection during winter. Our emphasis, in this context, is on a family of proteins, called \textit{Elips}. The second objective of this article is to draw attention of the cold-hardiness research community to the importance of using realistic cold acclimation protocols in controlled environments that will approximate natural/field conditions to be better able to draw meaningful conclusions about the biology of cold acclimation and ultimately improve freeze resistance. Results from our work with \textit{Rhododendron} (deciduous azaleas and broad-leaved evergreens), blueberry, and that of other researchers are discussed to support these objectives.

COLD ACCLIMATION

Freezing is a major environmental stress that can inflict injury to plant tissues. Plants, being poikilothermal, cannot escape damaging effects of extracellular ice and resultant cellular dehydration. Subfreezing temperatures, therefore, limit the productivity and geographical distribution of many wild and crop species. Considerable effort has been invested in understanding the cellular physiology, genetics, and molecular biology of plants’ response to subfreezing temperatures (Bertrand and Castonguay, 2003; Gusta et al., 2009; Guy, 1990, 2003; Moffatt et al., 2006; Pearce and Fuller, 2001; Smallwood and Bowles, 2002; Wisniewski et al., 2003; Xin and Browse, 2000). Review of this literature indicates that understanding the cellular basis of cold acclimation (CA) or cold hardening has been one of the major approaches in the past few decades to study plants’ response to subfreezing temperatures.

CA is the genetic ability of many species such as temperate-zone woody perennials to increase their freezing tolerance when non-acclimated (NA) tissues are exposed to inductive cues such as short days and/or low temperatures (as in the fall). It is one of the most-studied and best-characterized responses using both model systems and economically important species. Classic genetic studies have demonstrated that CA is a multigenic trait (Arora et al., 2000; Pan et al., 1994; Stone et al., 1993; Sutka, 1981). In recent years, systems biology research (using comparative transcriptomics, proteomics, and metabolomics with NA and CA tissues) coupled with mutational analysis of freezing tolerance (using mutants of model plants that are either “constitutively” or transgenically freeze-tolerant or lack CA ability) have significantly advanced our understanding of the signaling pathways and complex networks of molecular changes important for CA process (Browse and Lange, 2004; Hannah et al., 2005; Kaplan et al., 2006; Renaut et al., 2006; Van Buskirk and Thomashow, 2006; Warren, 1998).

WINTER-HARDINESS AND CLIMATE CHANGE: IMPORTANCE OF DEACCLIMATION RESISTANCE AND REACCLIMATION ABILITY

For winter survival, woody perennials not only must acclimate to cold, but also must resist premature deacclimation as a result of
unseasonable, midwinter warm spells. Deacclimation (DA) refers to reduction/loss of freezing tolerance originally attained through CA and, in nature, happens typically in early spring with the rise of temperatures. The topical interest in climate change particularly emphasizes the scenario of “premature or untimely DA” and resultant freeze damage. Temperate winters are becoming progressively milder as a result of climate warming, and temperature patterns are becoming increasingly irregular with risk of unseasonable warm spells (Arctic Climate Impact Assessment, 2005; Intergovernmental Panel on Climate Change, 2007), which can transiently deharden the plant tissues rendering them vulnerable to the risk of subsequent freezing injury (Kalberer et al., 2006). The historic 2007 Eastern US Spring Freeze, one of the most devastating in recent memory, exemplifies this phenomenon (Gu et al., 2008). Depending on the depth of DA, it may be either irreversible or reversed by subsequent exposure to low temperatures that may cause re-acclimation (RA), i.e., the restoration of, at least, a portion of the lost tolerance. Thus, in addition to the existence of sufficient and efficient CA ability, high DA resistance and RA capacity are also important components of winter survival of plants (Kalberer et al., 2006). Despite its critical importance to winter-hardiness, however, research on DA and RA has not received due attention.

LIGHT STRESS IN OVERWINTERING EVERGREENS

Another aspect of winter-hardiness that has not been well investigated, particularly in the context of broad-leaved evergreens, is cellular mechanisms of photoprotection during cold winters. Cold temperatures during winter can inhibit the enzymatic reactions of photosynthesis while not affecting the light absorption ability by overwintering evergreens. This can potentially result in photon flux in excess of that required for photosynthetic evolution of O2 [photosystem II (PSII) reaction centers] or assimilation of CO2. This excess energy, if not dissipated as heat or fluorescence, may cause inhibition of PSII reaction centers and/or photooxidative damage (Oquist and Huner, 2003). Evergreen species have evolved several mechanisms of photoprotection at suboptimal temperatures, and researchers have begun to address some of these only during the last decade (Adams et al., 2004; Demming-Adams and Adams, 2006; Niyogi, 1999; Verhoeven et al., 2005).

DEACCLIMATION, MIDWINTER-HARDINESS, AND ACCLIMATION CAPACITIES: ARE THEY LINKED?

As for the contribution of various components of overall winter-hardiness in woody perennials (as introduced earlier), it may be reasonable to assume that overwintering plants with high midwinter-hardiness would also exhibit a high degree of DA resistance and that a large maximal acclimation capacity and high DA resistance both represent evolutionary responses to low minimum temperatures, and the mechanisms responsible for these two processes might even be linked. To explore this notion, we initiated a study using deciduous azalea (Rhododendron L.) to examine the effect of genotypic or ecotypic biogeography on midwinter-hardiness and dehardening kinetics.

Nine azalea genotypes (species and varieties) were used in this study that represented eight seed provenances and multiple USDA cold hardiness zones (and midwinter-hardiness levels) making them good choices for comparative physiology. Natural habitats of these azaleas could be classified into three types: the southeastern lowlands, the Appalachian highlands, and the northeastern coastal region (Table 1) (Kalberer et al., 2007a). Buds from naturally (field) cold-acclimated plants were used (December to February) to evaluate midwinter freeze tolerance and deacclimation kinetics after exposure to a laboratory-controlled dehardening regime that included various durations (days) of deacclimation (DD; Kalberer et al., 2007a, 2007b). To investigate whether there was an association between spring temperature fluctuations and dehardening kinetics, the variation in temperature to which each genotype was exposed historically in its seed provenance was calculated (Kalberer et al., 2007a); temperature range of a provenance was calculated for each month by taking the difference between the monthly maximum and minimum 30-year temperature averages (National Oceanic and Atmospheric Administration, 2002).

Results on DA kinetics revealed two categories of azalea genotypes: slow deacclimators (high DA resistance) and fast deacclimators (low DA resistance) (Fig. 1). Data further indicated that no unequivocal relationship between proclivity to deharden (or lack thereof) and either the minimum temperature of habitats or the midwinter-hardiness could be established. For example, Rhododendron prinuliflorum (Small) Millais showed both low midwinter-hardiness and high DA resistance, whereas R. canadense (L.) Torr. had high midwinter-hardiness (the highest among all genotypes investigated) but low DA resistance; R. canadense (native to the north Atlantic states and freeze-hardy to −28.0 °C in December) deacclimated faster than the more sensitive Georgian species R. prinuliflorum (freeze-hardy to −24.6 °C) (Kalberer et al., 2007a). The recorded USDA hardness zones for R. canadense are 3b to 7 and the minimum average temperature for this provenance is −10.9 °C, whereas that for the provenances of R. prinuliflorum is warmer than −3 °C.

Hence, whereas the degree of midwinter-hardiness reflected the latitude and minimum temperatures of habitats, azaleas originating in cold climates and with high midwinter-hardiness did not always exhibit high DA resistance, which is perhaps related to other climatic and developmental factors. Others have also made similar observations; among blueberry cultivars (Rowland et al., 2005), potato species (Vega et al., 2000) and filbert varieties (Corylus L. spp.) (Hummer et al., 1986), high deacclimation resistance was not always associated with large acclimation capacities or high midwinter-hardiness. Similarly, ‘Concord’ grape (Vitis labrusca L.) had higher midwinter-hardiness than ‘Cabernet Sauvignon’ (Vitis vinifera L.), whereas the former deacclimated more rapidly (Wolf and Cook, 1992). Our recent work with Hydrangea species also indicates that a relatively harder species, H. paniculata Sieb., is a relatively faster deacclimator than the less-hardy H. macrophylla (Thunb.) Ser. (Pager et al., 2008, 2011). Furthermore, studies with cultivated potato indicated that the ability to acclimate rapidly is not always associated with a high CA capacity or high DA resistance; Solanum commersonii Dunn. attained higher levels of hardness (−9.3 °C) than five other potato species but acclimated more slowly than some species and deacclimated faster than others (Vega et al., 2000). Observations outlined previously suggest that high DA resistance, high midwinter-hardiness, and the rate of CA represent distinctly different attributes and are perhaps inherited independently. Therefore, they should be evaluated separately when selecting and breeding for improved winter-hardiness.

SELECTION FOR DEACCLIMATION-RESISTANCE IN PLANTS: WHAT MIGHT BE THE IMPETUS?

The question arises as to which environmental or biological factors likely promote high DA resistance in plants. Conceivably, DA resistance is not determined by a single parameter but is possibly a function of the degree of temperature fluctuations (frequency and magnitude), particularly during the late winter and early spring to which plants are exposed in their native habitats rather than the low temperatures per se. Conceivably, plants growing under relatively stable conditions would experience little evolutionary pressure to develop DA resistance to transient increases in temperature. In our azalea study, we noted that, on the whole, genotypes originating in the Appalachian mountains had relatively high DA resistance. We believe that the more variable temperatures of the Appalachians, in comparison with the coastal plains, may promote the evolution of higher DA resistance. On the other hand, R. canadense, the hardiest species investigated and that mostly inhabits lowland, coastal regions of the northeastern United States and the maritime provinces of Canada had low DA resistance; dehardening would be less of a threat under cool and stable temperatures during winter and early spring and thus would be little selective pressure for DA resistance. It may also be argued that the genotypes with greater midwinter-hardiness may experience less selective pressure for strong DA resistance than do less-hardy ones, because the former can safely lose a relatively large amount of acclimated hardness before becoming vulnerable to cold injury. Alternatively, high DA resistance could be prevalent in plants with delayed spring
development or deeper endo-/ecodormancy, because resumption of growth can lead directly or indirectly to irreversible deacclimation (Leinonen et al., 1997). Ample evidence exists to show that growth and development are not conducive to the retention of hardiness and that tissue hardiness is often negatively correlated with development (Mahfoozi et al., 2001a; Rapacz, 2002a, 2002b). Consistent with this notion, flower development of *R. prunifolium*, one of the slow deacclimator (DA-resistant) species used in our study, is slower than other native deciduous azaleas with anthesis in late summer, and a slow transition to active growth in these buds could explain the long-term maintenance of hardiness. The mechanism by which onogenetic development modulates hardiness has been the subject of some speculation but relatively little research. Growth and development could have a negative effect on hardiness by altering the subcellular structure. For example, the increased cellular water and reduced cytosol to vacuole ratios that accompany cell expansion can render plants more susceptible to mechanical damage from extracellular ice and even promote intracellular freezing, which is invariably lethal (Levitt, 1980; Stitt and Hurry, 2002; Strand et al., 1999). Alternatively, active growth might interfere with deacclimation resistance by competing for energy resources (Levitt, 1980; Stitt and Hurry, 2002; Strand et al., 1999). Conversely, rapid reacclimation may not necessarily always be deleterious to winter survival, especially if sufficient RA can quickly occur and somewhat mitigates the consequences of rapid dehardening. Even more desirable for a plant would be to have sufficient and efficient RA capacity even after a relatively long dehardening duration or substantial DA. Of course, for RA capacity to play a beneficial role in winter-hardiness, it is critical that the return of cold after premature DA is not too severe (in both the extent and suddenness) so as to allow sufficient reacclimation, which may not always happen in nature.

In our azalea study, the capacity to reharshen to various extents was observed in diverse species from three regions after various amounts of dehardening: the southeastern lowlands (*R. viscosum* (L.) Torr. *serrulatum* (Small) Millais), the Appalachians (*R. arborescens* (Pursh) Torr.), *R. viscosum* var. *montanum* Rehd., and the northeastern coastlands (*R. canadense*) (Kalberer et al., 2007a, 2007b). Our results also indicated that only 24 h of low-temperature exposure (12 h at 2°C followed by 12 h at −2°C) was sufficient to reharshen azalea buds post-DA and that RA occurred even after substantial dehardening in several genotypes (Kalberrer et al., 2007a, 2007b). Given that seasonal CA is a relatively gradual process, it is remarkable that dehardened buds of *R. canadense* and of the two latitudinal ecotypes, *R. viscosum* var. *montanum* and *R. viscosum* var. *serrulatum*, could regain up to ≈5, 11, and 9°C of freeze tolerance, respectively, after only 1 d of exposure to a reacclimation regime (Kalberer et al., 2007a). Our data also indicated that the RA capacities in these azalea genotypes did not have a significant association with DA resistance. For example, two of the fast-deacclimator genotypes, *R. canadense* and *R. viscosum* var. *serrulatum*, exhibited highly divergent RA capacities; the former regained ≈5°C in response to a 24 h RA regime after 1 DOD, whereas the latter recovered only 0.6°C with an identical treatment. No conclusions could be drawn about associations, if any, between the natural habitat and RA capacities in this study, and greater sample size will need to be examined to explore these relationships. Nevertheless, these results highlight that genetic variation, both inter- and intraspecific, for RA capacity exists in woody perennials and should be exploited in the breeding programs for improving winter-hardiness.

### Table 1. Deciduous azalea (*genus Rhododendron*) genotypes used in this study with common names and seed provenances (when applicable).

<table>
<thead>
<tr>
<th>Rhododendron genotype (common name)</th>
<th>Hardiness zones</th>
<th>Seed provenance</th>
<th>Provenance minimum temp. (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>canadense</em> (rhodora)</td>
<td>3b to 7</td>
<td>Long Pond in Poconos Mountains, PA</td>
<td>−10.9</td>
</tr>
<tr>
<td><em>prinophyllum</em> (rose shell)</td>
<td>4 to 8</td>
<td>Spruce Knob, WV</td>
<td>−8.6</td>
</tr>
<tr>
<td><em>viscosum</em> var. <em>montanum</em> (upland swamp)</td>
<td>—</td>
<td>Pisgah Swamp in Pisgah National Forest, NC</td>
<td>−5.0</td>
</tr>
<tr>
<td><em>arborescens</em> (sweet)</td>
<td>5a to 8b</td>
<td>Wayah Bald near Franklin, NC</td>
<td>−4.4</td>
</tr>
<tr>
<td><em>calendulacum</em> (flame)</td>
<td>5a to 8b</td>
<td>Little Canada near Wolf Creek Reservoir, NC</td>
<td>−3.9</td>
</tr>
<tr>
<td><em>canescens</em> (piedmont)</td>
<td>6b to 9</td>
<td>Moody Springs, SC</td>
<td>−2.9</td>
</tr>
<tr>
<td><em>atlanticum</em> (coastal)</td>
<td>6b to 8b</td>
<td>Western border of Delaware</td>
<td>−2.8</td>
</tr>
<tr>
<td><em>prunifolium</em> (plumleaf)</td>
<td>6b to 9a</td>
<td>Providence Canyon Park, GA</td>
<td>0.6</td>
</tr>
<tr>
<td><em>viscosum</em> var. <em>serrulatum</em> (hammock-sweet)</td>
<td>7a to 9a</td>
<td>Provenance unknown</td>
<td>—</td>
</tr>
</tbody>
</table>

*aUSDA hardiness zones represent species distributions assigned by Galile (1974).*

*Seed provenance information was provided by Natural Landscapes Nursery (West Grove, PA).*

*Minimum temperatures represent 30-year averages of the lowest monthly temperatures recorded during January of each year. Measurements are taken from the weather station closest to the provenance (National Oceanic and Atmospheric Administration, 2002), and the station is always located in the same state as the provenance.*

*No distribution of *R. viscosum* var. *montanum* is available in the scientific literature, but *R. viscosum* is distributed from 4a to 9a (Galile, 1974).*

Source: Kalberer et al., 2007a.

*Fig. 1. Logarithmic-linear models of frost hardiness (LT50) as a function of days of dehardening (DOD) for nine azalea genotypes. *Rhododendron canadense*, *Rhododendron canescens*, and *Rhododendron viscosum* var. *serrulatum* had dehardening kinetics representative of low dehardening resistance. The dehardening kinetics of *Rhododendron arborescens*, *Rhododendron atlanticum*, *Rhododendron calendulacem*, *Rhododendron prinophyllum*, *Rhododendron prunifolium*, and *Rhododendron viscosum* var. *montanum* represented high dehardening-resistance (Source: Kalberer et al., 2007a).*
Research shows that although reacclimataation ability may be present in many overwintering plants, the RA capacity becomes more limited as the degree or duration of warm temperature exposure increases and DA advances. Apple bark tissues could reacclimate only to the hardiness level exhibited on the day previous to the final day of DA (Howell and Weiser, 1970). Gusta and Weiser (1972) exposed Korean boxwood to alternating temperature cycles (25 °C followed by −10 °C) and found that although DA was fully reversible after one such cycle, RA capacity declined with subsequent cycles. Whether such loss of RA capacity is the result of a lack of energy-producing substrates (required for acclimation), or irreversible developmental changes after deacclimation, or some other factor, or the combination of these factors, has not been determined.

DEHYDRIN METABOLISM AND DEACCLIMATION/REACCLIMATION CYCLING

Dehardening reverses physiological adaptations acquired during exposure to cold that endow plants with enhanced frost-hardiness. Numerous studies have reported an accumulation of dehydrins, hydrophilic members of the late embryogenesis abundant class of proteins, in bark, leaves, buds, and other tissues during CA of woody perennials ( Marian et al., 2003). Dehydrin protein and transcript abundance declined to levels found in non-hardened plants during the dehardening of floral buds of blueberries (Vaccinium L.). (Arora et al., 1997) and bark tissues of peach (Prunus persica (L.) Batsch) ( Artlip et al., 1997). Levels of a 60 kDa dehydrin decreased on resumption of growth in needles of Scots pine (Pinus sylvestris L.) ( Kontunen-Soppela et al., 2000). The rationale for the functional significance of such accumulation is that dehydrins, as a result of their hydrophilicity and putative chaperone properties, likely protect cellular membranes, enzymes, and macromolecules from freeze desiccation (Danylik et al., 1998; Koag et al., 2003; Rime et al., 1999). We observed that, in general, dehydrin levels fell and rose in response to changing ambient temperatures and were associated with loss and gain in bud freeze tolerance during dehardening and rehardening, respectively (Fig. 2) ( Kalberer et al., 2007a). These observations are consistent with the putative role of dehydrins in freeze tolerance and provide support for potential use of dehydrin metabolism as a biochemical marker in dehardening and rehardening studies.

DEACCLIMATION RESISTANCE, REACCLIMATION CAPACITY, AND WINTER PROGRESSION

Another important question that needs to be examined is: how does winter progression, i.e., early versus late winter, influence DA resistance and RA capacity. To address this, we refer again to the azalea studies, with selected genotypes, in December (early winter) and in February (deep winter). In the Dec. 2004 study, rehardening in several azalea genotypes was observed even after only 1 DOD. However, in the Feb. 2006 study, rehardening did not occur until after 10 to 15 DOD. For example, in Dec. 2004, the buds of R. viscosum var. montanum rehardened by ≈4 °C even after 1 DOD (i.e., after dehardening by ≈3 °C), whereas they rehardened by ≈2 to 3 °C only after 10 to 15 DOD by which time they had lost ≈14 to 16 °C in freeze tolerance. Similarly, R. canadense buds were able to reharden in December by ≈5 °C after losing ≈9 °C in freeze tolerance after 1 DOD compared with 1.4 °C of rehardening in the February study after 5 DOD and losing 18 °C in freeze tolerance (Kalberer et al., 2007a, 2007b). The ability of buds to reharden only when exposed to a relatively long dehardening duration suggests that, during the late winter, buds reharden only after a “threshold” level of dehardening is reached. Perhaps levels of certain proteins or metabolites may need to decrease, or water content may need to increase, beyond a certain point during dehardening before rehardening can be induced. Alternatively, it may take a longer period of time to reprogram the tissues to cold-acclimate once they have begun to resume growth and development. Therefore, could the difference in RA capacities of buds between the Dec. 2004 and Feb. 2006 studies be related to differences in the depth of endodormancy? In other words, could it be that the greater depth of dormancy in December (as a result of less fulfillment of chilling requirements), compared with in February (i.e., shallower dormancy resulting from greater accumulation of chill-units), favorably influences rehardening even after low levels of dehardening?

DORMANCY AND DEACCLIMATION RESISTANCE OR REACCLIMATION CAPACITY

Dormancy can be defined as a temporary suspension of visible growth of any plant structure containing a meristem (Lang et al., 1997). As understory evergreens in the deciduous forests, leaves of most broad-leaved rhododendrons are commonly exposed to a combination of freezing temperatures and light during their natural habitat during winter. Our expressed sequence tag (EST) study with NA and CA leaf tissues of R. catawbiense Michx. indicated a downregulation of several photosynthesis-related genes (e.g., RuBisco small subunit precursor, RuBisco activase, plastidic fructose bisphosphate aldolase, chloroplast precursor of plastocyanin) in overwintering leaves suggesting that it could potentially lead to light energy harvested by the leaves to be in excess of what can be processed by photosystems. This research also revealed that cDNAs
encoding Elip homologs were the most abundant class in an EST library generated from winter-collected (CA) leaf tissues and no Elips were detected in the EST data set for NA (summer-collected) leaves (Wei et al., 2005); these Elips ESTs were later catalogued into seven distinct RcElips (for R. catawbiense Early light induced proteins) (Peng et al., 2008). Elips are nuclear-encoded, light-stress-induced proteins located in thylakoid membranes and belong to the chlorophyll a/b-binding protein family with a wide distribution among plant species (Adamska, 2001). It is proposed that Elips may transiently bind the released chlorophylls under high light stress and prevent the formation of free radicals and/or function in energy dissipation (Adamska, 2001; Montané and Klopstech, 2000). Elip accumulation, therefore, may constitute an adaptive response to winter conditions (cold and high light) in evergreens and play a key role in the protection of photosynthetic apparatus from excess light. Our more recent work revealed a significant upregulation of the transcripts of seven RcElips from August to December in two Rhododendron species, a less hardy R. ponticum L. (midwinter leaf-freeze tolerance of \(\approx -20 \, ^\circ\text{C}\)) and superhardy R. catawbiense (leaf-freeze tolerance of \(\approx -50 \, ^\circ\text{C}\)) (Wang et al., 2009). Further investigation of the seasonal profiles at the protein level should be beneficial to uncover the possible role of RcElips in these two species. Moreover, a strong positive correlation between the degree of CA ability and corresponding increment in Elips accumulation in the two species divergent in their freezing tolerance (Fig. 3) suggests that an enhanced ability of leaves to protect chloroplast from excess light may be one of the key components of a multifactorial cold acclimation process in evergreen rhododendrons. Others have also reported an accumulation of Elips proteins in winter-collected leaves of several evergreens—subalpine fir and lodgepole pines (Zarter et al., 2006a) and bearberry (Zarter et al., 2006b) but little to no accumulation in the summer-collected samples. Collectively, these studies point toward a potential role of Elips in the photoprotection scheme by overwintering evergreens and further research is warranted to fully understand their functional significance. However, other possible, heretofore unknown, functional roles of Elip upregulation in conferring freezing tolerance cannot be ruled out. Interestingly, the rate of seasonal increase in Elip abundance (per unit change in leaf freeze tolerance calculated as the LT_{50}) for R. catawbiense was twice that for R. ponticum, suggesting that R. catawbiense perhaps needs more efficient upregulation of photoprotection systems than R. ponticum. This notion is, curiously, bolstered by the fact that R. catawbiense exhibits thermonasty, whereas R. ponticum does not. Thermonasty refers to temperature-induced leaf movements, a phenomenon in which leaves droop and curl at freezing temperatures in winter (Nilsen, 1987). One of the proposed adaptive benefits (among others) of thermonastic leaf movement is the avoidance of high light stress in the winter by reducing leaf exposure to light (Bao and Nilsen, 1988) and thereby facilitating relatively faster recovery of PSII efficiency in spring (Russell et al., 2009). Lack of thermonastic behavior in R. ponticum, therefore, suggests that this species may perhaps be more tolerant of light stress in winter than R. catawbiense. Our results on the seasonal patterns of PSII efficiencies in the two species indicate that R. catawbiense leaves underwent significantly greater photoinhibition during fall and winter compared with R. ponticum, indicating relatively higher sensitivity to light stress of the former (Wang et al., 2009).

### Natural (Field) Versus Artificial (Controlled Environment) Cold Acclimation: Appreciating the Differences in Context of Research Outcomes

Another issue to address, as we try to further our understanding of the development of cold-hardiness in plants, and in woody perennials in particular, is the choice of experimental protocols for achieving CA. In the literature, we often find that researchers acclimate plants under artificial regimes such as...
as placing potted plants in growth chambers, typically at a constant \(4{\degree}C\), with fairly low light levels, for 1 to several weeks. This protocol has the advantage of being fairly "controllable" and may be sufficient for herbaceous annuals, which primarily acclimate in response to above-freezing, low temperatures over a short duration and by only a few degrees. Woody perennials, on the other hand, generally achieve greater levels of cold-hardiness than do herbaceous annuals, being able to withstand lower freezing temperatures over a longer duration. In woody perennials, CA proceeds in stages and is triggered by several environmental cues—short photoperiod, low temperatures (both above and below freezing), available moisture, etc. In nature, these factors change gradually—decreasing daylengths, declining temperatures into the freezing and below-freezing ranges, and declining moisture. Also, in nature, there are generally warmer day and cooler night temperatures. The light intensity is typically higher than in cold room acclimation regimes, and the light quality is different and varies seasonally.

In our studies on a typical northern highbush blueberry cultivar, Bluecrop, we have found that flower buds of field plants (in Beltsville, MD) have an LT50 of \(\approx 13{\degree}C\) in October and reach a minimum flower bud LT50 (maximum level of cold-hardiness) of \(\approx 27{\degree}C\) by mid-December, when plants have accumulated \(\approx 600\) chilling-units (Fig. 4). Cold-hardiness levels tend to decline in February and March with the return of warmer temperatures. For the southern rabiteye blueberry cultivar Tifblue, flower buds have an LT50 of \(\approx 12{\degree}C\) in October and an LT50 of \(\approx 25{\degree}C\) in mid-December (Fig. 4). Cold-hardiness levels begin declining in January. In contrast, under cold room conditions of constant 4 \(\degree\) C and a constant photoperiod (10 h light/14 h dark), 'Bluecrop' and 'Tifblue' plants reach a maximum level of cold-hardiness of \(\approx 24{\degree}C\) and \(\approx 17{\degree}C\), respectively (3 to 8 \(\degree\)C less hardy than field-acclimated plants) by \(\approx 500\) h (\(\approx 3\) weeks). Cold-hardiness levels remain constant as long as plants are kept in the cold room (Arora et al., 1997).

In recent years, we have taken a genomic approach to identify genes associated with CA in blueberry by generating ESTs from cDNA libraries prepared from NA and CA flower buds of the northern highbush cultivar Bluecrop and using these ESTs to construct a microarray. The original microarray (for blueberry) included \(\approx 2400\) cDNAs, 1200 from each of the NA and CA libraries. This microarray was first used to compare changes in gene expression of 'Bluecrop' plants acclimated under two different regimes—naturally under field conditions and in pots in a cold room environment with constant temperature (4 \(\degree\) C) and a constant photoperiod (10 h light/14 h dark). Changes in levels of gene transcripts were examined in flower buds at multiple times during cold acclimation under field (\(\approx 0, 70, 400, 800,\) and 1200 chill-units) and coldroom conditions (0, 500, and 1000 chill-units) (Dhanaraj et al., 2007; Rowland et al., 2008a). First, results indicated that many of the same genes were induced under both environments. These included many genes previously identified as part of the cold-response pathway in Arabidopsis (Fowler and Thomashow, 2002; Hannah et al., 2005; Seki et al., 2001) such as genes encoding galactinol synthase, beta amylase, LEAs, dehydrins, and ELIPs and also genes not previously observed to be cold-induced in Arabidopsis such as genes encoding protein kinase PINOID involved in...
auxin-mediated signaling, pectate lyase, and S-adenosylmethionine decarboxylase proenzyme. However, marked differences in gene expression were also observed under the two CA regimes. In general, the number of cold-induced genes was higher and the number of cold-suppressed genes was lower in the cold room than in the field (Fig. 5). Indeed, the number of genes induced under cold room conditions was approximately twice that induced under field conditions, although ‘Bluecrop’ field plants are 3°C more cold-hardy than cold room plants, indicating that all the induced genes in the cold room environment likely do not contribute to increased hardness. It is possible that some of the genes induced in the cold room may be associated with low-temperature growth acclimation rather than the freezing tolerance per se. Many of the genes induced under cold room conditions that were not induced under field conditions could be divided into three major groups: 1) general stress tolerance genes; 2) genes encoding glycolytic and tricarboxylic acid cycle enzymes; and 3) genes encoding protein synthesis machinery. Some of the genes induced under both environments, particularly genes associated with light stress, were induced to higher levels under field conditions than a cold room. Furthermore, many of the genes suppressed under field conditions, that were not suppressed under cold room conditions, reached maximum suppression by the 400 chill-unit sampling, at which point average outside temperatures had dropped below freezing to –2.1°C from an average of 6.3°C the week before. Thus, it is possible that these genes may have been suppressed in response to freezing temperatures, explaining their lack of suppression in the cold room environment. Many of these suppressed genes were in fact the same general stress genes that were induced in the cold room setting such as genes encoding heat shock proteins, phospholipid hydroperoxide glutathione peroxidase, and low temperature-induced 78 kD protein. From this, it is interesting to speculate that, in ‘Bluecrop’, cold room conditions may result in induction of a larger general stress response, whereas freezing conditions in the field result in more selective expression.

Just to complicate matters even more, we have also now compared gene expression of the southern rabbiteye cultivar ‘Tifblue’ under field and cold room acclimation regimes (unpublished data). Surprisingly, in contrast to ‘Bluecrop’, more genes were induced in ‘Tifblue’ under field conditions than under cold room conditions (Fig. 5). Some of the genes induced in ‘Tifblue’ in the field, that were not induced in the cold room, were genes that were highly induced in both environments in ‘Bluecrop’ such as genes encoding galactinol synthase, a cell wall glycine-rich protein, and water stress ER-5. Thus, not only do plants respond differently under these different CA regimes, but their responses are genotype-, or perhaps ecotype-, dependent. It appears that the southern cultivar ‘Tifblue’ may require lower temperatures, or a different light regime than was provided in the cold room, to induce its complete suite of cold-responsive genes.

Using wheat as a model, Herman et al. (2006) found that subzero acclimation or SZA (an additional 3 to 5°C increase in freezing tolerance upon exposure to –3°C) is accompanied by...
changes in physiology, cellular structure, the transcriptome, and the proteome distinct from those observed during a “typical” CA regime (freezing tolerance acquired at above freezing temperatures). In terms of the transcriptome, they found that many unknown and stress-related genes were upregulated, whereas many photosynthesis and plastid-related genes were downregulated during SZA as compared with “typical” CA. From a proteome standpoint, they noted a loss of many proteins as opposed to the appearance of new proteins in SZA samples. Thus, like in blueberry, subfreezing temperatures appeared to result in much suppression of protein expression.

In summary, these studies highlight the need for researchers to begin using more realistic CA protocols in controlled environments that more accurately approximate natural/field conditions. For woody perennials, in particular, a subzero temperature regime should be included to achieve full CA potential and the full complement of CA-related gene expression.

CONCLUDING REMARKS

Predicted global climate change may have dramatic effects on the extent and location of winter injury because the frequency of extreme and unseasonable weather events are likely to increase in the future. Such scenarios could disturb the timing of seasonal CA and DA in overwintering woody perennials. Therefore, the plants that are able to resist premature, rapid DA, as well as quickly RA when cold returns, might have a better chance of surviving cold winters. Discussion outlined here suggests that winter survivability of perennials is a complex, multicomponent response in which early initiation of acclimation, high midwinter-hardiness, high DA resistance, and efficient RA capacity all represent distinctly different components of this complex. For a successful breeding program focused on improving winter-hardiness, it is important to determine which aspect of winter-hardiness, if not all (acclimation timing and speed, midwinter-hardiness, DA resistance, RA capacity), might be the most limiting for a given species and a location and accordingly select appropriate parents to introduce these traits. Another layer of complexity that should be carefully considered, in this context, is that endodormant status (shallow versus deep) of the reproductive/vegetative apices can significantly impact these components of winter-hardiness.

Winter survival, especially by woody evergreens, is not only coping with the cold (and associated desiccation), but also requires tolerance of other environmental factors that may become stressful even when present at ambient levels; a case in point: light, which can result in photo-oxidative damage at cold temperatures when the enzymatic facet of photosynthetic processes is sluggish. Overwintering evergreens have evolved with various photoprotection strategies to manage such stress, including xanthophyll cycle, antioxidant systems, certain adaptive anatomical as well morphological features, etc. Accumulation of Elips in overwintering evergreens during winter represents a relatively novel strategy to cope with light stress. In-depth investigations on the mechanistic physiology and genetic control of this strategy deserve attention in the future.

Finally, recent genomic research comparing plants under different CA regimes (natural/field versus artificial/cold room and low/above freezing versus subzero freezing temperatures) indicates that there are major differences in gene expression under the different environments, and the differences may be genotype-dependent. To draw meaningful conclusions about the biology of CA and ultimately improve freeze resistance under real-life conditions, more efforts should be made by the cold-hardiness research community to design experiments to sort out the major reasons for these differences such as presence of subfreezing temperatures and higher light levels under field conditions, as implicated in the experiments described here. Once this is determined, more realistic CA protocols in controlled environments can be developed and implemented.

Literature Cited


