Impact of Temperature on Seed Dormancy

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Seed dormancy is an ecological adaptation that allows seasonal timing of germination for seeds in a population. Several environmental stimuli can trigger dormancy release, the most important being seed moisture content, light, and temperature. Of these, temperature is arguably the most important. The objective of this brief review is to consider the various ways temperature impacts seed dormancy release. It will cover general aspects of seed dormancy, temperature effects on primary dormancy and considerations for conducting and interpreting seed dormancy research.

Propagators of cultivated plants have long recognized that germination-delaying mechanisms exist in seeds. The first recorded discussion of seed dormancy was written by Theophrastus in ≈300 B.C. (Evenari, 1981). He recognized that germination of most seeds declined during storage (seed deterioration), while germination in some seeds increased (dormancy release).

One problem with discussing seed dormancy is that there is no single recognized terminology to describe the many different types of seed dormancy. Crocker (1916) described seven types of dormancy based on treatments used to overcome them. Subsequently, Nikolaeva (1977) defined dormancy based primarily upon physiological controls. More recently, Lang (1987) proposed the terms eco-, para-, endo-dormancy to simplify terminology. This system is currently utilized in American Society for Horticultural Science journals. However, this terminology is not sufficient to adequately describe all the types of dormancy found in seeds. Baskin and Baskin (1998) have developed the most complete set of terms to describe seed dormancy. They have extended the dormancy classifications of Nikolaeva to include additional specialty types. In this review, I will use a system based on the work of Nikolaeva, as modified by Baskin and Baskin (Hartmann et al., 2002). It is a classification system that I feel fits both the ecological and horticultural aspects of seed dormancy and is accepted by most of the scientists working in the field of seed biology (Table 1).

Table 1. Seed dormancy categories

<table>
<thead>
<tr>
<th>Type of Dormancy</th>
<th>Description</th>
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<tbody>
<tr>
<td>Primary dormancy</td>
<td>Dormancy that occurs in seeds that were previously non-dormant, but have become dormant because the environment was unfavorable for germination. Primary dormancy includes exogenous dormancy, endogenous dormancy, and combination dormancy. Secondary dormancy includes thermodormancy and conditional dormancy. These types of dormancy and the requirements to overcome them are listed in Table 1. Detailed descriptions of these categories can be found elsewhere (Baskin and Baskin, 1998; Geneve, 1998; Hartmann et al., 2002). In this review, I will limit each category to a brief description and emphasize dormancy conditions that are affected by temperature.</td>
</tr>
<tr>
<td>Primary exogenous dormancy</td>
<td>In exogenous dormancy, the tissues enclosing the embryo impact germination by either inhibiting water uptake, modifying gas (O2) exchange, or possibly contain germination inhibitors (Bewley and Black, 1994). Seeds of species with exogenous physical dormancy fail to imbibe water because of properties of the seed coverings. This form of seed dormancy occurs in 15 plant families (Baskin et al., 2000). Of these, most of the species displaying physical dormancy are found in the Malvaceae and Fabaceae. The anatomical structures preventing water uptake can be the seedcoat (testa) or endocarp (Baskin et al., 2000). In most species, there are elongated palisade cells in the outer layer of the seedcoat (exocotyledon) that prevent imbibition. Mechanical abrasion or chemical degradation of the seed coverings and submergence of the seed in hot water are the most common horticultural practices to overcome exogenous dormancy. Collectively, these treatments are termed scarification. However, in nature, it appears that temperature is the major factor determining water uptake in seeds with physical dormancy. Temperature impacts dormancy release for seeds with exogenous physical dormancy by affecting the seed coverings. For instance, some seeds require high temperature or daily fluctuations (&gt;15 °C change) in temperature to allow imbibition. This requirement is postulated as a way for seeds to detect whether they are in open or protected areas (Baskin and Baskin, 1998). Higher seeds with physical dormancy, as well as a greater day/night fluctuation, would occur in an open area, indicating less competition from other plants after germination. The coverings of seeds with physical dormancy may also be cracked by temperature fluctuations, alternate freezing and thawing and in some species by fire. For many seeds with physical dormancy, a specialized location on the seed coverings can act as an “environmental sensor.” In the Fabaceae, it is usually the lens (strophiole) (Manning and van Staden, 1987; Morrison et al., 1998); and in the Malvaceae, it is the chalazal plug (Egley, 1989). These structures are disrupted by temperature and become the site of water entry into the seed. For example, Quinlivan (1968) demonstrated that seeds of Lupinus varius L. became permeable to water at the lens after exposure to fluctuating temperature (i.e., 65 °C day temperatures with night temperatures down to 25 °C).</td>
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Primary endogenous dormancy

The second major category of primary seed dormancy is endogenous seed dormancy. Seeds with endogenous dormancy fail to germinate because of factors associated with the embryo. There are two types of endogenous dormancy—a morphological and physiological.

Morphological dormancy. Seeds with morphological dormancy have an embryo that is not fully developed at the time of seed disemination. Seeds where the embryo fills less than half of the seed are considered to have morphological dormancy (Baskin and Baskin, 1998). Enlargement of the embryo occurs after the seeds have imbibed water, but usually before germination begins. The process of embryo enlargement is influenced by temperature. Atwater (1980) distinguished three categories of morphological dormancy based on the embryo type found in herbaceous flower crops. These are rudimentary, linear, and undifferentiated embryo types. Rudimentary embryos are little more than a proembryo embedded in a massive endosperm. These are found in seeds of various families, such as the Ranunculaceae and Araliaceae. Germination-inhibiting chemicals may occur in the endosperm and become active at high temperatures. Methods for inducing germination include: (a) exposure to temperatures of 15 °C or below; (b) exposure to alternating temperatures; and (c) treatment with chemicals such as potassium nitrate or gibberellic acid. Seeds with linear embryos are torpedo-shaped and up to one-half the size of the seed. Important families and species in this category include the Apiaceae, Ericaceae, Primulaceae, and Gentianaceae. Conditions such as semipermeability of the inner seedcoats and internal germination inhibitors may be involved. Temperature >20 °C favors germination, as does treatment with gibberellic acid.

Some tropical species have seeds with small embryos that require an extended period at warm temperatures for germination to take place. For example, seeds of some palm spe- cies require 1 to 3 months at high temperatures (≥35 °C) to complete germination (Nagao et al., 1980). Other examples include Actinidia sp. and Annona squamosa L., whose seeds require 2 or 3 months at warm temperatures, respectively, to complete germination (Nikolaeva, 1977).
at the time theseed sheds from the plant

Chemical Inhibitors in seed coverings Removal of seed coverings (fruits)

1. Primary dormancy

<table>
<thead>
<tr>
<th>Causes of dormancy</th>
<th>Conditions to break dormancy</th>
<th>Representative genera</th>
</tr>
</thead>
<tbody>
<tr>
<td>Physical</td>
<td>Impermeable seedcoat</td>
<td>Scariﬁcation</td>
</tr>
<tr>
<td>Chemical</td>
<td>Inhibitors in seed coverings</td>
<td>Removal of seed coverings (fruits)</td>
</tr>
<tr>
<td>Morphological</td>
<td>The embryo is not fully developed</td>
<td>Warm or cold stratiﬁcation</td>
</tr>
<tr>
<td>Rudimentary</td>
<td>Small undifferentiated embryo</td>
<td>Cold stratiﬁcation and potassium nitrate</td>
</tr>
<tr>
<td>Linear</td>
<td>Small differentiated embryo &lt;1/2 size of seed</td>
<td>Warm stratiﬁcation and gibberellic acid</td>
</tr>
<tr>
<td>Physiological</td>
<td>Factors within embryo inhibits germination</td>
<td>Red light</td>
</tr>
<tr>
<td>Nondeep</td>
<td>Positively photodormant (requires light)</td>
<td>Darkness</td>
</tr>
<tr>
<td>Intermediate</td>
<td>Embryo germinates if separated from the seedcoat</td>
<td>Short period of dry storage</td>
</tr>
<tr>
<td>Deep</td>
<td>Embryo does not germinate when removed from seedcoat or will form a physiological dwarf</td>
<td>Moderate periods (up to 8 weeks) of cold stratiﬁcation</td>
</tr>
<tr>
<td>c. Combinational</td>
<td>Combinations of different dormancy</td>
<td>Long periods (&gt;8 weeks) of cold stratiﬁcation</td>
</tr>
<tr>
<td>Morphophysiological</td>
<td>Embryo and physiological dormancy</td>
<td>Cycles of warm and cold stratiﬁcation</td>
</tr>
<tr>
<td>Epicotyl</td>
<td>Radicle begins growth when temperature and water permit, but epicotyl is dormant</td>
<td>Warm followed by cold stratiﬁcation</td>
</tr>
<tr>
<td>Epicotyl and radicle (double dormancy)</td>
<td>Radicle and epicotyl require chilling stratiﬁcation, but radicle is released during first year and then</td>
<td>Cold stratiﬁcation followed by warm followed by a second cold stratiﬁcation</td>
</tr>
<tr>
<td>Exo-endodormancy</td>
<td>Combinations of exogenous and endogenous dormancy conditions. Example: physical (hard seedcoat) plus intermediate physiological dormancy</td>
<td>Sequential combinations of dormancy releasing treatments. Example: scariﬁcation followed by cold stratiﬁcation</td>
</tr>
</tbody>
</table>

2. Secondary dormancy

a. Thermodormancy

After primary dormancy is relieved, high temperature induces dormancy

b. Conditional dormancy

Change in ability to germinate related to time of the year

Orchid seeds have undifferentiated embryos when shed from the mother plant. They are not considered dormant in the same sense as others in this category because they lack substantial seed storage materials.

Physiological dormancy. The second type of endogenous dormancy is physiological dormancy. This type of dormancy involves physiological changes within the embryo allow the radicle to escape the restraint of the seed coverings. Physiological dormancy includes nondeep, intermediate, and deep categories. Endogenous, nondeep, physiological dormancy is the most common form of seed dormancy (Baskin and Baskin, 1998). This type of dormancy can be broken by light or darkness (photodormancy), short periods of chilling stratiﬁcation, or “after-ripening” (dry storage).

Seeds from species with endogenous, nondeep, physiological dormancy (especially small seeded species) often require light or darkness for germination. Light sensitivity in seeds is a photochrome response (Casal and Sánchez, 1998). For some seeds, there is a distinct light x temperature interaction regarding dormancy and germination. A light requirement can be offset by cool temperatures and sometimes, by alternating temperatures. Seeds of some cultivars of lettuce generally require light to germinate; however, they can germinate in darkness at temperatures below 25 °C (Hattangy, 1972). For years, birch (Betula sp.) seeds were thought to require moist chilling to permit germination. However, nonchilled seeds will germinate in light (Vanhatalo et al., 1996).

Freshly harvested seeds of some herbaceous plants display endogenous, nondeep physiological dormancy (Association of Ofﬁcial Seed Analysts, 1993; Atwater, 1980; Baskin and Baskin, 1998). This type of dormancy is often transitory and disappears during dry storage (after-ripening). For most cultivated cereals, grasses, vegetables, and flower crops, nondeep physiological dormancy may last for 1 to 6 months and disappears with dry storage during normal handling procedures (Geneve, 1998).

Temperature affects the time required to after-ripen seeds. For example, cultivated cucumber (Cucumis sativus var. sativus L.) has been selected over many years of cultivation for a short dormancy period. Freshly harvested seeds lose dormancy in dry storage at room temperature after several weeks (15 to 30 d). The hardwickii cucumber [Cucumis sativus var. hardwickii (Royle) Alte.] is considered a wild progenitor species of the cultivated cucumber and seeds can remain dormant for 60 to 270 d (Weston et al., 1992). Dormancy release occurred much earlier in hardwickii seeds held in dry storage at warmer temperatures (180 d at 17 °C compared to only 75 d at 37 °C). Roberts (1965) showed that there was a negative linear relationship between after-ripening time and temperature that was consistent within a species. He showed that time on a log scale to reach 50% germination was linear with temperature.

Moist chilling stratiﬁcation can relieve dormancy in species with nondeep, intermediate, or deep physiological dormancy (Fig. 1). It is often difﬁcult to differentiate between seeds that display nondeep vs. intermediate physiological dormancy. In general, seeds with nondeep dormancy require only short periods (days or up to several months) of chilling stratiﬁcation to relieve dormancy. Seeds with intermediate physiological dormancy usually require at least 2 months of chilling or gibberellin application can substitute for chilling. Embryos isolated from their seed coverings in nondeep or intermediate physiologically dormant seeds will germinate promptly and produce normal seedlings. In contrast, seeds with deep physiological dormancy require long periods (>3 months) of chilling stratiﬁcation to relieve dormancy and generally do not respond to exogenous gibberellin application. Embryos isolated from these seeds either will not germinate or will grow into abnormal seedlings with a dwarf phenotype, termed physiological dwarfs (Flemion and Waterbury, 1945).

In most cases, seeds with endogenous physiological dormancy respond in a similar manner with regard to stratiﬁcation temperature. A temperature near 4 °C has maximum effect, while below freezing temperatures or...
temperatures above 14 to 16 °C are generally not effective for relieving dormancy (Seeley, 1997). This observation has led to the concept of stratification degree hours for predicting the time required to relieve dormancy (Seeley and Damavandy, 1985). One hour at optimum stratification temperature (4 °C) is equal to one "stratification degree hour." Partial stratification degree hour values are assigned to warmer or cooler temperatures, but for durations below freezing and above 16 °C have no effect toward dormancy release. In addition, time above 16 °C can negate previous chilling.

Some species have seed lots where some seeds may not require stratification to germinate but the rate of seedling emergence is improved for all seeds by brief exposure to chilling temperatures. This phenomenon has been referred to as a facultative form of physiological dormancy (Geneve, 1998). Genera in this group include Anistruthium, Eustoma, and Impatiens (Ecker et al., 1994; Montero et al., 1990; Simmonds, 1980) and various conifers (Jones and Gosling, 1994). For example, in Cercis canadensis (Horovitz et al., 1975), the response of the an undeveloped (linear) embryo to chilling temperatures can induce germination (horovitz, 1975).

**Workshop**

Primary combinational dormancy

Combinational dormancy combines two or more kinds of dormancy, such as morphophysiological dormancy, where there is an underdeveloped embryo and physiological dormancy (Ilex) or exo-endodormancy that combines seedcoat dormancy and endogenous physiological dormancy (Cercis canadensis L.). To induce germination, all blocking conditions must be eliminated in proper sequence.

**Morphophysiological dormancy**. The most common form of combinational dormancy is morphophysiological dormancy. Currently, eight different types of morphophysiological dormancy are recognized (Baskin and Baskin, 1998; Nikolaeva, 1977). Those with horticultural interest include simple and epicotyl types.

Seeds with simple morphophysiological dormancy usually require warm (>15 °C) followed by cold (1 to 10 °C) conditions during which time the embryo develops during the warm temperature cycle and then breaks physiological dormancy during the chilling cycle. Various temperate zone herbaceous and woody plants fall into this category, including windflower (Anemone), twinleaf (Jeffersonia), ash (Fraxinus), yew (Taxus), and holly (Ilex) (Nikolaeva, 1977). In nature, these seeds are usually shed from the plant with coverings, but then require 1 to 3 months of subsequent chilling to enable the epicotyl to grow and prolonged survival of weed seeds in soil in the seasonal rhythms (dormancy cycling) and prolonged duration of dormancy periods when environmental conditions are not favorable for germination (Bewley and Black, 1994; Crocker, 1916; Karssen et al., 1983; Khan, 1981). These conditions can include unfavorable temperature, prolonged light or darkness, water stress, and anoxia. These conditions are particularly involved in the seasonal rhythms (dormancy cycling) and prolonged survival of weed seeds in soil (Baskin and Baskin, 1998; Egley, 1995).

Chilling temperatures can induce secondary dormancy in nondormant seeds. Coreopsis lanceolata L. seeds were relieved of nondeep physiological dormancy by dry storage for 6 to 18 months. The stored seeds germinated at high percentages at 15 and 25 °C, but entered secondary dormancy if held at 5 °C (Banovetz and Scheiner, 1994).

In some cases, seeds that did not require chilling stratification to satisfy primary dormancy may require it for release from secondary dormancy. For example, Nemophila insigne Doux. Ex Bent. seeds require darkness to germinate. If these seeds are exposed to light for a period of time, they enter secondary dormancy and will not germinate in the dark without a chilling treatment (Chen, 1968).
A high-temperature environment for germination can induce a common form of secondary dormancy termed thermodynamic. Thermodynamic dormancy can develop in species such as apple (Malus), lettuce (Lactuca), celery (Apium), Schizanthus, and pansy (Viola) if the germination temperature is too high (>25 °C). This phenomenon should not be confused with the thermal inhibition most seeds experience when the temperature exceeds the maximum temperature for germination. Seeds experiencing thermodynamic dormancy will not germinate when the temperature returns to near optimum temperatures, while thermally inhibited seeds will germinate when temperatures are lowered.

Seeds of some species of ash (Fraxinus) display morphophysiological dormancy that requires extended time (10 to 18 weeks) of warm stratification followed by additional time at 5 °C (=12 weeks) to relieve primary dormancy (Young and Young, 1992). Stratified seeds showed considerable secondary dormancy when germinated at constant 20 °C or alternating 20/30 °C (Piotto, 1994). Interestingly, there was no secondary dormancy seen in a widely fluctuating 25/5 °C germination environment. This and other studies suggest that caution should be taken when interpreting laboratory experiments where germination temperatures are held constant or day/night temperature fluctuations are minimized compared to the outside environment (Baskin and Baskin, 1998; Hilhorst, 1998).

Apple seeds require chilling stratification to relieve primary endogenous dormancy. Following release from dormancy, they are sensitive to induction into secondary dormancy at germination temperatures above 30 °C (Visser, 1954). Ozga and Dennis (1991) determined that abscisic acid content was not well correlated with induction of secondary dormancy. Hillhorst (1998) presents a convincing case for considering temperature-associated changes in membranes being responsible for release from dormancy, especially in seeds displaying secondary dormancy. Membranes adjust to varying temperature to maintain their fluidity, which directly impacts integral membrane proteins. These changes in the membrane may be related to release from primary dormancy or induction into secondary dormancy.

**Germination models**

A number of attempts have been made to develop mathematical models that predict seedling emergence from dormant seeds (Bradford 1996; Christensen et al., 1996; Franco, 1998; Kebab and Murdoch, 1999; Pritchard et al., 1996; Seeley and Damavandy, 1995). Some models that consider only temperature effects on dormancy release have been remarkably effective. For example, Bauwmeester and Karsen (1992) were able to predict seedling emergence in the field for a number of weed species using a “germination temperature window” based on the previous exposure of these seeds to various temperatures. Although temperature is implicated in most aspects of seed dormancy release, it is apparent that additional features such as population effects, seed moisture levels, and time must be considered when developing an effective model for dormancy release under field conditions (Hilhorst, 1998).

Genetic and environmental (primarily temperature factors) affect seed dormancy release. The genetic component can influence entire populations of seeds or individual seeds within a seed lot. An example where entire populations of seeds show different depths of dormancy is illustrated in Prunus serotina J.G. Ehr. seeds collected from different climatic zones (Farmer and Barnett, 1972). Seeds from ecotypes collected from higher altitudes required longer periods of chilling stratification to relieve dormancy and were slower to germinate at permissive germination temperatures following stratification compared to seeds collected from lower altitudes.

Genotype differences need to be considered when applying models that may have been generated using only one ecotype. Perez (1997) compared stratification requirements between low bud chilling peach accessions from subtropical regions with those of high-chilling accessions. Low-chilling genotypes showed dormancy release at temperatures as high as 14 °C, whereas this temperature had no impact on germination in high-chilling genotypes.

There are also differences in the depth of dormancy observed for seeds within a seed lot. The complexity of this genetic component becomes apparent when considering the correlation between seed-chilling requirements and bud-chilling requirements of plants (Powell, 1987). In studies with almond (Prunus dulcis L.), a high quantitative correlation was observed between the mean time for bud and seed dormancy release in seedling populations and the mean for both the seed and pollen parents (Kester, 1969). However, there was a low correlation between the time required to release dormancy in each individual seed and the subsequent chilling requirements for buds of the new plant developing from that embryo (Kester et al., 1977). This difference suggests that dormancy involves both a genetic component within the embryo and a maternal component from the seed parent. As a result, a great deal of variability can exist in the time to dormancy release in individual seeds within a given seed lot and between different seed lots of the same species collected in different years and different locations.

This maternal vs. paternal inheritance factor can be illustrated in reciprocal crosses of petunia (Girard, 1990). Seed coverings (maternal tissue) have an important influence on dormancy release (Hartmann et al., 1992) and these tissues are ostensibly maternal tissue. In petunia (Petunia x hybrida Hort. Vilm.-Andr.), the requirement for light was maternally inherited, while endogenous dormancy within the embryo was under paternal control.

Whether controlled by environmental factors during development or genetic factors within the embryo or seed coverings, the time required for dormancy release in individual seeds within a given seed lot is about normally distributed (Fig. 2). This suggests that release from seed dormancy could be described using a population-based thermal time model similar to well-characterized thermal time models for germination in nondormant seeds (Bradford, 1996). Accordingly, Pritchard et al. (1996) used thermal time to describe dormancy loss in horsechestnut (Aesculus hippocastanum L.) seeds. A negative linear relationship was observed between dormancy release with chilling over a range of stratification temperatures.

A second factor affecting predictive models for seed dormancy release is the interaction between temperature and seed moisture content. Chilling stratification is not effective unless seeds are hydrated. In nature, the degree of seed hydration varies depending on the environment. Therefore, there is a critical moisture content below which seeds would not be positively affected by chilling for dormancy release. In several conifer species, the critical moisture content appears to be ~25% moisture on a fresh weight basis (Gosling and Rigg, 1990). About 33% seed moisture allows dormancy release to proceed without

![Fig. 2. Percentage of peach seeds that germinate at 20 °C as a function of time under chilling stratification at 4 °C. (redrawn from data in Seeley and Damavandy, 1985).](image-url)
allowing germination during prolonged storage (Jones and Gosling, 1994). Downie et al. (1998) also observed that dormancy release in spruce [Picea glauca (Moench.) Voss.] seeds was achieved at a moisture content starting at \( \approx 25\% \). At this moisture content, seeds were at the boundary between water binding regions 3 and 4 as determined by moisture sorption isotherms (Vertucci and Farrant, 1995). In this condition, cellular components are hydrated, but not sufficiently to support turgor-driven cell expansion. Based on cellular properties in these moisture ranges, they suggested that protein hydration and possibly synthesis was required for release from dormancy in seeds requiring chilling stratification (Downie et al., 1998). Interestingly, freshly harvested seeds that require after-ripening (nondeep physiological dormancy) are released from dormancy in water binding region 2 (for example, wild oats, Foley, 1994). This example indicates the internal processes responsible for dormancy release are probably different for seeds with endogenous physiological dormancy that experience dormancy loss due to dry storage compared to those requiring chilling stratification.

Finally, the most interesting problem with models that attempt to describe the time required for dormancy release at a given temperature is conditional dormancy. This problem goes directly to the question "what constitutes dormancy release?". Dormancy cycling, as observed in many species, is a function of conditional dormancy (Baskin and Baskin, 1998). It is a transitional state between the dormant and nondormant seed condition. This transition can be observed by evaluating germination over a range of germination temperatures. Nondormant seeds germinate rapidly over a wide range of temperatures, while conditionally dormant seeds germinate only under the permissive (optimum) temperature (Vegis, 1964). In most laboratory studies, seeds are exposed to a dormancy releasing treatment (e.g., chilling temperatures) and the time to dormancy loss as indicated by germination at an optimal temperature is recorded. In reality, this measurement is an indication of the time required to move from a dormant to conditionally dormant state. The time required to achieve a fully nondormant state would be indicated by the ability to germinate over a range of temperatures (Baskin and Baskin, 1998). In nature, seeds can go through years of dormancy cycling, each cycle containing periods of dormancy, nondormancy and conditional dormancy. When other factors (such as light) are not limiting, germination occurs only when the degree of dormancy release corresponds to an appropriate germination temperature range. Therefore, models that attempt to predict germination in dormant seeds in nature must consider the impact of temperature on the degree of dormancy release (conditional dormancy) and adjust the model to account for corresponding permissive germination temperatures.

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