Bud endodormancy in deciduous fruit trees: advances and prospects

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Abstract
Bud endodormancy is a complex physiological process that is indispensable for the survival, growth, and development of deciduous perennial plants. The timely release of endodormancy is essential for flowering and fruit production of deciduous fruit trees. A better understanding of the mechanism of endodormancy will be of great help in the artificial regulation of endodormancy to cope with climate change and in creating new cultivars with different chilling requirements. Studies in poplar have clarified the mechanism of vegetative bud endodormancy, but the endodormancy of floral buds in fruit trees needs further study. In this review, we focus on the molecular regulation of endodormancy induction, maintenance and release in floral buds of deciduous fruit trees. We also describe recent advances in quantitative trait loci analysis of chilling requirements in fruit trees. We discuss phytohormones, epigenetic regulation, and the detailed molecular network controlling endodormancy, centered on SHORT VEGETATIVE PHASE (SVP) and Dormancy-associated MADS-box (DAM) genes during endodormancy maintenance and release. Combining previous studies and our observations, we propose a regulatory model for bud endodormancy and offer some perspectives for the future.

Introduction
Dormancy refers to a temporary suspension of visible growth of any plant structure containing a meristem and is a biological characteristic of higher plants adapted to seasonal environmental changes through long-term natural selection. Bud dormancy is an important physiological process that helps plants survive harsh winter weather and determines growth resumption and flowering in the following spring. Bud dormancy is generally divided into three types: para-dormancy, endodormancy, and ecodormancy according to various factors that cause dormancy. This classification has been widely accepted by researchers in the study of bud dormancy. In this review, we mainly discuss endodormancy.

In autumn, with decreased temperature and/or day length, buds (both terminal buds and axillary buds) are induced to enter endodormancy. In this stage, the buds cannot sprout, even under favorable conditions. After the fulfillment of a certain period of chilling accumulation (termed chilling requirement, CR), endodormancy is released, and buds acquire the potential capability to resume growth. Because the release of endodormancy generally occurs at the depth of winter, buds cannot break immediately because of unfavorable conditions and are forced into ecodormancy (Fig. 1). With increasing temperature, bud growth resumes in the spring. According to Howe et al. and Singh et al., endodormancy is further divided into three stages: induction (establishment), maintenance, and release (Fig. 1). Endodormancy maintenance refers to the process from the beginning of chilling accumulation until endodormancy release. In this stage, the buds could not burst unless chilling accumulation was satisfied. With prolonged chilling accumulation, the factors required for endodormancy maintenance are eliminated, and endodormancy is released.

During the past decade, significant progress has been made in studying the mechanism of growth cessation and
bud endodormancy in various species, especially in the model plant poplar (Populus spp.). Growth cessation is a necessary process prior to bud endodormancy. With the shortened day length in autumn, poplar ceases growth and forms apical buds containing leaf primordia and shoot apical meristems. Böhlenius et al. found that two genes, CONSTANS (CO) and FLOWERING LOCUS T (FT), which normally regulate flowering time, also regulate the induction of growth cessation and bud set under short days. FT2 is expressed in the leaves, and the FT2 protein is translocated to the apex, where it promotes growth; downregulation of FT2 in leaves results in reduced levels of the FT2 protein in the apex. Genes related to the circadian clock also participate in growth cessation. Ding et al. showed that the circadian clock gene GI (GIGANTEA) directly regulates the expression of FT2 independent of CO, thereby regulating short-day-induced growth cessation and apical bud formation in poplar. Downregulation of LHY and TOC1 delays growth cessation and bud set, and LHY2 is necessary and sufficient to cause FT2 downregulation in a night-length-dependent manner. FT2 downregulation results in inhibition of the expression of the downstream components LAP1 (the homologous gene of APETALA1, a characteristic gene of Arabidopsis floral meristem) and AIL1 (the homologous gene of AINTEGUMENTA in Arabidopsis), which in turn eventually leads to growth cessation and bud formation. Recent studies have shown that in addition to the CO/FT pathway, BRC1 (homologous to Arabidopsis gene BRANCHED 1) can form a feedback loop controlling seasonal growth in response to short days (Fig. 2). In addition, the content of the plant hormone abscisic acid (ABA) increases under short days, promoting ABI3 expression and bud development.

After growth cessation, buds enter endodormancy in response to the autumn environment. Abscisic acid (ABA) is centrally important in establishing bud endodormancy. Studies on poplar indicate a clear network of endodormancy regulation. ABA mediates callose deposition and plasmodesmata closure and blocks the transport of growth-promoting substances such as FT1 to shoot meristems, thus leading the buds to endodormancy; however, ABA does not affect growth cessation. This result further proves that growth cessation and bud endodormancy are two different processes. After prolonged chilling accumulation, GA biosynthesis genes (e.g., GA20ox8 and GA3ox1) are upregulated, and callose is degraded, allowing FT to be transported to shoot meristems, where it induces endodormancy release and bud break. In poplar, the MADS-box gene SVL (SHORT VEGETATIVE PHASE LIKE), which is induced by ABA, is a crucial transcription factor promoting bud endodormancy. SVL mediates short-photoperiod- and low-temperature-induced terminal bud endodormancy and directly promotes the expression of the ABA synthesis gene NCED3, forming a positive feedback regulation with ABA signaling. In addition, SVL promotes the expression of CALSI, a callose synthetase that promotes callose deposition, resulting in the closure of
plasmodesmata, thus blocking intercellular communication. At the same time, SVL also inhibits the expression of \( \text{GA20ox1} \) and \( \text{GA20ox2} \) and promotes the expression of \( \text{GA20ox8} \), thus reducing the content of gibberellin (GA) and deepening endodormancy \(^{19,20}\). Therefore, SVP/SVL can integrate the metabolism of ABA and GA, regulating the maintenance of endodormancy. Reducing the GA content by overexpressing the GA catabolism gene \( \text{GA2ox8} \) also promotes apical bud formation and delays endodormancy release\(^{19,21}\). EARLY BUD BREAK 1 (EBB1), which suppresses SVL expression, is positively regulated by low temperature, leading to the upregulation of \( \text{EBB3} \). EBB3 directly activates \( \text{CYCD3.1} \), promoting cell division and bud break\(^{22}\). In addition to transcriptional regulation, epigenetic regulation, including DNA methylation, is also reported to be involved in growth cessation and bud endodormancy processes in poplar. The DNA demethylase \( \text{DEIMITER-LIKE} \) is chilling responsive and mediates poplar bud formation and bud break\(^{23,24}\).

Although these findings in the model plant poplar help us greatly understand the biology of the bud dormancy cycle, most of the studies have focused on vegetative buds. However, in the case of fruit trees, the dormancy of floral buds should be given more attention because it is a critical component in fruit production. In this review, we focus on bud endodormancy of deciduous fruit trees, summarizing the recent progress on endodormancy regulation and discussing the scientific questions still to be addressed to further understand the mechanism of bud endodormancy.

**Endodormancy induction in deciduous fruit trees**

In many deciduous fruit trees, such as pear and apple, growth cessation and flower bud differentiation often occur in early summer (May–June in China)\(^{25}\) when the days become longer and the temperature higher\(^{25}\), while the establishment of endodormancy happens in late autumn when the days become shorter and the temperature lower. Hence, the growth cessation of shoots in some deciduous fruit trees is not strictly caused by short-day conditions. Heide and Prestrud\(^{26}\) showed that growth arrest and bud formation in pear and apple could be induced by artificial low temperatures rather than short days. However, under natural conditions, growth cessation occurs in summer at relatively high temperatures. Therefore, the factors regulating the growth cessation of deciduous fruit trees under natural conditions need further study, as Cooke et al.\(^{27}\) also commented.

Induction of bud endodormancy often occurs in autumn. In peach, bud endodormancy is induced by both short-day and low-temperature conditions\(^{28}\), while in grape, bud endodormancy was shown to be induced by short-day length or low temperature in different grape genotypes\(^{29}\). Heide and Prestrud\(^{26}\) concluded that endodormancy of European pear and apple could be induced by low temperature (\(<12{\degree}\text{C}\) rather than a short-day length under artificial conditions. However, growth cessation and bud endodormancy were not clearly distinguished in their study. It is difficult to determine whether plants in their study were in the endodormancy stage or had just ceased growth, because even if the trees ceased growth and formed apical buds, the apical buds might sprout when early defoliation occurs in summer\(^{30}\).

According to our observations, under natural conditions in Hangzhou, China (\(30°\ 06′\ N; 119°\ 55′\ E\)), the bud-break percentage of ‘Cuiguan’ pear (\(\text{Pyrus pyrifolia}\)) under forcing conditions decreased from September and reached its lowest in late October. During this period, the daily mean temperature decreased from c. \(25°\text{C}\) to c. \(14°\text{C}\), much higher than the temperature used in Heide and Prestrud’s experiments\(^{26}\) (Fig. 3). A similar phenomenon

![Fig. 3 Daily temperature and bud break percentage of ‘Cuiguan’ pear in Hangzhou](image)

**Fig. 3** Daily temperature and bud break percentage of ‘Cuiguan’ pear in Hangzhou. Daily mean temperature in Hangzhou (a) and bud-break percentage indicating the dormancy status of ‘Cuiguan’ pear (b) in autumn and winter of 2019. The endodormancy stage is indicated between the vertical dashed red lines. Horizontal dashed red lines indicate temperatures as shown (a) and 50% bud breakage (b). Branches were sampled at the time points shown and were kept in floral foam in forcing conditions for 21 days for measuring the bud-break percentage. When the trees entered endodormancy (start of the period with the lowest bud-break percentage), the daily mean temperature was still ~\(14°\text{C}\), suggesting that the critical temperature to induce endodormancy of ‘Cuiguan’ pear is possibly not lower than \(14°\text{C}\). When dormancy was released, the temperature was just approximately \(7°\text{C}\), suggesting that the \(7°\text{C}\) model might not be suitable for calculating the chilling requirement of ‘Cuiguan’ pear in the Hangzhou area.
was observed in another pear cultivar, ‘Suli’ (P. pyrifolia White Pear Group), in Dangshan County, China (34° 28’ N; 116° 29’ E)\(^3\). It is generally believed that there is a critical day length or temperature for each species or even ecotype to induce bud endodormancy. However, for most fruit trees, the critical temperature/day length for endodormancy induction still needs further study. In addition to a low temperature and short photoperiod changes in other environmental cues (e.g., air humidity or soil moisture content) might also be included as putative endodormancy inducers. Therefore, the environmental signals for bud endodormancy induction are still obscure in most fruit tree species and may differ with the local environment.

In addition to environmental inducers, exogenous ABA treatments have been found to promote the establishment of bud endodormancy in deciduous fruit trees\(^3\)\(^1\)\(^–\)\(^3\). During the endodormancy induction stage of grape buds, endogenous ABA increased\(^3\).^4 In grape, overexpression of \(CYP707A4\), a gene encoding an ABA 8′-hydroxylase caused a lower ABA content and increased lateral bud out-growth and shoot length\(^3\)\(^5\), suggesting that ABA might be the main regulator of growth cessation in grape. However, the effects of ABA on endodormancy induction have not been determined\(^3\)\(^5\). In our observations over several years, the ABA content in pear buds peaked during the endodormancy maintenance stage rather than the induction stage\(^3\)\(^1\),\(^3\)\(^3\). Therefore, whether bud endodormancy of deciduous fruit trees is induced by endogenous ABA under natural conditions remains unclear because of the lack of a genetic analysis approach.

**Endodormancy maintenance and release in deciduous fruit trees**

The maintenance and release of bud endodormancy are consecutive processes. After the induction of endodormancy, with increasing chilling hours, the factors that maintain bud endodormancy are gradually inhibited, and bud endodormancy is eventually released. Whether the endodormancy of buds can be released on time to enable commercial fruit production is a matter directly related to bud burst and flowering time and therefore affects yield and fruit quality.

**Chilling requirement**

The chilling requirement (CR) is directly related to endodormancy maintenance and release and closely related to flowering time. Therefore, an accurate evaluation of the CR is important, and various CR models have been developed. Weinberger\(^3\)\(^6\) demonstrated that temperatures below 45 °F (\(-7.2 °C\)) are effective for breaking peach endodormancy. Although a 7.2 °C model has been used for many years, there are some problems because different fruit trees may have their own specific low temperature. Thus, Richardson et al.\(^3\)\(^7\) proposed a Utah model, demonstrating that different extents of chilling have different effects on endodormancy. The 7.2 °C model and Utah model are widely used at present, while the 0–7.2 °C model, 0–14 °C model, positive Utah model, and modified Utah model, which are all modified from the 7.2 °C and Utah models, are suitable for special areas or climates\(^3\)\(^8\). The CR differs among plant species and cultivars and is also affected by climate and other environmental factors. For example, ‘Suli’ pear floral buds need approximately 800–1000 hours of chilling\(^3\)\(^9\),\(^4\)\(^0\), while ‘Cuiguan’ pear floral buds only need approximately 200–400 chilling hours to break endodormancy\(^3\)\(^9\). However, it is often difficult to determine the start time of chilling accumulation, resulting in inaccurate CR determination. In our recent observations, endodormancy of ‘Cuiguan’ pear was released when the daily mean temperature was above 7.2 °C in Hangzhou, China (Fig. 3), while the Utah model indicated that the CR was 113 chilling units (CU), much lower than that in previous studies (c. 300 CU)\(^3\)\(^9\),\(^4\)\(^1\). This phenomenon suggests that CR models are not universal for different conditions, and CR data acquired under natural conditions might still need to be modified by applying artificial chilling accumulation at a fixed temperature. As 4–5 °C is efficient for the endodormancy release of many trees, researchers often use that temperature to break endodormancy\(^3\)\(^8\),\(^4\)\(^0\),\(^4\)\(^2\).

Thus, we can calculate the CRs more accurately by using artificial chilling accumulation.

Obviously, CR is a typical quantitative trait and therefore amenable to study by QTL analysis. Fan et al.\(^4\)\(^3\) located CR-related QTLs in peach on linkage groups G1, G4, G5, G6, G7, and G8 using a population of 378 F2 individuals. Interestingly, the major QTLs in G1 were also associated with flowering time. Rowland et al.\(^4\)\(^4\) identified two CR-associated QTLs in blueberry from a genetic linkage map using simple repeat sequence (SSR) markers. Castède et al.\(^4\)\(^5\) identified dormancy- and flowering-related QTLs and screened 57 out of 79 candidate genes for colocalization with these QTLs. Among them, \(LHY\), which was reported to regulate growth cessation and bud break in poplar, was located on LG2 in the CR-related QTL but not in the QTL related to flowering time and heat requirements. \(HYL1\) of sweet cherry was located on LG1 together with \(DAM5\) and \(DAM6\) in the QTL related to flowering time, which was consistent with findings in peach, showing that this genomic region is conserved between sweet cherry and peach\(^4\)\(^5\). Recently, QTLs related to the downregulation of \(PmI\)\(DAM6\), endodormancy release, and bud burst in Japanese apricot, were identified, further demonstrating that the downregulation of \(DAM\) genes is involved in bud endodormancy release and bud burst\(^4\)\(^6\). Gabay et al.\(^4\)\(^7\),\(^4\)\(^8\) identified QTLs associated with bud burst in European pear and found collinearity of the
genetic regions controlling the CR between apple and pear. Recently, these researchers further carried out RNA-seq and metabolic profiling, combined with QTL analysis, and identified α-linolenic acid as an important metabolite in relation to the CR. Although much progress has been made in CR research in different species, we are not yet able to fully explain the mechanistic basis of the CR.

**Roles of phytohormones**

Phytohormones play an important role in the dormancy–growth cycle. Among them, ABA and GA are the two most important hormones that antagonistically regulate bud endodormancy induction, maintenance, and release. A high level of endogenous ABA is the primary factor in maintaining bud endodormancy, while GA is responsible for endodormancy release. Treatment with exogenous GA caused delayed endodormancy release, while ABA plays a crucial role in endodormancy maintenance, which is consistent with studies on poplar. Over-expression of the ABA-responsive transcription factor PpyABF3 caused growth inhibition in pear calli with downregulation of CYCLIN-D and EXPANSIN A1, indicating that ABA promotes endodormancy maintenance, possibly by inhibiting cell division and cell expansion. Recent studies on pear showed that the expression of PpyCYP707A3, a gene related to ABA catabolism, increased continuously and linearly during the chilling accumulation process. At the same time, the ABA content decreased, and bud break increased, which is similar to findings in peach. In addition, overexpression of VvCYP707A4 reduced the ABA content in grape buds, promoting endodormancy release and leading to enhanced regrowth.

In contrast to ABA, GA may be related to endodormancy release. The GA content in peach buds increased with accumulated chilling hours. During the process of endodormancy release, the expression of gibberellin-related genes, such as GA3OX, GA20OX, and GASA, was significantly upregulated, and the GA content in buds increased significantly in grape and pear, which is similar to observations in poplar. Exogenous GA treatments were effective in promoting endodormancy release only after a period of chilling accumulation in Japanese apricot and pear. However, the mechanism of chilling-induced GA accumulation remains obscure, and how GAs function in endodormancy release in deciduous fruit trees is still not clear.

Moreover, the regulation of ABA and GA in the dormancy cycle usually occurs through their crosstalk. Our recent study showed that pear GAST1, which promotes GA biosynthesis, is inhibited by ABA. When ABA is degraded with the upregulation of CYP707A during prolonged chilling accumulation, GAST1 is released from ABA inhibition, resulting in an increase in the GA content. ABA-responsive ABF3 in pear also induced the expression of GA2OX1, which might lead to a lower GA content. Thus, ABA seems to promote GA catabolism and inhibit GA biosynthesis, thus maintaining bud endodormancy in pear, which is similar to the results in poplar. However, the transcription factors at the intersection of ABA and GA during bud endodormancy are still obscure.

In addition to GA and ABA, other phytohormones are also involved in the endodormancy process. Studies on tea plants showed that the IAA content in a terminal and lateral buds increased gradually during endodormancy release, while studies on the adventitious buds of leafy spurge showed that the IAA content was higher in the paradormancy and endodormancy stages and lower in the ecodormancy stage. A study in grape showed that ethylene might also participate in the process of bud endodormancy release. However, there is still a lack of genetic approaches that would provide solid evidence to establish the regulatory roles of these phytohormones in bud endodormancy.

**Molecular network of DAM/SVP-centered bud endodormancy regulation**

Recent studies have shown that ABA regulates endodormancy, probably through MADS-box transcription factors that belong to the AGL24/SVP subfamily. In non-Rosaceae deciduous fruit trees, these MADS-box proteins are usually referred to as SVPs, which are clustered in the same evolutionary branch as Arabidopsis SVPs (Fig. 4). The functions of these proteins during the endodormancy process have been verified using transgenic plants in several species, such as poplar and kiwifruit. In poplar, SVP-like (SVL) acts as a central transcription factor integrating ABA signaling, ABA synthesis, GA synthesis, GA catabolism, and cell division, hence maintaining bud endodormancy. In rosaceous plants, these dormancy-associated MADS-box proteins are named DAMs and are clustered separately from other SVPs. Six DAM genes, which are tandemly located at the end of chromosome 1 of *Prunus* species, were first identified in the peach *evergreen* mutant. Ectopic expression of *PmDAM6* in poplar promoted the formation of dormant terminal buds and inhibited growth. Zhao et al. found protein–protein interactions among DAM1–DAM6 proteins in *Prunus mume* and different expression patterns during bud endodormancy,
indicating that DAMs might form different dimers during endodormancy. Wu et al. identified DAM and SVP genes in apple and found that overexpression of MdDAMb and MdSVPa caused delayed bud break. Ubi et al. and Saito et al. successively cloned three DAM genes (PpyMADS13-1/-2/-3) from Japanese pear ‘Kosui’ and showed that they are located on chromosomes 8 and 15. Afterwards, Liu et al. and Niu et al. also identified the same DAM genes from Chinese pear ‘Suli’ as those in Japanese pear and numbered them. Tuan et al. named PpyMADS13-I PpyDAM1, corresponding to Niu’s DAM3. To better understand the evolution of SVP and DAM genes among different species, we unified the names of pear DAM genes according to the order of the chromosomes based on the new pear genome (unpublished data) and constructed a phylogenetic tree for SVP and DAM proteins in deciduous tree species (Fig. 4). DAM genes are SVP-like genes regulating the endodormancy process in Rosaceae, and they have not been found in grape, poplar, or Arabidopsis.

DAM genes are well studied in pear, peach, apple, and Japanese apricot. A molecular network associated with DAMs has already been demonstrated. Studies on pears have shown that low-temperature- and ABA-induced C-repeat binding factors (CBFs) directly bind to the promoters of DAM genes and activate their expression, promoting pear bud endodormancy while ABA-induced HD-ZIP protein HB22 also directly binds to the promoter of DAM1 independently of CBF. In Japanese apricot, CBFs not only bind to the promoter of DAM6 but also form dimers with the DAM6 protein. In the ABA signaling cascade, ABA-responsive AREB1 inhibits the expression of DAM genes, while ABF3, an ABA-responsive transcription factor, induces DAM3 expression, and ABF2 (a homologue to AREB1) disrupts activation by interacting with ABF3. These interactions
indicate the fine-tuned control of DAM gene expression, enabling plants to release endodormancy on time. In peach, the transcription factor TCP20 interacts with ABF2 and binds to the DAM5/DAM6 promoter, inhibiting its expression. In addition to transcription factors, miRNA6390 might be involved in the post-transcriptional regulation of DAM genes, promoting the degradation of DAM transcripts. At the same time, DAM promotes the expression of NCED, which encodes a key enzyme in ABA synthesis; elevated NCED expression maintains a high ABA concentration in buds. Together, these roles of DAM form a negative feedback mechanism regulating pear bud endodormancy. Furthermore, the expression of DAMs can negatively regulate the expression of FT2, a gene that promotes growth and flowering, during endodormancy. In our recent study, overexpression of DAM3 in pear calli resulted in downregulation of Cyclin-D and EXPA1, suggesting the potential regulation by DAMs of cell division and expansion, which is similar to the role of SVL in poplar, indirectly inhibiting Cyclin-D3 via EBB3. However, other target genes of DAMs in deciduous fruit trees remain obscure.

On the basis of previous studies in deciduous fruit trees and poplar, we summarized a DAM/SVP-centered endodormancy regulation network (Fig. 5). However, this molecular network cannot fully explain the mechanism of endodormancy release induced by chilling accumulation and the differences in the CR among different cultivars.

### Epigenetic regulation of bud endodormancy maintenance and release

Epigenetic regulation plays an important role in plant growth and development and in the response to environmental stresses. Epigenetic modification refers to reversible but heritable changes in gene expression and gene function without changes in the nucleotide sequence of the gene via mechanisms, including histone modification, DNA methylation, long noncoding RNA (lncRNA), and chromatin remodeling. An increasing number of studies have shown that epigenetic modifications are deeply involved in the processes of both vernalization and seed dormancy, which are somewhat similar to bud endodormancy in the requirement for a certain period of chilling accumulation to finish vernalization or release seed dormancy. FLOWERING LOCUS C (FLC), which belongs to the MADS-box family, is an important gene regulating vernalization in Arabidopsis and has been proven to be regulated by epigenetic modification. Prolonged chilling during vernalization promotes the transcription of an antisense lncRNA named COOLAIR in the FLC locus and further alters histone modifications at the FLC gene locus through the polycomb-silencing complex pathway, resulting in decreased H3K4me3 levels and increased H3K27me3 levels. These changes in histone methylation downregulate the expression of FLC, and the degree of downregulation was positively correlated with the duration of chilling accumulation. Epigenetic regulation plays an important role in plant growth and development and in the response to environmental stresses. Epigenetic modification refers to reversible but heritable changes in gene expression and gene function without changes in the nucleotide sequence of the gene via mechanisms, including histone modification, DNA methylation, long noncoding RNA (lncRNA), and chromatin remodeling. An increasing number of studies have shown that epigenetic modifications are deeply involved in the processes of both vernalization and seed dormancy, which are somewhat similar to bud endodormancy in the requirement for a certain period of chilling accumulation to finish vernalization or release seed dormancy. FLOWERING LOCUS C (FLC), which belongs to the MADS-box family, is an important gene regulating vernalization in Arabidopsis and has been proven to be regulated by epigenetic modification. Prolonged chilling during vernalization promotes the transcription of an antisense lncRNA named COOLAIR in the FLC locus and further alters histone modifications at the FLC gene locus through the polycomb-silencing complex pathway, resulting in decreased H3K4me3 levels and increased H3K27me3 levels. These changes in histone methylation downregulate the expression of FLC, and the degree of downregulation was positively correlated with the duration of chilling accumulation.
modifications, including histone methylation and histone acetylation of genes involved in ABA and GA metabolism and signal transduction (e.g., CYP707A, GA2ox), also act in regulating seed dormancy release and germination\textsuperscript{84–88}. In addition, DELAY OF GERMINATION1 (DOG1), involved in seed dormancy in ABA signaling\textsuperscript{89}, is also regulated by H3K27me3 and H3K4me3 histone modifications during the seed dormancy cycle\textsuperscript{90}. Therefore, endogenous hormone balance is critical for seed dormancy maintenance and release, and epigenetic regulation might activate and silence hormone-related genes.

Because the model of transcriptional regulation does not fully explain the mechanism of bud endodormancy release caused by subsequent chilling accumulation, some researchers have proposed that DAM genes are further regulated by epigenetic modifications similar to FLC during bud endodormancy. Santamaria et al.\textsuperscript{91} found differential expression of the histone modification-related genes HUB2 and GCNSL in dormant and germinating buds in chestnut. Research in leafy spurge has shown that there was also a decrease in H3K4me3 and an increase in H3K27me3 in the DAMI promoter region during endodormancy release\textsuperscript{92}, which is similar to the epigenetic modification of FLC during vernalization. Changes in H3K4me3, H3K27me3, and H3ac modifications of the promoter, coding region, and the second intron region of DAM6 in two peach cultivars, were closely related to their specific chilling requirements and the date of endodormancy release\textsuperscript{93}. During natural endodormancy, the reduction in H3K4me3 and H3ac modification of AcSVP2, which can delay bud break in kiwifruit, resulted in downregulation of AcSVP2, causing endodormancy release\textsuperscript{94}. Recently, combined analysis of ChIP-seq with RNA-seq in sweet cherry revealed that the enrichment of H3K4me3 and H3K27me3 at target genes such as Pav-DAM5 was positively and negatively correlated, respectively, with the expression of these genes during endodormancy\textsuperscript{95}. In peach, downregulation of DAM genes was closely related to the increased level of H3K4me3 and CHH DNA methylation during chilling accumulation and during warmer conditions after chilling exposure, while noncoding RNAs also participated in regulating epigenetic modifications\textsuperscript{76,96}. However, in poplar, SVL seems not to be a target for H3K27me3 histone modification, but the expression of the downstream gene EBB3 showed a negative correlation with H3K27me3 deposition after chilling exposure\textsuperscript{91,76}. These studies indicate that DAM/SVP genes in deciduous fruit trees are potential target genes for epigenetic regulation during endodormancy maintenance and release, and that the changes in histone modification might be related to the chilling requirements of plants. Nevertheless, in the absence of mutants that alter chromatin regulation and affect endodormancy, most of these studies in deciduous fruit trees have yet to establish a causal link between chromatin regulation and endodormancy regulation.

Moreover, genes related to hormone metabolism might also be epigenetically regulated during bud endodormancy, similar to processes in seed dormancy. During seed dormancy and germination, CYP707A, which encodes an enzyme of ABA catabolism\textsuperscript{86}, is most likely a target gene of epigenetic regulation. In pear, CYP707A3 is upregulated with prolonged chilling accumulation during bud endodormancy\textsuperscript{31,40}, suggesting that CYP707A3 might similarly be a likely target of epigenetic accumulation during bud endodormancy processes. In addition to histone modification, DNA methylation is also involved in the regulation of bud endodormancy\textsuperscript{99–101}. Law and Sutte\textsuperscript{102} found that potato tubers showed high methylation levels during both endodormancy induction and release. Li et al.\textsuperscript{103} found that endodormant blueberry floral buds had higher methylation levels than those in the ecdormancy stage. Studies on apple showed that the changes in DNA methylation in buds during chilling accumulation and endodormancy release were significantly correlated with altered expression of some genes\textsuperscript{104}. DNA methylation and small interfering RNA (siRNA) might be involved in the regulation of MADS-box genes in sweet cherry during the dormancy transition\textsuperscript{100}. More recently, Rothkegel et al. demonstrated that DNA methylation might act as an early sensor of low temperature, leading to genotype-specific reprogramming and thus changing gene expression\textsuperscript{105}.

In summary, epigenetic modifications such as histone modifications and DNA methylation are involved in bud endodormancy regulation, but how these modifications regulate endodormancy remains largely unknown, even in the model plant poplar. How the key genes related to bud endodormancy, such as DAM genes, are regulated by epigenetic modification still needs further study.

**Conclusions and perspectives**

Floral bud endodormancy in deciduous fruit trees is an essential process that is complex and different from that in the model plant poplar. Nevertheless, the results of poplar endodormancy can provide a basis for research on fruit tree dormancy in the future. Low temperature and a short photoperiod are supposed to be the main environmental factors promoting bud endodormancy. In autumn,
environmental changes cause the accumulation of ABA, leading to the differential expression of transcription factors (e.g., ABF, CBF, and DAM/SVP). DAM/SVPs in turn regulate the contents of phytohormones (ABA and GA) by regulating the expression of structural genes (e.g., NCED, GA20OX, and GA20OX) and forming a feedback regulation loop. At the same time, long-term exposure to winter low temperatures might cause epigenetic changes, which in turn result in changes in gene expression (e.g., DAM and CYP707A4), thus altering the contents of ABA and GA, leading to endodormancy release. However, there are still some major issues related to bud dormancy in fruit trees that need to be addressed in the future:

1. In some deciduous fruit trees, growth cessation occurs from early summer when the day length and temperature increase, but induction of endodormancy occurs in autumn when it is still warm in some regions. Although artificial low temperature and/or short-day length induce bud endodormancy, factors causing growth cessation and inducing endodormancy of deciduous fruit trees under natural conditions remain unclear.

2. Although DAMs/SVPs in some fruit trees, such as peach and kiwifruit, have been shown to promote endodormancy induction and maintenance, more genetic evidence is needed to verify the role of DAMs in bud endodormancy in other fruit trees. In addition, how DAM/SVP functions as a growth inhibitor remains largely unknown but might be clarified in the future by studying the target genes of DAMs/SVPs.

3. While much evidence has shown that epigenetic regulation participates in the dormancy cycle, how these epigenetic modifications (e.g., histone modification and DNA methylation) are enriched or reversed during dormancy processes remains unknown. It would be interesting to uncover how epigenetic marks are recruited to specific genes (e.g., DAM and SVP) and how those modifications are affected by chilling accumulation during bud endodormancy processes.

4. Chilling requirements are a complex quantitative trait. Although recent QTL analyses have provided useful information, the detailed genetic basis of CR traits remains obscure. The mechanisms underlying the formation of CR traits in floral buds need further study using newer genetic approaches, such as gene editing, genome-wide association studies, and genome-wide environmental association studies, which entail years of accumulated experience and materials.

Hormonal and transcriptional regulation can partially explain the dormancy cycle, while revealing the various epigenetic regulation mechanisms might explain the different CRs among different cultivars and under different environmental conditions. More genome-wide techniques, such as ChIP-seq and genome-wide association studies, should be used to reveal the genetic basis of the CR trait, while genetic transformation will allow us to verify the functions of key regulators of endodormancy.

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Conflict of interest
The authors declare no competing interests.

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