Distinct Pathways of Regulation Seed Dormancy and Germination by Abscisic Acid

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Abstract

Abscisic Acid (ABA) is the phytohormone inducing seed dormancy, whereas seed germination occurs at low ABA concentration. Two pathways of their regulation are considered. One represents ABA reception and its signal transduction to ABA-responsive genes to induce dormancy genes, a modification of this way leading to germination in the absence of ABA. Another way suggests direct ABA inhibition of plasma-membrane H+-ATPase in dormant seeds that prevents germination and its activation in the absence of ABA, thus promoting seed germination.

Keywords: Seed dormancy; Germination; Transcription of dormancy genes; Activation of plasma membrane H+-ATPase.

Introduction

After the discovery of ABA receptors belonging to PYR/PYL/RCAR protein family [1,2], and mechanism of their functioning [3], the components of signal system transducing the ABA signal to nuclear ABA-sensitive genes were identified [4,5]. According to the modern point of view, this regulatory pathway is common for the induction of primary seed dormancy and its maintenance. However, dormancy release includes the ABA degradation [6], and non-dormant seeds contain low ABA concentrations, which do not prevent seed germination [7,8]. In the absence of ABA, this signal system is modified to make the germination possible [9,10]. Figure 1 schematically describes such model of ABA regulation established on Arabidopsis seeds. ABA signaling system in seeds consists of three components, namely protein phosphatases, protein kinases and transcription factor ABI5.

<table>
<thead>
<tr>
<th>In the presence of ABA</th>
<th>In the absence of ABA</th>
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<tr>
<td>1. ABA interacts with the receptor and the complex * receptor-ABA-protein phosphatase PP2C* is formed.</td>
<td>1. Protein phosphatase PP2C forms a complex with protein kinase SnRK2.</td>
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<td>2. Protein phosphatase PP2C is inhibited that results in the activation of protein kinase SnRK2.</td>
<td>2. SnRK2 is inactivated by dephosphorylation.</td>
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<td>3. Protein kinase phosphorylates transcription factor ABI5.</td>
<td>3. Phosphoprotein phosphatase PP6 dephosphorylates ABI5 transcription factor and inactivates it.</td>
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<td>4. Phosphorylated ABI5 interacts with ABA response elements in promoters of ABA-sensitive genes and activates their expressions.</td>
<td>4. ABI5 transcription factor is degraded in proteasomes.</td>
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<td>5. ABA-sensitive genes determine seed dormancy, i.e. inability to germinate.</td>
<td>5. Genes are expressed that determine non-dormant seed germination.</td>
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Figure 1: ABA and regulation of dormancy and germination via transcription.

Protein phosphatases PP2C, known also as ABI1 and ABI2 factors, negatively regulate ABA responses. Arabidopsis seeds have six genes of PP2Cs, the expression of which results in the production of these proteins taking part in the formation of the ternary complex with ABA and its receptor and in modulating the phosphorylation activity of protein kinases. Three protein kinases SnRK2 take part in ABA signaling. They mediate phosphorylation and activation of transcription factor ABI5- the step critical in the dormancy and inhibition of germination.

ABI5 protein always has a phosphorylation site to be activated by SnRK2 protein kinases. This ABA-responsive activation is accompanied by its stabilization during dormancy induction and...
maintenance. The participation of ABI5 permits the expression of ABA-responsive dormancy-specific genes. The degradation of ABI5 is necessary for the initiation of germination. It begins by ABI5 dephosphorylation by another family of serine/threonine protein phosphatases, namely FYPP/PP6 phosphoprotein phosphatases, followed by its breakdown through the 26S proteasome-dependent pathway. ABI5 dephosphorylation is an essential prerequisite for eliminating the inhibitory effect of ABA on germination. The abolition of ABI5 transcription factor causing subsequent changes in gene expression is a key event for seed germination.

The successful functioning of ABA reception and subsequent transduction of the signal to the corresponding nuclear genes determines the so-called ABA sensitivity; any disturbance of receptor--signaling system, for example in mutants, results in a loss of ABA sensitivity. The list of genes involved in dormancy regulation was presented by Graeber et al. [11]. In dormant Arabidopsis seeds gene expression is enriched for proteins involved in ABA synthesis, gibberellin catabolism, protein kinases, protein phosphatases, numerous transcription factors, and typical maturation proteins, namely small heat shock proteins and Late Embryogenesis Abundant (LEA) proteins [12,13]. Special attention was paid to DOG1 protein regulating seed dormancy induction and delaying germination by interfering with PP2C protein phosphatase activity [14]. The translation of proteins occurs slowly in dormant seeds.

The set of germination proteins radically differs from the proteins translated during dormancy. In imbibing and germinating seeds, translational machinery operates much more rapidly than in dormant seeds [15]. Protein composition depends first of all on mRNAs stored during maturation [16,17]. At radicle emergence state and after germination, de novo transcription of other genes starts. As a result, germinating seeds contain the complete set of “house-keeping” enzymes involved in ABA synthesis, gibberellin catabolism, protein kinases, protein phosphatases, numerous transcription factors, and typical maturation proteins, namely small heat shock proteins and Late Embryogenesis Abundant (LEA) proteins [12,13]. Special attention was paid to DOG1 protein regulating seed dormancy induction and delaying germination by interfering with PP2C protein phosphatase activity [14]. The translation of proteins occurs slowly in dormant seeds.

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The activation occurs inside the terminal region of regulatory domain at the level of pentultimate threonine, the phosphorylation of which creates a binding site for 1-3-3 protein, which stabilizes the upregulated state of the enzyme, i.e. stabilizes its conformation [19]. Fusicoccin, a specific activator of plasma membrane H⁺-ATPase [22], binds to the enzyme simultaneously with 14-3-3 protein, its molecule being situated within the cavity in the interaction surface between H⁺-ATPase and 14-3-3 protein to tightly couple both partners [19]. In the upregulated state, the enzyme actively extrudes the H⁺ ions to cell walls [23]. The activity of plasma membrane H⁺-ATPase is measured either by rate of ATP hydrolysis, usually by the release of inorganic phosphate, or by H⁺ ion extrusion, estimated by acidification of ambient solution or by pH shift recorded by indicator color. In the experiments, fusicoccin is applied as a specific activator, whereas vanadate and diethylstilbestrol are applied as the inhibitors of ATP hydrolysis.

The capacity of plasma membrane H⁺-ATPase to acidify cell walls and promote cell elongation is well-known as an “acid growth” (acid-induced growth) exhibited in coleoptiles
and hypocotyls, the growth of which starts after radicle protrusion [24,25]. However, no data on the participation of plasma membrane H⁺-ATPase in seed germination is available in literature. We have observed the acidification of cell walls in broad bean, wheat and horse chestnut seeds prior to radicle protrusion that is prior to growth beginning in the embryos of germinating seeds [26,27]. Such acidification is similar to the effect of acid buffer, it is stimulated by fusicoccin that confirms the participation of this enzyme. Analogous activation of ATP hydrolysis occurs just before growth initiation, which is blocked by vanadate. These data indicate a direct participation of the enzyme in seed germination. The experiments with seed incubation in the solutions of fusicoccin stimulating radicle emergence or in diethylstilbestrol and vanadate inhibiting it have confirmed the conclusion.

In seeds, proteomic analysis failed to record the presence of plasma membrane H⁺-ATPase, because the method does not permit to see the proteins of high molecular weight. The mol. wt. of this enzyme is about 100 kDa. The enzyme can be identified by western blotting and application of specific antibodies. It was shown in broad bean embryo axes in the course of imbibition, radicle emergence and growth [27]. Similarly, in non-dormant horse chestnut seeds, the enzyme protein was identified in imbibing, protruding and growing embryo axes. In all samples, 14-3-3 protein was also present [27]. The plasma membrane H⁺-ATPase protein was also shown in the embryos of dry maize seeds, they exhibit a weak activity after short 5h-imbibition, not accompanied by protein synthesis [28,29]. The presence of the enzyme in dry seeds shows that it was synthesized and accumulated earlier, apparently during seed maturation. In seed embryo axes, the delay of imbibition by PEG-6000, osmoticum, retards the acidification, which needs about 68-70% water content for enzyme activation. Therefore, during imbibition the transition of autoinhibited form of H⁺-ATPase to an active one occurs. We followed this transition in horse chestnut seeds, which experience the state of deep physiological dormancy and then dormancy release to acquire the capacity of rapid germination after four months of cold wet stratification [29]. The long period of dormancy and dormancy release culminated in the enzyme activation and germination indicates rather prolonged time of the transition from autoinhibited form to an active one.

In dormant seeds, the rate of cell wall acidification in embryo axes is rather low at the initial stage of embryo dormancy, but twofold increased after 6th week of stratification up to the level typical of emerged embryos, when seeds experience only coat-imposed dormancy [29]. During embryo dormancy, fusicoccin does not exert any stimulating effect either on H⁺ ion extrusion into cell walls or on the size of embryos themselves. It means that during embryo dormancy plasma membrane H⁺-ATPase is still autoinhibited. Its activity and responsiveness to fusicoccin significantly rose after embryo dormancy, when water content in embryo increased and cell preparation for germination occurred. Such activation of the enzyme is not related to indolyl-acetic acid as it was shown for coleoptiles and hypocotyls [24,25], i.e. for postgerminative growth, because the dormancy release and germination of horse chestnut seeds do not respond to the treatments by indolyl-acetic acid or gibberellin.

The difference in plasma membrane H⁺-ATPase behavior over the whole period of dormancy and dormancy release is apparently due to the dynamics of ABA in embryo axis tissues. The content of endogenous ABA manifold increased during the period of embryo dormancy and then declined to very low level by the germination commencement [30]. ABA accumulation correlates in time with the autoinhibited state of the enzyme. The treatment of horse chestnut embryos with 10⁻⁷ M ABA did not inhibit the acidification over this time interval, 10⁻⁴ M influenced weakly, while 10⁻³ M ABA actively inhibited the H⁺-extrusion, apparently because the concentrations of endogenous and exogenous ABA summed. Therefore, the target of ABA inhibitory action on dormant seeds is plasma membrane H⁺-ATPase resulted in the cessation of H⁺ ion delivery to cell walls, the prerequisite of cell growth initiation. Decline of endogenous ABA content leads to the activation of the enzyme and commencement of germination. In non-dormant seeds lacking ABA, the transition of the enzyme to active state is related to an increase in water content.

The described pathway of ABA action on seed germination differs from that described above, namely from the receptor – signaling – transcription system, in two aspects: this pathway is shorter (no ABA-receptor activation) and it does not directly depend on the transcription of new mRNA in imbibing seeds, as follows from the experiments with α-amanitin, the inhibitor of the new mRNAs transcription. After radicle protrusion, when the growth of seedling organs occurs, a postgerminative program includes active operation of plasma membrane H⁺-ATPase [19,31]; the inhibitory action of ABA on their cell elongation is explained by ABA interaction with the receptor, ABA inhibition of protein phosphatase and ABA activation of protein kinase leading to the phosphorylation of an unknown amino acid residue in regulatory domain of the enzyme. This chain of events can result in the inhibition of enzyme, that in its turn causes K⁺ efflux from cytoplasm and its acidification due to the damage of H⁺ ion transport by the enzyme [31]. These observations might give insight into the transition from germination program of seeds to postgerminative program of seedling development.

Acknowledgements

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References


