Brassinosteroid regulation in rice seed biology

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Abstract

Rice (Oryza sativa L.) is not only a model monocotyledon plant, but also an important cereal seed crop. Improvements in seed-related traits is the key to obtaining high grain yield and quality, therefore attracting attention from both plant scientists and crop breeders. In higher plants, brassinosteroid (BR), a major growth-promoting hormone, plays an important role in regulating numerous agronomic traits associated with both vegetative and reproductive growth, thereby presenting huge application potential. Here, we review recent progress into BR regulation in rice seed biology. Both BR biosynthesis and signaling have been shown to regulate grain size, grain filling, grain number, seed germination and biosynthesis of seed components. Thus, considering the pleiotropic effects of BR, strategies aimed at genetic modulation of the BR pathway have been proposed to improve seed-related traits in rice, and therefore, enhance both yield and quality. This review not only strengthens our understanding of the underlying mechanism and regulatory network of BR-regulated key agronomic traits in rice, but also facilitates the future application of BR in rice breeding programs.


Introduction

Rice (Oryza sativa L.) is the staple food of more than half of the world’s population, and production is therefore critical to future food security. Research into rice seed biology is therefore attracting increasing attention from both plant scientists and crop breeders, offering the potential to enhance both yield and quality. The structure of mature rice seeds consists of a spikelet hull, seed coat, aleurone layer, endosperm and embryo, with the embryo containing most of the genetic information, and the endosperm storing nutrients and forming the edible component. Rice seeds showing excellent performance during grain development, seed dormancy and germination are critical in guaranteeing high yield and quality. Understanding the regulatory molecular mechanism of key rice seed traits is therefore essential in improving and breeding elite varieties. For example, high-vigor seeds are suitable for direct seeding production, reducing labor costs and increasing planting efficiency11, while high-yielding, high-quality rice varieties, with emphasis on taste and appearance, are preferred by farmers12.

In general, rice seed traits are controlled by both endogenous and environmental factors, with phytohormones playing a central role, not only as a critical intrinsic regulator, but also by conveying environmental input. The most well-known plant hormone family includes gibberellin (GA), abscisic acid (ABA), auxin, cytokinin, ethylene, brassinosteroid (BR), jasmonic acid (JA) and strigolactone3, all of which play essential roles in seed-related traits, such as ripening (ethylene)4, dormancy (ABA)5, and germination (GA)6. As a major plant growth-promoting phytohormone, BR is widely involved in a number of rice growth and developmental events, including plant growth, leaf angle, tillering, pollen fertility, grain size and seed germination (Fig. 1). Considering the potential application of BR in improving important agronomic traits in rice, this study reviews the role of this plant hormone in seed biology, such as regulation of grain size, grain filling and seed germination7−9. The findings provide valuable information for genetic modification of specific hormone pathways aimed at optimizing seed-related traits in rice.

The most well-known effect of BR is the promotion of plant growth. BR-deficient rice mutants resulting from mutation of BR biosynthesis genes, such as BRD110, D21, D112, exhibit dwarf and compact phenotypes, while blocking of BR signal transduction was found to trigger a similar phenotype, with reduced plant height and leaf angle11. At present, the BR signaling pathway is one of the most understood signal transduction pathways in plants, especially in the model plant Arabidopsis113. Although progress in BR-related studies in rice is lagging behind that of Arabidopsis, the primary signaling pathway is conserved. Briefly, BR signals are first sensed by the receptor OsBRI1 and coreceptor OsBAK1, and then transmitted to OsBSK3 kinase and an unidentified phosphatase, thereby suppressing OsGSK2 kinase activity114,10. OsGSK2, a GSK3/HAGGY-like kinase, is the core negative regulator of the BR signaling pathway, inhibiting BR signals by phosphorylating downstream key transcription factors, such as OsBZR1 and DLT17−19. Rice plants overexpressing OsGSK2 or presenting DLT mutation showed similar phenotypes to the BR receptor mutant d6117,20, highlighting the essential role of BRs in normal growth and plant architecture. In addition, BRs also play multiple roles in rice seed growth and development15.
**BR regulation of grain size**

Grain size is an important agronomic trait closely related to rice yield and quality. As a complex quantitative trait, grain size is controlled by a series of genes, of which more than 80 have so far been cloned[2]. Based on the regulatory pathways, these genes can be divided into six categories: plant hormones, mitogen-activated protein kinase (MAPK) signaling, the G-protein pathway, ubiquitin-proteasome pathway, HAIKU (IKU) pathway and transcriptional regulatory factors[21]. At least four plant hormones; namely, BR, auxin, cytokinin and GA, have been reported to be directly or indirectly involved in the regulation of rice grain size[22], and of these, BR is a major regulator, with more than 20 genes having so far been cloned. According to their specific roles in the BR pathway, these genes can be further divided into three categories: components of BR biosynthesis or signaling, targets of BR signaling pathway, and regulators of the BR pathway (Fig. 2).

**Components of BR biosynthesis and signaling**

Mutation of the BR synthesis-related genes *BRD1*, *D2* and *D11* produced small grains, suggesting that BR plays positive roles in controlling rice grain size[10–12]. A similar small-grained phenotype was also observed in BR insensitive mutants[7], such as *d61* and *bzr1[23]*, while large rice seeds were generated following overexpression of *OsBSK3* and knock-down of *OsGSK2*, respectively[17,24]. Recently, a kinesin-13a protein BHS1 was identified to be a new potential BR signaling component. BHS1 regulates rice grain length by negatively modulating BR signaling downstream of *OsBRI1[25]*. These results further suggest that BR plays a positive role in regulating rice seed size.

In addition, *OsBSK2*, a BR-signaling kinase functioning downstream of *OsBRI1*, positively regulates rice grain size independent of the BR signaling pathway[26]. As the central negative component of the BR signaling pathway, the activity and status of Glycogen synthase kinase 3 (GSK3) family members are also closely related to the regulation of rice grain size. Manipulation of GSK3 family members, such as *OsGSK2*, *OsGSK3* and *OsGSK5*, was found to alter rice grain size[17,19,27,28]. For example, *qGL3* encodes the protein phosphatase OsPPKL1, which contains two Kelch domains, and affects BR signal transduction by dephosphorylating and stabilizing OsGSK3 kinase[27]. Meanwhile, a rare allelic variation of *qgl3* in the second Kelch domain was found to result in a long-grained phenotype[29]. Furthermore, OsAK3, which interacts with *qGL3* both in vivo and in vitro, results in a longer grain length by controlling cell expansion in rice spikelet glumes[30]. In addition, *DLT* and *BZR1*, downstream transcription factors directly regulated by *OsGSK2*, were found to function as positive regulators of rice grain size[7,17,31]. Recent research further suggests that *qGL3* can induce phosphorylation of the 14-3-3 protein OsGF14b, consequently inhibiting OsBZR1 function by promoting cytoplasmic retention and suppressing transcriptional activation activity, which negatively regulated grain length in rice[32]. In addition to its role in the BR signaling pathway, OsPPKL1 is proven to be a cryptic inhibitor of cytokinin phosphorelay which regulates rice grain size, suggesting that PPKL1 may have dual roles in modulating both hormones[33]. Furthermore, disrupting the function of OsCP3, a member of the RNA Pol II CTD phosphatase-like family, will increase *OsGSK2* abundance and decrease *OsBZR1* levels, resulting in changes in rice grain size[34].

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**Fig. 1** Brassinosteroid regulates a series of key agronomic traits in rice, including plant height, pollen fertility, leaf angle, tiller number, grain size and seed germination.
**Targets of the BR signaling pathway**

In addition to the key elements of BR biosynthesis and signaling, other downstream components of the BR signaling pathway, notably those directly or indirectly controlled by OsGSK2 kinase, also play essential roles in regulating rice seed size. For example, OsGSK2 directly interacts with several members of the OsOFP family proteins, such as OsOFP1, OsOFP3 and OsOFP8, modulating their phosphorylation status, protein stability or activity, and ultimately affecting grain size \[35-37\]. Interestingly, the activity of phosphorylated OsOFP1 and OsOFP8 was found to decrease \[35,37\], while the stability and protein accumulation of phosphorylated OsOFP3 increased \[36\]. In addition, overexpression of OsOFP1 and OsOFP8 significantly increased grain length in rice, while OsOFP3 overexpression had the opposite effect. These results imply that OsGSK2 either enhances the accumulation of negative regulators such as OsOFP3 or suppresses the activity of positive regulators such as OsOFP1 and OsOFP8, thereby acting as a key negative regulator of grain length.

In addition to OsOFPs, several other OsGSK2-interactive proteins related to grain size have also been identified and cloned. For example, OsWRKY53, which is phosphorylated and suppressed by OsGSK2, was found to interact with BZR1 and synergistically control BR-regulated plant traits, including rice grain length \[38\]. Meanwhile, OsGAMYBL2, another OsGSK2-interacting protein, is destabilized by BR, with up- and down-regulated expression causing a decrease and increase in rice grain size, respectively \[39\]. Moreover, OsGSK2 was found to directly interact with GS2/GL2/OsGRF4, inhibiting transcription activation activity, and thereby altering cell size \[40\]. In addition, GS2 has been proven to be involved in the regulation of panicle length, seed shattering and cold resistance in rice \[41,42\]. As a target of OsmiR396, GS2 also interacts with transcriptional co-activators OsGIFs, forming a complex regulatory module, which predominantly affects cell expansion \[43,44\].

GS9 is a negative regulator of grain length and a positive regulator of grain width, with gs9 mutant rice seeds exhibiting a slender grain shape. Importantly, GS9 mutation had no effects on rice growth or development, thus improving grain shape without sacrificing yield or other major agronomic traits. Analysis of the underlying molecular mechanism revealed that GS9 directly interacts with OsOFP8 protein, the target of OsGSK2 kinase \[45\]. Thus, GS9 is indirectly regulated by OsGSK2 via OsOFP8, thereby affecting BR-induced regulation of grain size. Identification of genes downstream of BR that affect grain size would therefore enrich our understanding of the BR signal.
transduction pathway, and also contribute to the specific improvement of grain size in rice.

**Regulators of BR biosynthesis and signaling**

Some genes control seed size in rice by affecting BR synthesis or signal transduction. For example, **RAV6** and **XIAO** and **SLG** regulate rice grain size by modulating the expression of BR biosynthesis genes, such as **D2** and **D11**. In more detail, enhanced expression of **SLG** improved BR content and increased grain length, while knock-out of **XIAO** led to a distinct BR-deficient phenotype, with short grains. Meanwhile, the effects of **RAV6** are more complex. Elevated expression promoted BR content and rice leaf angle, but reduced grain size, possibly due to the complex down-stream target genes of **RAV6**. **CYP734A4** genes affect the level of bioactive BRs by degrading BRs encoding cytochrome P450 monooxygenases. Plants overexpressing **CYP734A4** as a result of a T-DNA insertion showed a typical BR-deficient phenotype, including dwarfing and grain shrinkage. On the other hand, **OsREM4** plays a key role in inducing seed dormancy and inhibiting seed germination under both normal and abiotic stress conditions is also the target gene in **RAmy3D** seed germination under both normal and abiotic stress. The results showed that both the **GSK2** overexpressing line and **bzz1** mutants germinated slower than their respective wild-type controls.

**BR regulation of seed germination**

Rice germination involves a series of orderly physiological and morphological changes following imbition and expansion of the seeds, which usually begins with rapid absorption of water and ends with radicle protrusion. Normal seed germination requires a number of prerequisites, mainly light, temperature, water, nutrients and phytohormones. Of the various plant hormones, GA is dominant in promoting seed germination, with GA-deficient mutants in Arabidopsis showing strong dormancy and an abnormal germination phenotype in the absence of exogenous GA treatment. In contrast, ABA plays a key role in inducing seed dormancy and inhibiting seed germination, with ABA-deficient Arabidopsis mutants showing significantly faster seed germination than the wild-type. Moreover, GA and ABA are antagonistic in regulating seed germination, with spatial and temporal balance determining germination versus dormancy. In addition, GA also promotes seed germination by mobilizing stored starch in rice. This occurs mainly via GA synthesis and secretion into aleurone layer cells in the embryo tissue, which activates the transcription factor GAMYB, thereby initiating the expression of α-amylase and feedback to the endosperm, finally inducing starch hydrolysis into glucose and other small molecular sugars for embryo growth. However, the ABA-inducible **OsWRKY51** gene cooperates with the GA-repressible **OsWRKY71** gene to inhibit transcriptional activity of GAMYB in rice embryo and aleurone cells. Recently, several genes affecting seed germination and dormancy were cloned in rice mutants, with roles in GA biosynthesis (**OsKOT**), **ABA** signal transduction (**OsMFT2**, **ISA1**, **OsPP2C09** and **ABA** metabolism (**OsTPP**, **OsZmP09**). Compared to their respective wild-type controls, these mutants showed either strong germination or dormancy characteristics.

**Direct regulation of rice seed germination by BR**

In addition to GA and ABA, other hormones, including BR, play a role in fine-tuning the rice germination process. For example, shoot length and the germination rate of rice seeds was found to decrease significantly when treated with the BR synthesis inhibitor brassinazole. BR is not only an important regulator of seed germination, but also plays key roles in controlling other agronomic traits related to rice yield, such as grain size, leaf angle, and tiller number. Thus, further understanding of the mechanisms underlying BR-regulated seed germination in rice will aid breeding programs aimed at improving yield-related traits.

Although Arabidopsis and rice share a similar BR signaling pathway, little is known about the mechanism of hypocotyl elongation and germination in rice compared to Arabidopsis. In rice, BR-deficient mutants, such as **brd1**, **d2**, **gns4** and **nbg4**, exhibit a slower germination rate and shorter mesocotyls. Meanwhile, a germination experiment in the dark showed that **d61**, a BR-insensitive mutant, has a remarkably shorter coleoptile and radicle length. These studies indicate that both BR synthesis and signal transduction are involved in the regulation of rice seed germination and post-germination growth. However, the underlying molecular mechanisms remain largely unknown.

To determine the mechanism of BR-regulated seed germination, a series of BR biosynthesis and signaling related rice mutants were used in a germination assay. The results showed that both the **GSK2** overexpressing line and **bzz1** mutant germinated slower than their respective wild-type controls. Meanwhile, further experiments revealed that the **BZR1-Ramy3D** centered transcription module is critical in accelerating the degradation and utilization of starch in both the endosperm and embryo of rice seeds, thus promoting seed germination (Fig. 3). As the key **α-amylase** gene in rice, **Ramy3D**, which is also known as **αAmy3**, acts as a positive regulator of seed germination. Overexpression of **αAmy3** could promote rice seed germination under both normal and abiotic stress conditions. Interestingly, **Ramy3D** is also the target gene in responses to hypoxia and sugar starvation signals, with the regulatory pathway mediated by the **CIPK15-SnRK1A-MYBS1-Ramy3D** signaling cascade (Fig. 3). More importantly, since **Ramy3D** is highly expressed during early stages of seed germination, disruption of the **BZR1-Ramy3D** regulatory...
Co-regulation of rice seed germination via crosstalk between BR and GA

Notable progress has been made in determining the molecular mechanisms underlying BR-GA crosstalk during growth regulation in both rice and Arabidopsis [75,81–84]. Although low levels of BR promoted expression of the GA biosynthesis gene GA3ox-2 in rice, high levels inactivated GA by inducing expression of the GA inactivation gene GA2ox-3, with feedback inhibiting BR biosynthesis [85]. This is also the main reason why a high concentration of BR inhibits rice growth. Based on these studies, the mechanism of BR-GA co-regulated rice seed germination has been gradually revealed. First, GA was found to recover seed germination defects in BR-deficient and insensitive rice seeds, while an iTRAQ proteomic approach identified the differentially abundant target proteins involved in this process [85]. Accordingly, a total of 42 target proteins were identified, five of which are LEA family proteins. Expression of these LEA family members was further found to be suppressed by GA at both the transcript and protein levels. Meanwhile, genetic evidence suggests that LEAs function downstream of both the BR and GA pathways, with LEA mutation inhibiting rice seed germination [85]. Moreover, BR and GA were found to induce the expression of REP-1 in both the rice embryo and endosperm during seed germination, thereby affecting the turnover of storage proteins, including glutelin, which...
Conclusions and perspectives

BR, as a principal growth-promoting plant hormone, plays a wide range of roles in plant growth and development. In rice, BR has at least three potential applications in breeding practices. First, by attenuating BR biosynthesis or signaling to improve plant architecture. With the reduction in arable land and increasing demands for food, breeding of compact rice varieties complemented by intensive planting has been deemed a practical strategy for reducing competition for water, light and nutrients, thereby improving yield per unit area. Second, BR could also be used to regulate a series of seed-related traits, including grain size, grain filling, grain number and seed components, thereby increasing rice yield and quality. Third, BR involvement in the regulation of seed germination and PHS shows further potential, without affecting the germination rate, with the bzh1 mutant showing slightly delayed germination but strong resistance to PHS. However, due to the pleiotropic effects of BR, improving rice seed-related traits by enhancing the BR pathway also has negative effects on other agronomic traits; for example, increasing the leaf angle and loosening the overall plant architecture. Meanwhile, BR-deficient and insensitive rice mutants typically exhibit dwarfism and a compact plant phenotype, although their grain size is also smaller. At present, the most effective way of applying the positive aspects of BR while avoiding its negative effects is to use weak BR mutants or downstream target genes with specific roles in regulating certain agronomic traits. For example, osdwarf4-1, a weak BR-deficient mutant, causes only limited defects in rice morphology, including a slightly dwarfed stature and more erect leaves, while the yield was higher than that of the wild-type control under dense planting conditions. Another example is G59, a specific regulator of rice grain size functioning downstream of the BR signaling pathway. G59 mutation improved rice appearance quality, presenting a slender shape and reduced chalkiness, but without observable negative effects on any other agronomic traits.

Rice is one of the most important crops in the world, while its seed traits are directly correlated with yield and quality. Studies aimed at improving seed-related traits are therefore attracting increasing attention from both plant scientists and rice breeders. However, although BR plays comprehensive roles in regulating multiple seed traits, little is known about the underlying molecular mechanism and regulatory network. For example, BR interacts with GA and ABA to co-regulate rice seed germination, as demonstrated by a series of physiological and preliminary molecular studies; however, the exact integrative nodes remain unclear. Future application of powerful and accurate technologies, accompanied by additional genetic resources and a more complete understanding of the BR signaling pathway in rice will therefore help uncover the mechanisms underlying BR-regulated seed traits in rice.
Acknowledgments

This work was supported by the Hainan Yazhou Bay Seed Lab (B21HJ8105), the National Natural Science Foundation of China (32071984, 31971914), and the Programs from Jiangsu Province Government (BK20200045, JBGS[2021]001, SWYY-154 and PAPD).

Conflict of interest

The authors declare that they have no conflict of interest.

Dates

Received 16 May 2022; Accepted 6 July 2022; Published online 20 July 2022

REFERENCES


Brassinosteroid regulation of rice seed traits