Letter

OsGATA7 modulates brassinosteroids-mediated growth regulation and influences architecture and grain shape

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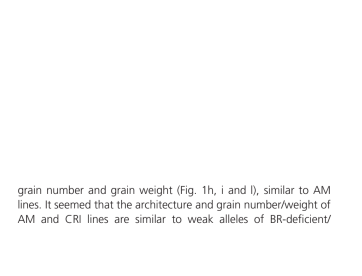


Figure 1 OsGATA7 modulates BR-mediated growth regulation in rice. (a) The expression pattern of OsGATA7.GUS staining assay of pOsGATA7-GUS lines shows GUS activity in stamen and pistil (i), lemma (ii), young root (iii), young leaf blade (iv), lamina joint (v), internode (vi). Bars = 2 cm. qRT-PCR analysis shows the relative expression level of OsGATA7. (b) The panicles of WT and AM lines. Bar = 4 cm. (c) Statistical analysis of primary branch number and grain number of WT and AM lines. Values are means \pm SE (n = 6–11). **P < 0.01, *P < 0.05. The Student's t-test was used to analyse the significant differences between wild type and AM lines (the same below). (d) The grains of WT and AM lines. Bars = 1 cm. (e) Statistical analysis of grain length and width of WT and AM lines. Values are means \pm SE (n = 35–54). **P < 0.01, *P < 0.05. (f) Statistical analysis of weight per 1000 grains. Values are means \pm SE (n = 3), **P < 0.01. (q) The adult plants of WT and CRI lines. Bar = 10 cm. (h) The panicles of WT and CRI lines. Bar = 3 cm. (i) Statistical analysis of primary branch number, secondary branch number and grain number per plant. Values are means ± SE (n = 9-29, 9-19 and 8-12). **P < 0.01, *P < 0.05. (j) The grains of WT and CRI lines. Bars = 1 cm. (k) Statistical analysis of grain length and width in WT and CRI lines. Values are means \pm SE (n = 27–32). **P < 0.01. (I) Statistical analysis of weight per 1000 grains, grain yield per plant (values are means \pm SE, n = 3–6, 9–19, **P < 0.01. *P < 0.05) and the results of grain yield per unit area. (m) Statistical analysis of root length of WT and AM lines after 12-day eBL treatment. Values are means \pm SE (n = 17–20). (n) The relative expression level of OsGATA7 in 6-day-old seedlings after 1 \times 10⁻⁶ M eBL treatment under light and in darkness. (o) The grains of WT and d61. Bars = 1 cm. (p) Statistical analysis of grain length and width in WT and d61. Values are means \pm SE (n = 10). *P < 0.05. (g) Statistical analysis of the outer epidermal cell number in the grain (length) of lemma by scanning electron microscope. Values are means \pm SE (n = 10). **P < 0.01. (r) Overexpression of OsGATA7 in Arabidopsis BR-insensitive mutant bri1-5. Bar = 1 cm.

signal and response, these lines have better growth condition than reported BR-related mutants, indicating OsGATA7 has not as severe influences as BR signalling in rice growth, especially in reproductive development. Even the grain number and weight are slightly lower in individual plant of AM and CRI1 lines, and the selfed-seed fertilities of CRI1 lines were slightly lower than WT (AM lines were similar to WT), the yield analysis still illustrated that the grain weight of CRI lines was higher (30.4% and 43.3% in CRI7-1 and CRI-5 line) than wild type in the same area (Fig. 1l) because of dense planting as these lines have compact architecture. Besides, the grains (as well as brown rice) produced from AM lines (Fig. 1d and e) and CRI lines (Fig. 1j and k) are longer and narrower than WT. Detailed analysis of the outer epidermal cell showed that AM lines had increased cell number (Fig. 1g) but not cell elongation, whereas BR was considered to mainly induce cell elongation. Grain shape is a preference which links to rice quality. Our statistical analysis (Fig. 1o and p) further demonstrated that BR-insensitive mutant d61 had shorter and round grain, suggesting that OsGATA7 played different roles in grain shape regulation with BR signalling. Taken together, OsGATA7 functions in controlling rice plant architecture and panicle/grain development and the knock-down/genome-edited lines have ideal traits of both architecture and grain shape. The CRI lines have enhanced grain yield than WT.

As well as AM and CRI lines, we also constructed the overexpression lines of OsGATA7 (pUbi-OsGATA7). There was no significant phenotype in adult plant of overexpression lines, which might be because that the increased OsGATA7 transcription level in wild type would not be enough to induce growth phenotypes or OsGATA7 worked with other partners and could not induce growth phenotypes by overexpressing it alone. Many literatures reported Arabidopsis BR-deficient or BR-insensitive mutants have phenotypes of dwarf, round leaves, shorter petioles and reduced reproductively. Overexpression of OsGATA7 could partially rescue phenotypes of Arabidopsis BR-insensitive mutant bri1-5, such as plant height (Fig. 1r), round leaves and short petioles, but not rosette leaf sides (important for Arabidopsis plant density), which further suggested that OsGATA7 enhanced BR signalling and partially modulated BR-mediated plant growth in some processes.

Above all, OsGATA7 is a multifunction gene regulating rice growth. The plant height, leaf inclination, panicle development and grain number/shape/weight of OsGATA7 knock-down/ genome-edited lines suggest OsGATA7 is involved in BRmediated growth regulation. Interestingly, BR positively

regulates OsGATA7 expression, and OsGATA7 also affects BR signalling and sensitivity, indicating BR and OsGATA7 synergistically regulate some processes of rice growth. But OsGATA7 has diverse functions in others processes, such as grain shape regulation, indicating that OsGATA7 has only partial overlap with BR signalling regulation of rice growth. Partially rescued phenotypes of Arabidopsis bri1-5 mutant by OsGATA7 overexpression further demonstrated that OsGATA7 modulated some BR-mediated growth regulations, and OsGATA7 also had BRindependent regulation in plant growth, especially in reproductive growth. AM and CRI lines are suitable for dense planting because of the ideal architecture, especially CRI lines (more compact than AM lines). The enhanced plant number in the same area, and no severe reduced grain weight and number in individual plant, increases the grain yield in the unit area obviously. And so far we did not observe that the AM and CRI1 lines are more sensitive to pathogen/herbivore or abiotic stress, which mean they would be putative candidate lines for breeding. Taken together, our work identified OsGATA7 functioning in BR-mediated architecture regulation, panicle development and grain shape/number/weight/vield. OsGATA7 modulates BR-mediated rice growth regulation and avoids sideeffects of BR signalling, which would be a putative candidate gene for potentially using in agriculture production as OsGATA7 knock-down/genome-edited lines have ideal architecture, better grain shape, and enhanced grain yield.

Acknowledgments

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References

Hudson, D., Guevara, D.R., Hand, A.J., Xu, Z., Hao, L., Chen, X., Zhu, T. et al. (2013) Rice cytokinin GATA transcription Factor1 regulates chloroplast development and plant architecture. Plant Ph. siol. 162, 132-144.

Reyes, J.C., Muro-Pastor, M. and Florencio, F.J. (2004) The GATA family of transcription factors in Arabidopsis and rice. Plant Ph siol. 134, 1718–1732.

- Sumiyo, T., Motoyuki, A., Shozo, F., Suguru, T., Shigeo, Y., Masahiro, Y., Atsushi, Y. *et al.* (2005) A novel cytochrome P450 is implicated in brassinosteroid biosynthesis via the characterization of a rice dwarf mutant, *d arf11*, with reduced seed length. *Plant Cell*, **17**, 776–790.
- Wang, L., Yin, H., Qian, Q., Yang, J., Huang, C., Hu, X. and Luo, D. (2009) NECK LEAF 1, a GATA type transcription factor, modulates organogenesis by
- regulating the expression of multiple regulatory genes during reproductive development in rice. *Cell Res.* **19**, 598–611.
- Yamamuro, C., Ihara, Y., Wu, X., Noguchi, T., Fujioka, S., Takatsuto, S., Ashikari, M. *et al.* (2000) Loss of function of a rice brassinosteroid insensitive1 homolog prevents internode elongation and bending of the lamina joint. *Plant Cell*, **12**, 1591–1606.