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Hemayet Ullah, Howard University

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Arabidopsis scaffold protein RACK1A modulates rare sugar D-allose regulated gibberellin signaling

Herman Fennell,1 Abdulquddri Olawin,1 Rahman M. Mizanur,2 Ken Izumori,3 Jin-Gui Chen4 and Hemayet Ullah1,*

1Department of Biology; Howard University; Washington, DC USA; 2US Army Medical Research Institute of Infectious Diseases (USAMRIID); Fort Detrick; Frederick, MD USA; 3Faculty of Agriculture; Kagawa University; Kagawa, Japan; 4Biosciences Division; Oak Ridge National Laboratory; Oak Ridge, TN USA

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As energy sources and structural components, sugars are the central regulators of plant growth and development. In addition to the abundant natural sugars in plants, more than 50 different kinds of rare sugars exist in nature, several of which show distinct roles in plant growth and development. Recently, one of the rare sugars, D-allose, an epimer of D-glucose at C3, is found to suppress plant hormone gibberellin (GA) signaling in rice. Scaffold protein RACK1A in the model plant Arabidopsis is implicated in the GA pathway as rack1a knockout mutants show insensitivity to GA in GA-induced seed germination. Using genetic knockout lines and a reporter gene, the functional role of RACK1A in the D-allose pathway was investigated. It was found that the rack1a knockout seeds showed hypersensitivity to D-allose-induced inhibition of seed germination, implicating a role for RACK1A in the D-allose mediated suppression of seed germination. On the other hand, a functional RACK1A in the background of the double knockout mutations in the other two RACK1 isoforms, rack1b/rack1c, showed significant resistance to the D-allose induced inhibition of seed germination. The collective results implicate the RACK1A in the D-allose mediated seed germination inhibition pathway. Elucidation of the rare sugar signaling mechanism will help to advance understanding of this less studied but important cellular signaling pathway.

Rare sugars—monosaccharides that rarely exist in nature—are known to regulate diverse physiological responses in both plants and animals. With the advent of the discovery of “Izumoring,” an in vitro enzymatic approach to the synthesis of all rare sugars,1,2 research in this area has implicated rare sugars in many physiological conditions. These include but are not limited to the immunosuppressive activity in liver transplantation,3 protection against liver ischemia reperfusion injury,5,6 protection from reactive oxygen species (ROS)7,8 and anticancer activity on different cancer cell lines.9–11 Although not abundant, rare sugars—D-allose, D-psicose, D-allitol, L-galactose, tagatose or their derivatives—have been found in the tissues of higher plants as well.12–18 It was found that D-psicose inhibits plant root growth via hexokinase-independent pathway; it also inhibits bacterial blight disease in rice.19–20 D-allose was found to inhibit the rice growth and kinase-independent pathway; it also inhibits bacterial blight disease in rice as well.21

Recently, Akimitsu’s group, a pioneer in rare sugar research, through the use of powerful genetics study has shown that the rare sugar D-allose suppresses GA signaling pathway in rice.22 D-allose strongly inhibited GA mediated α-amylase induction in embryo-less rice half seeds,22 implicating a negative role of D-allose in the GA pathway. Earlier, we have shown that Arabidopsis scaffold protein RACK1A (Receptor for Activating C Kinase 1) positively regulates GA signaling pathway as rack1a knockout seeds showed insensitivity to GA-induced seed germination.23 Here we tested our hypothesis that the suppression of GA signaling pathway by D-allose would impact the RACK1A mediated positive regulation of GA signaling pathway.

Scaffold protein RACK1 in metazoan plays a major role in coordinating different signal transduction pathways ranging from cell division to ion channel regulation by interacting with diverse proteins.24–28 RACK1 proteins with seven WD-40 repeats are highly conserved (70–80% at the protein level) in wide range of species, including plants, humans, rats, chickens, flies, nematodes, algae and yeasts. Although not recognized as such, the first RACK1 homolog in plants was identified in tobacco BY-2 suspension cells as a plant hormone auxin inducible gene29 and later in Arabidopsis, rice, rape and alfalfa.30–33 As opposed to a single gene in metazoan, the model plant Arabidopsis genome maintains three different RACK1 genes—termed as RACK1A, RACK1B and RACK1C.34 These Arabidopsis genes are found to regulate plant development with unequal genetic redundancy.34 The analysis of double and triple Arabidopsis rack1 mutants revealed that the difference in gene expression level and the cross-regulation

*Correspondence to: Hemayet Ullah; Email: hullah@howard.edu
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may determine the role played by the individual RACK1 gene in regulating plant development. In addition to regulating plant development, Arabidopsis RACK1 mediates multiple hormonal and stress responses.

In order to elucidate the role of D-allose in Arabidopsis seed germination pathways, age-matched seeds of designated genotypes were grown under continuous light (75 μmol/m²/sec) at 22°C in MS minus sugar liquid media containing either 10 mM D-allose or 10 mM D-allose and 10 μM of Gibberellins (GA) or 10 mM of D-glucose for two weeks. Compared with the no treatment control, D-allose inhibited WT seed germination and inhibition is enhanced in the rack1a knockout line. rack1b/rack1c double mutant line partially overcomes the D-allose mediated inhibition of seed germination. D-glucose control treatment does not affect the seed germination in any of the genotypes (lower panel).

**Figure 1.** Seed germination inhibition by D-allose is enhanced in rack1a knockout line. Age matched seeds from indicated genotypes were grown under continuous light (75 μmol/m²/sec) at 22°C in MS minus sugar liquid media containing either 10 mM D-allose or 10 mM D-allose and 10 μM of Gibberellins (GA) or 10 mM of D-glucose for two weeks. Compared with the no treatment control, D-allose inhibited WT seed germination and inhibition is enhanced in the rack1a knockout line. rack1b/rack1c double mutant line partially overcomes the D-allose mediated inhibition of seed germination. D-glucose control treatment does not affect the seed germination in any of the genotypes (lower panel).

significantly downregulated compared with that in the WT seedlings, RACK1A expression was not significantly downregulated in the rack1b/rack1c mutant background. If RACK1B/RACK1C collectively exert negative regulation on RACK1A expression, it will be quite informative to see the response of rack1b/rack1c double mutant to D-allose induced seed germination inhibition. Quite interestingly, it was found that neither D-allose nor D-allose plus GA could significantly affect the seed germination rate and the early seedling development in the rack1b/rack1c double mutant background (Fig. 1F and I). On the other hand, all the seeds without any D-allose application did germinate and showed normal growth and development (Fig. 1A–C). As D-allose is an epimer of D-glucose, we used D-glucose as a control to make sure that the effect is very specific to D-allose. As can be seen from Figure 1J–L, all three genotypes of the tested seeds (WT, rack1a-1 knockout and rack1b/rack1c double knockout) did not show germination inhibition from D-glucose treatment, effectively eliminating the possibility that the D-allose acts as a simple carbon source. The results from the WT seedlings, where a functional RACK1A is present, can be explained by the presence of the two more isoforms of RACK1A—which can potentially cross-regulate the RACK1A function. It is reported that the knockout phenotype of rack1a is enhanced in either rack1b or rack1c knockout combination, implying the role of RACK1B or RACK1C in the RACK1A function.

To understand the cellular mechanism of this signaling network, it is proposed that D-allose will not only negatively regulate GA-induced seed germination and growth, it may potentially downregulate RACK1A expression to attenuate the positive regulation of GA signaling by RACK1A. We utilized a transgenic line expressing GFP fused to the RACK1A gene which is driven by the RACK1A native promoter. As can be seen in Figure 2, the application of D-allose to the transgenic seeds (Fig. 2C and D), compared with the no-treatment control (Fig. 2A and B) did significantly downregulated RACK1A expression in the root tip region of the embryo isolated from seeds treated for 72 h. Application of GA could significantly enhance the RACK1A expression in the same tissue regions (Fig. 2E and F). D-Glucose was used as a control, which, contrary to the D-allose, slightly increased RACK1A expression (data not shown), indicating that the D-allose specifically downregulates RACK1A expression in the embryonic root tip region. Although the gene expression was restricted to the root tip growth area, it remains to be seen whether the similar regulation takes place in the shoot apical meristems of young seedlings. Even though it is known that D-allose exert its inhibitory role on GA signaling through hexokinase-dependent pathway, it will be an intriguing study to investigate the precise role hexokinase plays in the currently elucidated pathway. Figure 3 presents a possible working model of the pathway. The simple explanation for the presented data are that D-allose negatively regulates GA-mediated seed germination and early seedling development through the inhibition of RACK1A expression. Though RACK1A is reported to positively regulate GA signaling pathway, in the absence of epistasis studies between rack1a and known GA signaling mutants, it is not quite possible to indicate a downstream or upstream regulatory.

Figure 1.
position of RACK1A in the GA signaling pathway. However, the presumptive position of the RACK1A in the current model is partially consistent with the presented data but with future genetic studies the more concrete regulatory position in this pathway will be ascertained.

Conclusion

D-allose has shown its promise in diverse physiological and biological applications in animal and plant studies. The current study fills up a gap in the understanding of the prominent role of D-allose in the seed germination signaling pathway in Arabidopsis. Considering the huge economic values in understanding the seed germination signaling pathway, the current study results will help in advancing the molecular elucidation of this pathway. In addition, the role of growth hormone GA induced seed germination process is an intensely studied physiological processes and the precise understanding of the signaling pathways will potentially contribute to the better elucidation of the pathway. Understanding the D-allose mediated cellular signaling mechanism during the seed germination process, can potentially contribute to wider applied use of this rather under-estimated important physiological regulator.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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References


