

Plants recruit distinct chemical activities of coumarins under different soil pHs

Gatersleben, 29.11.2023 Iron is an essential micronutrient for plants and is generally found in large quantities in the Earth's crust. However, its availability to plants is highly dependent on the pH value of the soil. When suffering from iron deficiency, certain plant species release coumarins. Which type of these small secondary metabolites is mainly released depends on the pH value of the soil. Using the model plant *Arabidopsis thaliana* as an example, a research team led by the IPK Leibniz Institute has shown how different coumarins are used to maximize iron uptake under acidic and alkaline conditions. The results were published in the journal "The Plant Cell".

Plants have two main uptake mechanisms to obtain iron (Fe) from the soils. The type of strategy employed depends on the botanical classification of the plant. In the so-called strategy-I mechanism, plants must first reduce the trivalent iron (Fe³+) into bivalent iron (Fe²+). Only then can Fe²+ be taken up by roots as a free ion. This strategy is used by non-Poaceae plants such as oilseed rape and the model plant *Arabidopsis thaliana*. Grasses, which belong to the Poaceae familiy, employ a so-called strategy II. These plants secrete chelating compounds, which can be reuptaken once chelated with Fe³+. Thus, no reduction step is required for Fe import into root cells.

"Interestingly, some Strategy-I plants also release metabolites into the soil through their roots when they suffer from Fe deficiency. Some of these are coumarins," explains IPK scientist Dr. Ricardo Giehl, co-head of the "Molecular Plant Nutrition" working group. However, the physiological role of these coumarins has not yet been sufficiently clarified.

In their work, the researchers were able to show that the main function of two coumarins released in response to Fe deficiency is largely dependent on the external pH. Under slightly acidic conditions, coumarins, especially sideretin, help to sustain Fe³⁺ reduction. Here, sideretin works together with the membrane-bound enzyme FERRIC REDUCTION OXIDASE 2 (FRO2) to efficiently Fe²⁺ uptake into the roots. If the medium is alkaline, the biosynthesis of coumarins is shifted from sideretin to fraxetin, a response that the research team found to depend on the transcription factor MYB72. At alkaline pH, sideretin loses both its ability to reduce and even to solubilize Fe³⁺ from precipitated sources, while fraxetin retains a high Fe³⁺ mobilization capacity under such pH conditions. Therefore, rather than reducing Fe³⁺ directly, the main function of fraxetin is to provide soluble Fe(III)-chelates for FRO2-mediated reduction.

"Our study shows that by adjusting coumarin biosynthesis, plants recruit specific functions depending on the prevailing pH of the soil," says Dr. Ricardo Giehl. If the

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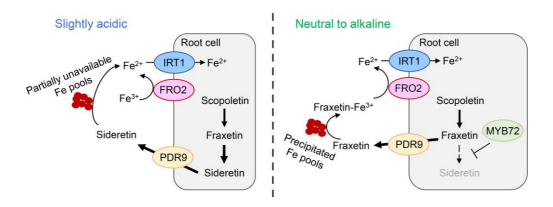
Media Contact Christian Schafmeister Phone: +49 39482 5461 schafmeister@ipk-gatersleben.de conditions are slightly acidic, plants favor the synthesis of the superior Fe³⁺ reductant sideretin, while at high pH they direct synthesis towards fraxetin, which retains high Fe³⁺ mobilization capacity even under alkaline conditions.

With their work, the researchers provide valuable insights into environment-dependent fine regulation of metabolite biosynthesis and thus help to further understand how plants adapt to different pH conditions in the soil. The results open up new possibilities for the targeted improvement of plant productivity and plant health under variable soil conditions.

Original publication:

Paffrath *et al.* (2023): A major role of coumarin-dependent ferric iron reduction in strategy l-type iron acquisition in Arabidopsis. The Plant Cell. DOI: <u>10.1093/plcell/koad279</u>

Graphic:



Coumarin biosynthesis in roots responds to environmental pH conditions. Under slightly acidic conditions (low pH), plants produce the superior ferric iron (Fe³⁺) reductant sideretin, which complements Fe³⁺ reduction by the membrane-bound reductase FRO2. At neutral to alkaline conditions, fraxetin is synthesized to maintain the mobilization of poorly soluble iron sources for further reduction by FRO2.