WAT1 (WALLS ARE THIN1) defines a novel auxin transporter in plants and integrates auxin signaling in secondary wall formation in Arabidopsis fibers

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Background
Our knowledge of signaling mechanisms involved in secondary cell wall (SCW) formation is quite limited. To discover novel markers of SCW, a genomics approach using Zinnia elegans xylogenic cultures was undertaken that identified hundreds of gene candidates expressed at the onset of secondary wall formation [1]. Arabidopsis homologs and the corresponding T-DNA mutants for each Zinnia gene were identified and the panel of Arabidopsis cell wall mutants was subjected to developmental and wall-related phenotyping.

Results and conclusion
Among the most interesting mutants was wat1 (walls are thin1). The most conspicuous phenotypic feature of wat1 was the severe reduction (sometimes to the extent of being inexistent) of SCW in xylary and interfascicular stem fibers. Interestingly, xylem vessel wall thickness and morphology were not modified by the mutation. In addition to the SCW phenotype, wat1 was characterized by 5-Me-tryptophan seedling toxicity, severely decreased auxin transport and content in stems, and massive down-regulation of auxin-related genes. These data led us to the conclusion that WAT1 acts as an upstream regulator of SCW deposition in fibers, presumably through an auxin-mediated mechanism [2].

Bioinformatic analysis of WAT1, annotated as ‘homo-log to a Medicago truncatula nodulin gene, MtNOD21, suggested that WAT1 encoded a putative transporter belonging to the Plant Metabolite Exporter family [3]. WAT1:GFP fusion protein experiments localized WAT1 on the tonoplast, confirming the prediction that WAT1 is a membrane protein. Although WAT1 is plant-specific, it shares structural similarities with bacterial amino acid transporters in that it consists of ten transmembrane domains encompassed within a tandem Domain of Unknown Function 6 (DUF6).

To characterize WAT1 function, we recently tested its capacity to transport tryptophan and/or auxin in both yeast and Xenopus oocytes. Neither WAT1-expressing yeast cells nor Xenopus oocytes were able to facilitate radiolabeled Trp import or export. However, we have been able to demonstrate that WAT1 facilitates auxin import in both expression systems. These results clearly place WAT1 among the ranks, along with PINs, AUX/LAXs and ABCB/MDR/PGPSs, as a novel, bona fide auxin transporter in plants.

This study constitutes the first functional characterization of any of the 46 members of the WAT1 gene family in Arabidopsis and our hope is that this discovery will help pave the way in identifying the functions of other family members. Moreover, the wat1 mutant will be an ideal tool to address the question as to how auxin subcellular homeostasis plays a role in fiber SCW formation in Arabidopsis. Our current efforts to understand poplar WAT1-mediated auxin signaling in wood formation in trees will also be discussed.
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