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Identifying the transporters of different flavonoids in plants

Elinor P. Thompson,* Julia M. Davies and Beverley J. Glover
Department of Plant Sciences; University of Cambridge; Cambridge, UK

We recently identified a new component of flavonoid transport pathways in *Arabidopsis*. The MATE protein FFT (Flower Flavonoid Transporter) is primarily found in guard cells and seedling roots, and mutation of the transporter results in floral and growth phenotypes. The nature of FFT's substrate requires further exploration but our data suggest that it is a kaempferol diglucoside. Here we discuss potential partner H⁺-ATPases and possible redundancy among the close homologues within the large *Arabidopsis* MATE family.

Plant flavonoids are becoming notorious for their wide and expanding range of possible functions. Beyond UV protection (itself not entirely without debate), further roles have been added in plant development; nodulation and interactions with pathogens; fertilisation; and auxin transport. For such a well-described biochemical network, it is interesting that few aspects of flavonoid function are clear-cut: perhaps it is the recently established link with auxin, so intimately involved in every aspect of plant development, that consigns them to multiple incompletely-known regulatory pathways. Knowledge is lacking, in particular, about the transport of flavonoids. Such transport is necessary¹ and we now know that selective uptake of flavonoids and movement of flavonoids through the plant occur.^{2,3} When naringenin, dihydrokaempferol and dihydroquercetin were added to the *Arabidopsis tt4* mutant [lacking the enzyme chalcone synthase (CHS) and thus all flavonoids] at root tip, mid-root or to cotyledons, they were converted to downstream products. Grafting on flavonoid-producing tissues to *tt4* could also complement the mutation.³

Kitamura⁴ and Buer et al.⁵ speculate that MATE transporters are good candidates to enable flavonoid transport at the membrane, allowing the necessary movement from one membrane system to another. The link between MATE proteins and flavonoid transport is justified by work in tomato⁶ and confirmed by the discovery of TT12.^{7,8} Also conforming to this premise is our recent work on FFT (Flower Flavonoid Transporter), a MATE protein probably situated in the tonoplast membrane that has a role in flavonoid transport in specialised guard cells and anthers.⁹

Auxin and Flavonoids

The phenotype of the *fft-1* mutant adds to the numerous recent descriptions of flavonoids influencing auxin transport. The mutant grew faster than wild type in all conditions we tested. Since *Arabidopsis* seedlings are very small the difference is subtle but was unvarying and statistically significant up to about 2 weeks post-germination, and culminated in *fft-1* flowering earlier than wild type. The most notable effects of the *fft-1* mutation are floral, namely lack of anther dehiscence and reduced pollen viability, but even these cannot be entirely dissociated from an auxin phenotype.¹⁰

We do not have exhaustive data on the substrate for FFT, but LC-MS comparing *fft-1* mutant and Col-0 *A. thaliana* showed significant changes in the level of a kaempferol diglucoside in some floral tissues, among a range of perturbed flavonol glycosides. Although FFT promoter-GUS reporter plants showed strong, specific staining in root tissues, the faster root growth of seedlings remains unexplained, since we did not find differences

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*Correspondence to: Elinor P. Thompson;
Email: et250@cam.ac.uk

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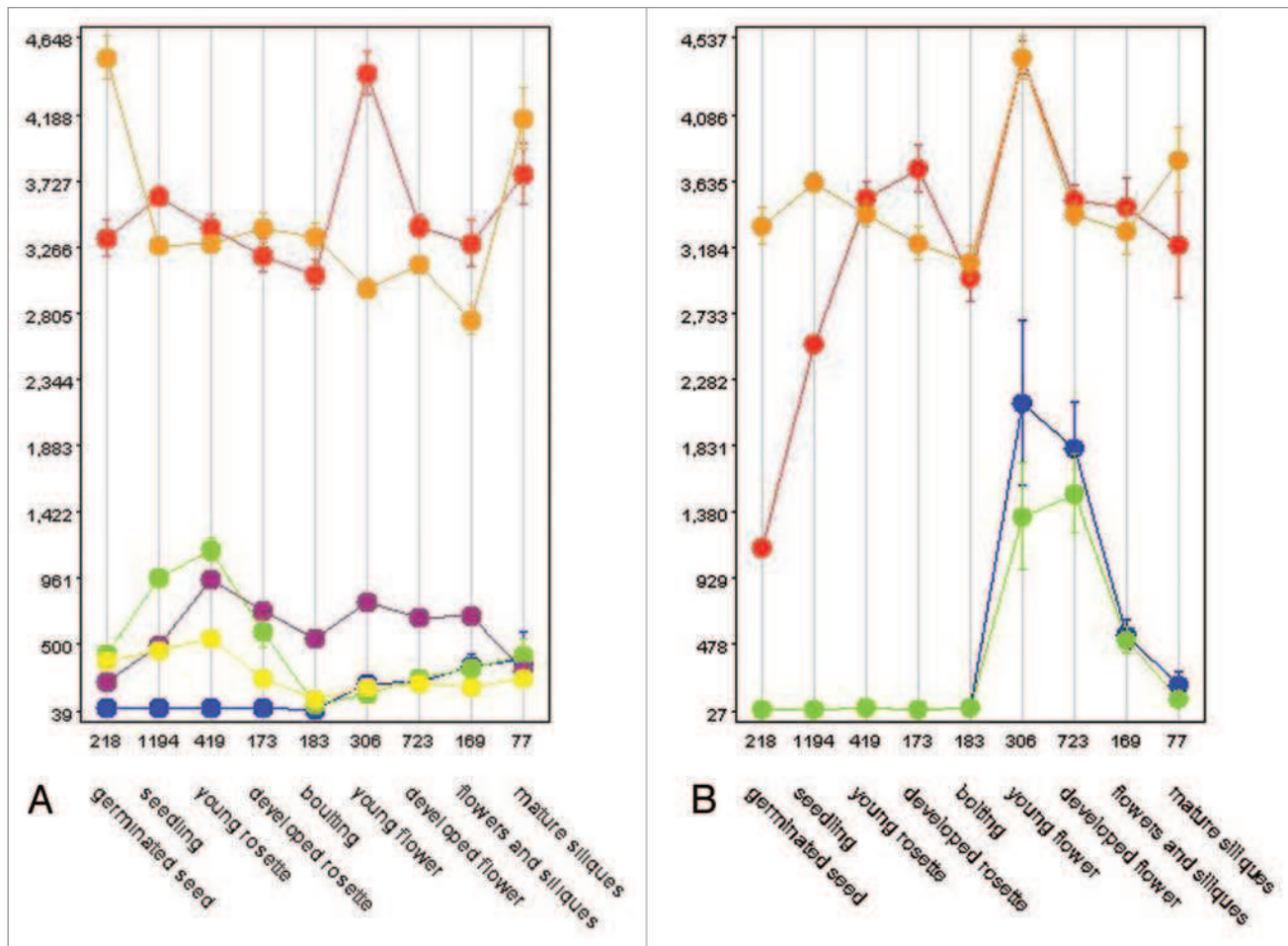


Figure 1. (A) Genevestigator developmental expression profiles of *FFT* (red), *At4g00350/DTX34* (blue), *At5g38030, DTX30* (green), *At1g47530/DTX33* (orange), *At3g26590/DTX29* ORFs (purple and yellow). (B) *FFT* (orange) with ATPases *At5g57350/ATPase 3* (red), *At2g07560/AHA6* (blue) and *At1g80660/AHA9* (green).

in flavonoids outside floral tissues in our analyses. Both roots and floral tissues require careful dissection and visualisation to confirm and quantify better the effects on the various kaempferol and quercetin compounds that were altered in *fft-1*. It would also be interesting to attempt to complement the mutation with a range of flavonoid compounds, and to visualise GUS expression following transformation of *fft-1* plants with a *dr5* auxin reporter construct.

Flavonoids and Pollen

As mentioned above, *fft-1* plants recovered over time, appearing wild type in the mature vegetative phase. Early flowers are profoundly affected by the lesion, however, with anthers frequently nondehiscent and containing little or no viable pollen.

Only low numbers of seeds are produced in siliques on the primary inflorescence. The importance of flavonoids for pollen development and fertility in maize and petunia is well known.^{11,12} In contrast, in a surprising paper in 1996, Ylstra et al. showed that the *tt4* mutant was fertile, with normal pollen development.¹³ Arabidopsis has been seen since as a rare exception among the flowering plants, in being able to produce viable pollen in the apparent absence of flavonoids. Reduced fertility in *fft-1* contradicts the received wisdom on flavonoids in Arabidopsis to some degree, in suggesting that perturbation of flavonoid metabolism is certainly capable of interfering with the plant's fertility. It is possible that the solution lies in the detail, with Arabidopsis having mechanisms to compensate for the absence of flavonoids, but lacking the mechanisms to

adjust to alterations in the types and proportions of flavonoids present.

Homologues

The MATE family of proteins clearly transport diverse compounds, from the organic cation tetraethylammonium by a human family member, antibiotics by prokaryotic forms, through to citrate secretion by *FRD3* in Arabidopsis.¹⁴ Now more whole-genome sequences are being completed, we can see homologues of *FFT* in many other plants. The most closely related proteins currently seen in plants are strikingly similar, the most similar of which are a 75% identical *Ricinus communis* putative multidrug resistance protein (*EEF49069*); the 72% identical *Solanum lycopersicum* putative anthocyanin permease discussed by Matthews et al. (2003)

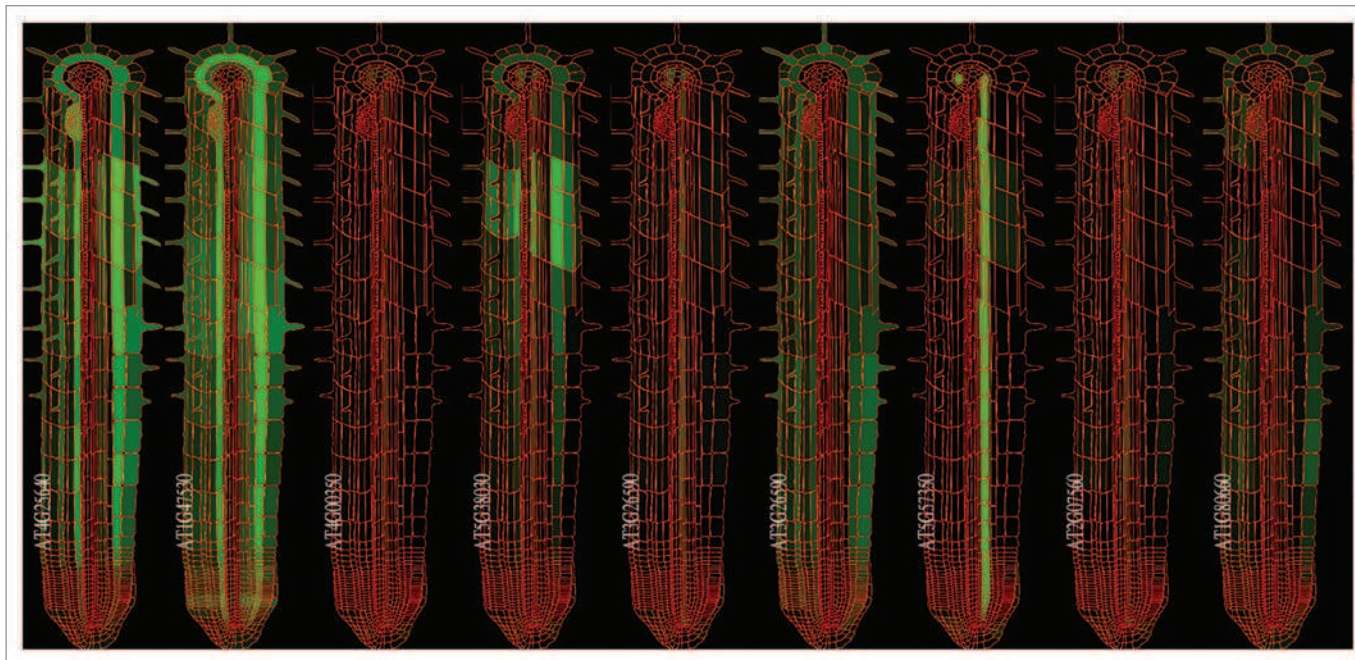


Figure 2. Arexldb expression profiles in roots of *FFT* (At4g25640), At4g00350/*DTX34*, At5g38030/*DTX30*, At1g47530/*DTX33*, At3g26590/*DTX29* MATE proteins with ATPases At5g57350/*ATPase 3*, At2g07560/*AHA6* and At1g80660/*AHA9*.

and in our paper;⁹ and a 70% identical *Vitis vinifera* hypothetical protein. Others are now seen in *Populus trichocarpa* and *Oryza sativa*. The most similar *A. thaliana* proteins to *FFT* (*DTX35*) are 67% identical (At4g00350; *DTX34*), followed by 51% (At1g47530, *DTX33*; tagged as a “ripening responsive protein”), 50% (At5g38030, *DTX30*), and 48% identical (At3g26590, *DTX29*) family members, but these are part of a large family of 50+ putative proteins, as often seen in this species (for review, see Yazaki, 2005).¹⁵

Redundancy

At1g47530/*DTX33* and *FFT* transcription profiles are similar in collected microarrays. The microarray data showing widespread transcription for *FFT* in the profiling application, Genevestigator,¹⁶ correspond with our RT-PCR, namely in the lateral root and root elongation zone; senescent and mature leaves; and floral tissues especially sepal, petal and siliques (Fig. 1). Although alike in many microarrays and throughout development, where *FFT* has a peak in the young flower, At1g47530/*DTX33* peaks in the developing seedling and appears, in this computer-based analysis, to be more highly

expressed in roots. In the Arabidopsis gene expression database (www.arexldb.org¹⁷) At1g47530/*DTX33* indeed shows very strong expression in the root cortex (Fig. 2).

Interestingly, the most similar Arabidopsis MATE to *FFT* at sequence level, At4g00350/*DTX34*, has a large peak of expression in pollen, contrasting both with the *FFT* bioinformatics and our experimental data, neither of which show *FFT* transcript in pollen. At4g00350/*DTX34* is described in TAIR (www.arabidopsis.org/index.jsp) as being found in mature and germinated pollen, petal differentiation and expansion stage, anthesis, and the globular stage embryo. In general, bioinformatics data suggest that At4g00350/*DTX34* expression resembles that of the other close homologue At3g26590/*DTX29*, although the latter is missing the peak in pollen. These possible transcriptional similarities are interesting when thinking of the potential redundancy seen in the *fft-1* phenotype and might guide future work. Notably, the expression patterns of many of this group of similar proteins seem to feature

in anthesis, corresponding with the phenotype of *fft-1*.

Flavonoids in Guard Cells and the Anther

A GUS-promoter construct showed that guard cells are the likely principal location of the *FFT* transcript in aerial tissues. This is not surprising since much of a plant’s phenylpropanoid and flavonoid content is found in the vacuoles of guard cells, which are central to UV protection and environmental regulation.^{18,19} The presence of *FFT* transcript in the hydathode, anther and nectaries is notable—are special roles required of flavonoids in these specialised guard cells? Of interest here too is the proposal that a partner for the TT12 MATE is the proton pumping-ATPase AHA10 (Arabidopsis H⁺-ATPase), which creates the gradient for active transport by TT12.²⁰ Notably, there is a similar H⁺-ATPase, AHA9, that is expressed exclusively in anther tissues.²¹ Whether this forms an equivalent system in anthers with *FFT*, like that of TT12/AHA10 in seedcoats, requires investigation.

We therefore paired *FFT* with *AHA* transcripts in Genevestigator, to find the most similar profile to that of *FFT* (Fig.

1). At1g80660/*AHA9* and *FFT* both display peaks of expression that correspond with development of the young flower, also matched by At5g57350/*ATPase 3* and At2g07560/*AHA6*. Arexldb shows a different picture in root transcription (Fig. 2).

It would be of interest to further analyse the temporal expression of *FFT* in the floral organs to discover any synchronization with anthesis. *FFT* transcription may also be affected by transcription factors such as CRABS CLAW which are required for anther development.²² More technically challenging would be finding which flavonoids occur in the anther endothecium (thus concerned with dehiscence), or the tapetum (i.e., compounds needed for microsporogenesis and pollen development). Any overlap in expression between different MATE proteins may allow redundancy in some cell types but not in others; for example, *fft-1*'s most profound phenotype is in the anther although its transcript is also found in guard cells in the vegetative tissue of young seedlings, where the mutation did not have any obvious effect (besides the subtle changes in germination and growth at this stage).

Overall, although much work remains to be done to establish the details of flavonoid transport in particular tissues, the transport of specific flavonoids by different transporters can be seen to add another

level of control to the complex pathways through which plant metabolites influence developmental programmes.

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