Overlapping expression patterns and differential transcript levels of phosphate transporter genes in arbuscular mycorrhizal, P$_i$-fertilised and phytohormone-treated *Medicago truncatula* roots

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**Abstract** A microarray carrying 5,648 probes of *Medicago truncatula* root-expressed genes was screened in order to identify those that are specifically regulated by the arbuscular mycorrhizal (AM) fungus *Gigaspora rosea*, by P$_i$ fertilisation or by the phytohormones abscisic acid and jasmonic acid. Amongst the identified genes, 21% showed a common induction and 31% a common repression between roots fertilised with P$_i$ or inoculated with the AM fungus *G. rosea*, while there was no obvious overlap in the expression patterns between mycorrhizal and phytohormone-treated roots. Expression patterns were further studied by comparing the results with published data obtained from roots colonised by the AM fungi *Glomus mosseae* and *Glomus intraradices*, but only very few genes were identified as being commonly regulated by all three AM fungi. Analysis of P$_i$ concentrations in plants colonised by either of the three AM fungi revealed that this could be due to the higher P$_i$ levels in plants inoculated by *G. rosea* compared with the other two fungi, explaining that numerous genes are commonly regulated by the interaction with *G. rosea* and by phosphate. Differential gene expression in roots inoculated with the three AM fungi was further studied by expression analyses of six genes from the phosphate transporter gene family in *M. truncatula*. While *MtPT4* was induced by all three fungi, the other five genes showed different degrees of repression mirroring the functional differences in phosphate nutrition by *G. rosea*, *G. mosseae* and *G. intraradices*.

**Keywords** Abscisic acid · Arbuscular mycorrhiza · *Gigaspora rosea* · *Glomus intraradices* · *Glomus mosseae* · Jasmonic acid · Phosphate transporter genes

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Abbreviations
ABA Abscisic acid
AM Arbuscular mycorrhizal
Gi Glomus intraradices
Gm Glomus mosseae
Gr Gigaspora rosea
JA Jasmonic acid
P_i Inorganic phosphate

Introduction

Fungi of the phylum Glomeromycota form mutualistic symbioses with 80% of the vascular land plants, called arbuscular mycorrhizas (Newman and Reddell 1987; Schüller et al. 2001). This widespread interaction is in the first instance characterised by nutrient exchange between the two partners (Smith and Read 1997). Arbuscular mycorrhizal (AM) fungi deliver mineral nutrients to the plant and obtain as heterotrophic organisms carbohydrates from their autotrophic partner (Smith and Read 1997). This is the general concept of the mycorrhizal symbiosis, but the extent of exchange can vary depending on fungal and plant species (Smith et al. 2003; Lerat et al. 2003). In addition to being biofertilisers, AM fungi possess bioregulating activities by changing the content and distribution of phytohormones (Ludwig-Müller 2000; Hause et al. 2007). This has been shown in different plant AM fungus combinations, but the role of AM fungi as bioregulators was only comprehensively analysed in the case of jasmonic acid (JA) and abscisic acid (ABA). Both, JA and ABA, are important for mycorrhizal establishment as shown by a reverse genetics approach diminishing JA biosynthesis (Isayenkov et al. 2005) or by use of an ABA-deficient mutant (Herrera-Medina et al. 2007), respectively. Moreover, for JA it has been demonstrated that the up-regulation of biosynthesis takes place in arbuscule-containing plant cells of M. truncatula (Isayenkov et al. 2005). The third important function of the AM symbiosis is based on its influence on plant responses to their biotic and abiotic environment. In most cases, mycorrhizal roots show enhanced resistance to pathogens (Borowicz 2001) and AM fungi confer tolerance to the plant under adverse soil conditions such as drought (Ruiz-Lozano 2003) or heavy metal contamination (Gohre and Paszkowski 2006).

For more than a decade, the AM symbiosis has also been a subject of molecular studies. Expression analyses using targeted and non-targeted approaches were carried out to gain insights into the processes underlying the development and the different functions of the symbiosis (Balestrini and Lanfranco 2006). A number of plant genes were identified which were known from other interactions showing similarities between the colonisation of the plant by mycorrhizal fungi, by pathogens or nodule-forming rhizobia (e.g. Gianinazzi-Pearson et al. 1996; van Rijn et al. 1997). This obvious overlap has not only been used to discuss the evolution of plant–microbe interactions (Parniske 2000), but also to explain the role of AM fungi as bioprotectors (Grunwald et al. 2004; Li et al. 2006). However, most of the studies were conducted to understand the molecular basis of nutrient transport from the fungus to the plant. In this respect, mycorrhiza-regulated phosphate transporter (PT) genes were isolated and analysed in different plants (Rausch et al. 2001; Harrison et al. 2002). Interestingly, in the legumes Lotus japonicus and Medicago truncatula down-regulation or mutations of LjPT3 and MtPT4 led to a reduction in mycorrhization (Maeda et al. 2006; Javot et al. 2007), while such a phenotype could not be observed with LePT4 indicating functional redundancy of PT genes in tomato (Nagy et al. 2005).

A broad overview of gene expression patterns in mycorrhiza has been obtained from large array analyses in the model plant M. truncatula and Oryza sativa. AM fungi of the genus Glomus were used in most cases and the results were compared with expression profiles in roots after fertilisation with inorganic phosphate (P_i), in nodules or during pathogenic interactions (Liu et al. 2003; Manthey et al. 2004; Hohnjec et al. 2005; Guimil et al. 2005). In order to extend knowledge about gene regulation in the AM symbiosis, the analysis for the present paper was extended to the interaction between Gigaspora rosea and M. truncatula. The fungus G. rosea differs in various aspects from Glomus species. It contains no RNA during dormancy, it only infects the plant through germinating spores, it reacts differently to compounds occurring in root exudates and it forms auxiliary cells (Franken and George 2006). However, diversity of the AM symbiosis has not only been observed for morphological aspects. AM fungi also seem to differ in the amounts of phosphate transferred to the plant (Smith et al. 2004) and in their response to phosphate enrichments in the soil (Cavagnaro et al. 2005). To gain a deeper insight into these mechanisms, the expression of the M. truncatula phosphate transporter gene family (five known genes and one newly identified) was analysed in roots colonised with the three AM fungi G. mosseae, G. intraradices and G. rosea. In addition, transcript levels were monitored not only after P_i fertilisation, but also after treatment of the roots with ABA or JA, in order to identify genes which are not directly regulated by the AM fungus, but indirectly by changes in the levels of these phytohormones. The results were compared with those obtained in this study for G. rosea and with results published for two Glomus species (Hohnjec et al. 2005).
Materials and methods

Plant growth and inoculation

*Medicago truncatula* cv. Jemalong A17 (seeds obtained from Thierry Huguet, CNRS-INRA Toulouse, France) was cultivated as described by Wulf et al. (2003). Briefly, seeds were subjected to a 10 min treatment with sulphuric acid and three times washing in sterile water, followed by 3 min sterilization in sodium hypochlorite (1.6%, w/v) and a final washing step in sterile water. Surface-sterilised seeds were germinated on wet filter paper in a Petri dish in the dark for 3 days at 4°C and for additional 2–4 days at RT. Seedlings were transferred to pots containing 0.7 l of a 1:1 mixture of vermiculite and expanded clay and grown at 25°C with 16 h light and 65% humidity. Plants were fertilised every 4 days with 10 ml Long Ashton nutrient solution (Hewitt 1966) containing reduced P concentration (0.13 mM).

In order to obtain mycorrhizal roots, inoculation was carried out by 1:10 mixing of commercial available inoculum of *Gigaspora rosea* BEG9, *Glomus mosseae* BEG12 or *Glomus intraradices* BEG141 (Biorize Sarl, Dijon, France) with the substrate when transplanting to pots. Plants were harvested 6 weeks after inoculation and mycorrhiza parameters were estimated after trypan blue staining according to the method of Trouvelot et al. (1986). For the +P treatment, plants were fertilised during the whole growth period with the fertiliser solution containing 1.3 mM P<sub>i</sub>. In order to challenge the plants with the phytohormones, 100 μM (+)-cis, trans-abscisic acid in 10 ml 2.5% acetone or 45 μM jasmonic acid in the 0.13 mM P<sub>i</sub> fertiliser solution were applied to the growth substrate 4 h before harvest. Control plants received the same amount of the corresponding solvents in the 0.13 mM P<sub>i</sub> fertiliser solution.

Array hybridisation

Plant material was frozen and grounded in liquid nitrogen. For each RNA extraction, 100 mg of grounded plant material was processed with the RNaseasy kit according to the protocol of the supplier (Qiagen, Hilden, Germany). DNA was removed by DNase I treatment (Bauer et al. 1993) and RNA quality and quantity was estimated with the LabChip-System RNA 600 Nano kit in the Bioanalyzer 2100 (Agilent Technologies, Boeblingen, Germany). Mycorrhiza-regulated gene expression was confirmed by cDNA synthesis (procedure see below) and non-quantitative PCR using primer pairs for *MtPT4* (Table S1 in Supplementary material) and *MtHA1* (Krajinski et al. 2002). For hybridisation of the M6okRIT macro- and microarrays, cDNA probes were synthesised from two biological replicates with each two technical repetitions and labelled with 33P, Cy3- or Cy5-dUTP (Amersham, Freiburg, Germany), respectively (Küster et al. 2004). Hybridisation signals were obtained using the Storm Phosphorimager (Molecular Dynamics, Ismaning, Germany) equipped with the ImageQuant software or by the ScanArray 4000 microarray scanner (Perkin-Elmer, Boston, MA, USA), and the data were statistically analysed using the EMMA 1.0 software (Dondrup et al. 2003).

PCR cloning

mRNA sequences from the phosphate transporter genes *MtPT1-5* were obtained from the NCBI database (AF000354, AF000355, EF016358, AY116210, EF016359). The gene *MtPT6* was deduced from two cDNA sequences (TC99939, BQ145041) and one genomic sequence (clone MTH2-177C12 on chromosome 3). Using the primers MtPt6for, MtPt6rev and MtPt6-5′ (Table S1 in Supplementary material), cDNA fragments were amplified, sequence analysed and the complete coding sequence was assembled. The whole *MtPT6* cDNA sequence has been submitted to the EMBL database and can be obtained under the AC number AM982728. Amino acid sequences were deduced from the coding regions and aligned with a selection of phosphate transporters from other plants and from the AM fungus *Glomus versiforme* as outgroup by ClustalW (Thompson et al. 1994). The amino acid alignment was phylogenetically analysed by the program PUZZLE (version 4.0.2; Strimmer and von Haeseler 1996) and the results were displayed as dendrogram with the program TREEVIEW (version 1.5.3; Page 1996).

Real-time PCR

RNA was extracted and DNase treated as described above. cDNA was synthesised from 4 μg total RNA in a volume of 20 μl using oligo(dT) following the protocol of the MMLV-reverse transcriptase supplier (Promega, Mannheim, Germany). The absence of contamination with genomic DNA was confirmed by PCR using primer pairs spanning an intron sequence as described (Wulf et al. 2003). Real-time PCR reactions were carried out in 25 μl volume using 2 μl of cDNA (1:10 diluted, three technical repetitions) as template, 0.15 μM primer pairs of the *MtPT* genes (Table S1 in Supplementary material) or of the *MtTef1* gene (Wulf et al. 2003) and the QuantiTect™ SYBR® Green PCR Kit (Qiagen, Hilden, Germany). For the standard curve, a dilution series of 10<sup>8</sup> to 10 molecules of the different gene fragments was prepared by PCR and checked by agarose gel electrophoresis. Detection of the amplified fragments and recording of the melting curve was done on an iCycler (Bio-Rad Laboratories, München, Germany) with 15 min at 95°C and 45 cycles (15 s at 95°C; 30 s at 56°C; 30 s at 72°C) followed by heating from 56 to 95°C with a rate of...
Overview of array screening results

Concentration of P_i

For soluble P_i concentration measurements, 100 mg fresh shoot material (four biological repetitions) were sampled and macerated in 1.2 ml extraction buffer (45 mmol/l sodium acetate buffer, pH 5.0). The suspension was centrifuged at 12,000 g for 30 min at 4°C and P_i concentration was determined by carrying out the phosphorus-molybdate blue colour reaction and measuring absorbance at 700 nm (Murphy and Riley 1962).

Results and discussion

Gigaspora rosea-regulated transcript levels

Medicago truncatula roots inoculated with the AM fungus Gigaspora rosea and harvested 6 weeks after inoculation showed in two independent experiments a mycorrhizal colonisation degree of 37 and 41% and a relative arbuscule frequency of 80 and 82%, respectively. The arbuscules resembled the Arum-type. This pattern has been already described before in the interaction between M. truncatula and G. rosea, but is different from the type found in tomato-G. rosea mycorrhizas (Burleigh et al. 2002). Non-quantitative RT-PCR showed that the AM-specific genes MtHA1 (Krajinski et al. 2002) and MtPT4 (Harrison et al. 2002) were clearly expressed in the roots colonised by G. rosea, but not in the controls (data not shown). Screening of the Mt6kRIT microarray with cDNA probes from mycorrhizas and controls resulted in significant expression values for 1,300 genes (Table 1). For 118 genes a significant up-regulation between 2- and 20-fold could be detected, while 222 were up to 25-fold repressed (Table S2a in Supplementary material). This was much less than the number of Glomus intraradices-regulated genes identified amongst the same set of cDNA fragments (Manthey et al. 2004), but in that study, P < 0.075 and M ≥ 0.7 or ≤ −0.7 were accepted as being significant. Based on the same values, only slightly higher percentages between 12 and 20% were obtained in another experiment where the regulation by the AM fungi Glomus mosseae and G. intraradices was analysed on Mt16kOLI1 microarrays of M. truncatula (Hohnjec et al. 2005). Far fewer genes (4%) were detected in an experiment with Glomus versiforme, although inserts of a cDNA library from M. truncatula mycorrhizal roots were hybridised (Liu et al. 2003). Significant signals for the AM-specific genes MtPT4 and MtHA1 could not be detected on the arrays, although they are induced in G. rosea-colonised roots. This has been also observed in other array hybridisation experiments (Liu et al. 2003, 2007) and is probably due to the low overall expression of these genes.

Data for G. rosea from the analysis presented here were compared with those for G. mosseae and G. intraradices previously published (Hohnjec et al. 2005). This revealed 163 genes with significant results for all three fungi (Table S3 in Supplementary material), whereof 57 genes are regulated by G. rosea (Table 2). Amongst the 163 genes, which could be compared between the three fungi, 7% showed an overlapping pattern of regulation between G. rosea- and G. mosseae- and 9% between G. rosea- and G. intraradices-colonised roots, but 22% were similarly up- or down-regulated in mycorrhiza of all three AM fungi are of unknown function but one putatively encodes an alpha-xylosidase precursor (Table 2). Interestingly, a common regulation pattern for Glomus- and Gigaspora-mycorrhizal roots has been shown for a xyloglucan endotransglycosylase gene, which is also involved in modification of xyloglucan polymers (Maldonado-Mendoza et al. 2005). A more recent analysis compared expression patterns in M. truncatula mycorrhiza with a Glomus and a Gigaspora species using the Mt16kOLI1 and 10% of the G. intraradices-induced genes were also regulated in roots colonised by Gigaspora gigantea (Liu et al. 2007). This is more close to the 9% we have found between G. rosea and G. intraradices than to the 22% identified as overlapping between the two Glomus species. Much more genes with overlapping expression were found in an investigation, where 10 out of 14 selected genes showed a common induction in M. truncatula mycorrhiza formed by different AM fungal isolates (Massoumou et al. 2007). However, the genes were pre-selected as mycorrhiza-regulated, and the

| Samples | N genes (P < 0.05) | N induced genes (M value) | N repressed genes (M value) |
|---------|-------------------|--------------------------|---------------------------|
| G. rosea | 1,300 | 118 (4.31 ≥ M) | 222 (−4.64 ≤ M) |
| ABA | 1,202 | 121 (3.32 ≥ M) | 205 (−4.17 ≤ M) |
| JA | 433 | 193 (4.68 ≥ M) | 71 (−4.29 ≤ M) |
| P_i | 1,221 | 139 (3.9 ≥ M) | 143 (−4.65 ≤ M) |

The number of genes which gave significant results after hybridisation of different samples versus control is shown.

The numbers of at least two fold induced or repressed genes (log2 ratio $M \geq 1$ and $\leq -1$) and the maximum and minimum $M$ values are indicated.
### Table 2
Clusters, annotations and $M$ values of *G. rosea*-regulated genes

| Cluster          | Description                                      | Gr  | Gm  | Gi  |
|------------------|--------------------------------------------------|-----|-----|-----|
| **Cell wall**    |                                                   |     |     |     |
| MtC45430         | Alpha-xylosidase precursor                       | 1.95| 1.11| 1.47|
| MtC45398         | Endo-beta-1-4-D-glucanase                        | −1.18| 0.96| 0.63|
| MtC00798         | Arabinogalactan-protein precursor                | −1.33| −1.01| −0.38|
| **Cytoskeleton** |                                                   |     |     |     |
| MtC00684         | Tubulin/ftsz protein                             | −1.28| 0.90| 1.37|
| MtC20031         | Tubulin/ftsz protein                             | −1.30| 0.76| 1.14|
| **Membrane transport** |                                           |     |     |     |
| MtC20051.1       | Envelope calcium ATPase                          | −2.29| 1.66| 0.51|
| MtC20233         | Vacuolar ATP synthase                            | −2.87| −0.61| −0.43|
| **Vesicular trafficking** |                                         |     |     |     |
| MtC93021.1       | Importin alpha subunit (NLS receptor)            | 1.23| 0.61| 0.75|
| **Primary metabolism** |                                             |     |     |     |
| MtC90582         | Albumin 2                                        | −1.92| 0.92| 1.44|
| MtC00680.1       | Narbonin                                         | 1.57| 1.81| 1.30|
| MtC00519.2       | NADH-ubiquinone oxidoreductase                   | −1.32| 0.32| 1.16|
| MtC50697         | Blue copper protein                              | 1.74| 0.82| 0.75|
| MtC30172         | Biotin carboxyl carrier protein                  | −3.15| 0.52| 0.50|
| MtC00606         | Epimerase/dehydratase                            | −1.69| −0.47| −0.33|
| MtC93062         | NADH-plastoquinone oxidoreductase               | −2.50| −0.46| −0.65|
| **Secondary metabolism** |                                         |     |     |     |
| MtC50990         | Cytochrome P450                                  | −2.03| −0.37| 1.17|
| MtC10905         | Squalene synthase                                | 1.95| 0.20| 0.41|
| **Chromatin and DNA metabolism** |                                   |     |     |     |
| MtC10278         | Histone H2b                                      | −1.50| 0.86| 1.12|
| **Gene expression and RNA metabolism** |                                               |     |     |     |
| MtC40096         | Protein disulphide isomerase                     | −2.80| 0.88| 1.77|
| MtC30151         | WRKY-type DNA binding protein                    | −4.17| 0.66| 0.48|
| MtC92170         | MYB-related transcription activator              | 1.40| 1.85| −1.01|
| **Protein synthesis and processing** |                                             |     |     |     |
| MtC00024         | 40S ribosomal protein S11                        | 3.34| 0.59| 1.38|
| MtC10400         | Chaperonin CPN10                                 | −1.69| 0.66| 1.20|
| MtC93070         | Subtilisin-like serine protease                  | 1.01| 0.99| 0.42|
| MtC00760         | Ubiquitin-conjugating enzyme                     | −1.69| −0.82| −0.36|
| MtC40177         | Serine protease                                  | −1.56| 0.51| −0.61|
| **Signal transduction** |                                         |     |     |     |
| MtC00080         | Calmodulin                                       | −2.26| −0.37| −0.58|
| MtC20202         | Ser/Thr protein phosphatase                      | 1.60| −0.42| −0.65|
| **Miscellaneous** |                                                   |     |     |     |
| MtC10112         | Peroxidase precursor                             | −2.75| 1.21| 1.27|
| MtC40023.1       | Peroxidase                                       | 0.84| 0.50| 1.14|
| MtC50998         | C2 domain-containing protein                     | −1.08| −0.64| −1.51|
| MtC00724.1       | Peroxidase                                       | −1.04| −0.58| −2.17|
| MtC10717         | Peroxidase precursor                             | −1.32| 0.66| 1.47|
| MtC00252.2       | Stress-inducible protein                         | −1.22| 0.62| 0.54|
| MtC00227         | PR10                                             | −1.67| −0.19| −1.75|
| MtC10312         | Chitinase                                        | −2.67| −2.10| −2.58|
| MtC30250         | Stress related protein                           | −1.15| −1.01| −0.73|
| MtC00640.1       | Germin-like protein (oxalate oxidase)            | 1.02| −0.78| −1.43|
analysis was carried out using the sensitive semiquantitative RT-PCR technique. For none of the genes, data were obtained in the present analysis or in the analysis by Hohnjec et al. (2005). In another analysis using M. truncatula mycorrhizal roots colonised with the AM fungi G. intraradices, G. mosseae and Scutellospora castanea (Feddermann et al. 2008) six out of ten mycorrhiza-regulated genes were only induced by S. castanea, but not by the two Glomus species. It is difficult to compare the analyses amongst each other, because different environmental conditions influence the outcome of the experiments. Nevertheless, general expression patterns suggest that the phylogenetic distance between AM fungal isolates is not only restricted to the morphology of spores, the mode of colonisation or the amount of phosphate which is transported from the fungus to the plant (for review, see Franken and George 2006), but is also mirrored by the plant genes which are regulated during mycorrhization.

Phytohormone-regulated transcript levels

Alterations in the homeostasis of plant hormones have been implicated in the process of mycorrhization. There are numerous reports showing that the levels of plant hormones such as cytokinin, JA, auxin, auxin-related compounds and ABA actually change upon the establishment of AM (for review, see Hause et al. 2007). Regarding JA and ABA, both phytohormones are well known for its important signalling and regulatory roles that enable plants to survive a variety of abiotic and biotic stresses (Finkelstein and Rock 2002; Wasternack 2007). To assess whether or not mycorrhizal-regulated genes could also be induced or repressed by phytohormones, non-inoculated plants were challenged 4 h before root sampling for RNA extraction with ABA or JA. cDNA probes synthesised from this RNA were used to hybridise Mt6kRIT arrays. Based on the same parameters as described above, 121 of 1,202 genes were induced by ABA and 205 were repressed (Table 1). Similar analyses have been carried out for example in rice and in sugarcane (Rabbani et al. 2003; Rocha et al. 2007) and numerous genes identified as responsive to abiotic stress in different plants were shown to be also regulated by ABA (for review, see Yamaguchi-Shinozaki and Shinozaki 2006). Such stress-regulated genes were found amongst the sequences showing an increased hybridisation signal on the array, but the number was not particularly high (Table S2b in Supplementary material). Comparing the ABA and G. rosea-regulated expression patterns, the overlap was very small (Table 3). This indicates that the increase of this phytohormone during mycorrhization is too low for a regulation of a significant number of genes or that the response is restricted to a very few cells. Nevertheless, ABA is necessary for the proper formation of arbuscules and for a sustained colonisation of the plant root as shown by the

Table 2 continued

| Cluster   | Description                  | Gr  | Gm  | Gi  |
|-----------|------------------------------|-----|-----|-----|
| Unknown function |                              |     |     |     |
| MtC00311   | –                            | 1.24| 4.95| 6.42|
| MtC93285   | Growth-related protein-like   | 1.44| 5.85| 6.21|
| MtC10058   | USP domain                   | 1.11| 1.05| 2.51|
| MtC93028   | Leucine-rich repeat           | 2.44| 1.73| 1.83|
| MtC20198   | –                            | −1.49| 0.61| 1.73|
| MtC10256   | –                            | −1.75| 0.64| 1.24|
| MtC40200   | Methyltransferase            | −2.74| 0.82| 0.96|
| MtC20055   | Tumour differentially expressed protein | −1.82| 0.48| 0.43|
| MtC00686   | Asp/Glu rich protein         | −1.77| −0.35| 0.36|
| MtC45607   | Basic helix-loop-helix domain | 3.44| −0.55| −0.31|
| MtC30081   | Phosphodiesterase            | −4.62| 0.58| −0.80|
| MtC10195   | Lys/Glu/Ala rich protein     | −1.64| −0.49| −1.02|
| MtC30420   | –                            | −1.15| −2.16| −2.15|
| MtC10662   | Lipoxygenase Lh2 domain      | 3.65| −1.49| −2.24|
| MtC00372.1 | –                            | −2.21| −0.63| −3.22|
| MtC50619   | Aldose 1-epimerase           | −1.54| −0.85| −3.97|
| No homology |                              |     |     |     |
| MtC91938   | –                            | −1.42| 0.52| 1.06|
| MtC91288   | –                            | 2.00| −0.77| −1.08|
| MtC00275   | –                            | 2.11| −1.09| −1.53|

Shown are only those genes where data for the AM fungi G. mosseae (Gm) and G. intraradices (Gi) are available (Hohnjec et al. 2005). M values >1 or < −1 (more than two times regulated) are indicated in bold.
Table 3 Overlapping expression patterns

|        | ABA+ | ABA- | JA+ | JA- | P+  | P-  |
|--------|------|------|-----|-----|-----|-----|
| Gr+    | 2%   | 3%   | 1%  | 0%  | 21% | 5%  |
| Gr-    | 4%   | 3%   | 6%  | 13% | 1%  | 31% |
| ABA+   | 2%   | 4%   | 3%  | 2%  |     |     |
| ABA-   | 5%   | 7%   | 3%  | 4%  |     |     |
| JA+    | 3%   |     |     |     |     |     |
| JA-    | 3%   |     |     |     |     |     |

The table shows the percentage of gene numbers with identical regulation (induced or repressed) to the total gene numbers with significant results in two treatments. + and – mean significantly induced or repressed by the particular treatment.

Analysis of AM colonisation of the tomato mutant *situs*, in which ABA levels are drastically reduced in comparison to wild-type tomato plants (Herrera-Medina et al. 2007). Interestingly, the present analysis revealed that two leghemoglobin-encoding genes showed induced transcript levels by *G. rosea* and ABA (Table S4 in Supplementary material). It has been suggested that leghemoglobins play a role in detoxification in mycorrhizal and nodulated roots as well as also under non-symbiotic conditions by binding nitric oxide (Vieweg et al. 2005). Since nitric oxide is involved in ABA signalling in stress responses (Ruan et al. 2004), leghemoglobin expression could be part of an auto-regulatory circuit involved in the fine-tuning of the system during ABA response.

In the case of JA, Mt6kRIT macroarrays were screened and significant data were obtained for 433 genes, of which 193 seemed to be up- and 71 down-regulated (Table 1). When compared with the other treatments, a significant fraction of the induced genes putatively encodes components of secondary metabolism and stress response (Table S2c in Supplementary material). This is in accordance with the proposed function of jasmonates in the plant reaction to various stress factors (Wasternack and Hause 2002) and corresponds to data obtained from a transcriptional analysis of sugarcane roots treated with JA methyl ester (Bower et al. 2005). Stress response was also the category, where the expression patterns in mycorrhiza mostly overlapped with those in JA treated roots (Table S4 in Supplementary material). This suggests at least some of the genes involved in this reaction in mycorrhizal roots could be indirectly regulated by changes in the JA level.

P$_i$-regulated transcript levels

Mycorrhizal fungi are generally able to increase the P$_i$ level in plants (Bucher 2007). Additional array hybridisation experiments have therefore been carried out, in order to distinguish genes directly regulated by the AM fungus from those, which could be induced or repressed indirectly by the improved phosphate nutrition. Plants were grown in parallel to those mentioned above, but were fertilised with sufficient amounts of P$_i$. RNA was extracted and used for probe synthesis and hybridisation. Amongst the 1,221 genes with statistically significant results (Table 1), 139 showed a greater than two times induction by P$_i$ and 143 were repressed (Table S2d in Supplementary material). This was less than has been observed on Mt16kOLI1 microarrays (26 and 16%; Hohnjec et al. 2005), but much more than amongst the genes identified in a mycorrhizal cDNA library (0.7 and 0.5%; Liu et al. 2003). Comparison of the expression patterns revealed that amongst 218 genes, where significant data were obtained for both treatments, 21 or 31% were similarly induced or repressed in *G. rosea*-colonised and P$_i$-fertilised roots (Table 3). This was relatively high and contrasts the overlapping expression values of 4 or 9% found between roots fertilised with P$_i$ and roots inoculated with *G. mosseae* or *G. intraradices*, respectively (Hohnjec et al. 2005).

An additional experiment was carried out to analyse, whether this difference in gene expression patterns could be based on different symbiotic activities of the three AM fungi. *M. truncatula* seedlings were inoculated in parallel with *G. mosseae*, *G. intraradices* or *G. rosea* or were mock-inoculated and grown under phosphate-limiting conditions. When the mycorrhizal symbioses were fully established (60–80% colonisation), soluble P$_i$ concentrations were measured in the shoots, since they are discussed to be mainly involved in the regulation of genes in the roots (Franco-Zorrilla et al. 2004). Comparing the P$_i$ concentrations in the mycorrhizal plants with the corresponding controls revealed that they were not much different in case of the two *Glomus* species, but showed a two-fold increase in *G. rosea* inoculated plants (Fig. 1a). This might be due to the fact that in plants interacting with *Glomus* species increased phosphate uptake was converted to biomass production, while in *G. rosea* plants this was not the case. In contrast these plants showed a slightly lower biomass than the corresponding controls (Fig. 1b) so that the same phosphate uptake resulted in the higher P$_i$ concentration. Such different mycorrhizal effects on *M. truncatula* biomass were observed before (Smith et al. 2003) and are probably based on the variations in carbohydrate drain of AM fungi in pot cultures, because they need different amounts of energy for building up their extraradical mycelium (Lerat et al. 2003). A recent study showed that *Gigaspora* species in general lead to higher P$_i$ concentrations in leaves of *Plantago lanceolata* than *Glomus* species do (Maherali and Klironomos 2007). In the present experiment, the higher P$_i$ concentrations in the shoots of *G. rosea*-colonised plants can explain the high overlap in the expression pattern with those plants which were fertilised with P$_i$ and the differences to the results obtained before with *Glomus* species.
Transcript levels of MtPT genes

To analyse whether patterns of overlapping Pi- and mycorrhiza-regulated transcript levels can be extended to genes directly involved in Pi nutrition of the plants, databases were screened for cDNA sequences with homologies for phosphate transporter genes. Apart from MtPT1, MtPT2 and MtPT4, which are known from previous studies (Liu et al. 1998; Harrison et al. 2002), a number of other putative PT-encoding genes were identified. Primers were designed based on sequence alignment and used for a first non-quantitative RT-PCR experiment. Four of these putative genes showed RNA levels in mycorrhizal roots (data not shown). Full length cDNA sequences were available for MtPT3 and MtPT5 and the characterisation of these two genes was recently published (Liu et al. 2008). The other two putative genes could only be deduced from different cDNA and genomic sequences in the database. Further PCR cloning experiments revealed a sixth gene (MtPT6), while one tentative consensus (TC107305) turned out to encode only a truncated phosphate transporter. In order to analyse the phylogenetic relationship between the phosphate transporters, amino acid sequences were deduced and aligned with each other and with those from other plants (Fig. 2). This revealed that amongst the class II

Fig. 1 Pi concentrations and fresh weights. Medicago truncatula plants were cultivated and fertilised with 0.13 mM (−P) or 1.3 mM P_i (+P) or fertilised with 0.13 mM P_i and inoculated with the AM fungi Glomus mosseae (Gm), Glomus intraradices (Gi) and Gigaspora rosea (Gr). Six weeks after inoculation, soluble P_i concentrations and fresh weights were measured and the ratio of the P_i concentrations (a) and the fresh weights (b) in 1.3 mM P_i-fertilised or inoculated plants to contents in the control treatments were calculated. Significant differences are indicated by asterisks (one-way ANOVA; P < 0.05; n = 4)

Fig. 2 Phylogenetic tree of plant phosphate transporters. Amino acid sequences of phosphate transporters from Medicago truncatula (MtPT1: AAB81346, MtPT2: AAB81347, MtPT3: ABM69110, MtPT4: AAM76744, MtPT5: ABM69111, MtPT6: CAQ42956), from tomato (LePT1: AAB82146, LePT2: AAB82147; LePT4: AAV97730), from potato (StPT1: CAA67395, StPT3: AAV97729, StPT4: AAWS1149), from rice (OsPT6: AF536969, OsPT9: AF536969, OsPT11: AF536971, OsPT12: AF536972, OsPT13: AF536973) and from Glomus versiforme as outgroup (GvPT1: AAC49132) were aligned using ClustalW. Based on the alignment, phylogeny was reconstructed with the program PUZZLE and the distance tree was displayed by the program TREEVIEW.
of phosphate transporters (Nagy et al. 2005), MtPT1, 2, 3 and 5 group closely together, while MtPT6 occupies a branch together with StPT1 and StPT3 from potato and StPT1 from tomato. As already shown (Nagy et al. 2005), MtPT4 belongs to the class I which members are all exclusively expressed or highly up-regulated in mycorrhizal roots. Based on the sequence similarities, all phosphate transporter genes are probably located at the plasma membrane and this has been shown for at least some of them (Liu et al. 2008).

Based on the sequences of the six different transporter genes, primer pairs were designed and used for quantitative real-time RT-PCR experiments using as template RNA extracts from roots fertilised with high amounts of P$_i$ (1.3 mM P$_i$) or inoculated with G. rosea, G. mosseae or G. intraradices (60–80% mycorrhization) or from control roots of this experiment (0.13 mM P$_i$). The results were calibrated by the levels of the constitutively expressed MtTef1 gene and $M$ values were calculated for the ratios of the relative transcript levels in the roots fertilised with 1.3 mM P$_i$ or inoculated with the different AM fungi to the values obtained from the 0.13 mM P$_i$ controls (Fig. 3). This revealed that MtPT4 is only regulated by mycorrhization as it has been shown many times before and that this is true for all three AM fungi used in the experiment. The expression is specific for cells harbouring fungal arbuscules and cannot be seen in any other tissue (Harrison et al. 2002) and the gene is essential for the function of the symbiosis (Javot et al. 2007). MtPT1 is down-regulated in roots fertilised with P$_i$ or colonised by G. intraradices and G. rosea (Fig. 3). MtPT1 is root epidermis-specific expressed (Xiao et al. 2006) and therefore likely to be responsible for P$_i$ uptake from the soil. Interestingly, colonisation by G. mosseae does not lead to a repressed transcript levels of the gene confirming the result obtained before (Hohnjec et al. 2005). MtPT2 is significantly down-regulated by high P$_i$ only (Fig. 3). MtPT2 promoter activity has been located in the vascular cylinder (Xiao et al. 2006). Therefore, the activity of the resulting protein might be also required in a mycorrhizal root explaining why the expression level is not very much affected in the symbiosis. The gene MtPT3 is down-regulated by P$_i$ and by all three AM fungi (Fig. 3), which is consistent with the results obtained by Liu et al. (2008). MtPT5 is significantly repressed only in roots colonised by G. intraradices, but not in those containing hyphae of G. mosseae and G. rosea (Fig. 3). Inoculation with G. versiforme did also not lead to a reduced RNA level of MtPT5 and it was shown that the gene is expressed in all parts of the root except the vascular tissues, but including the cortex of the colonised region (Liu et al. 2008). Concerning regulation by P$_i$ contradictory results were obtained. Growing plants in a substrate under low or high P$_i$ conditions only slightly influences MtPT5 (Fig. 3), but resupplying the mineral element to roots in a nutrient solution immediately leads to a sharp decrease of MtPT5 RNA (Liu et al. 2008). This indicates that the down-regulation of the gene by P$_i$ is only transient. MtPT6 is only regulated by the three AM fungi (Fig. 3). This clearly supports the suggestion by Liu et al. (2008) that the down-regulation of PT-encoding genes in mycorrhizal roots is not mediated by an increased P$_i$ content in the plant.

Conclusion

Comparison of gene expression in roots colonised with the AM fungus G. rosea and treated with the two phytohormones ABA and JA revealed only a very small number of
genes with a common pattern of regulation. If mycorrhiza-regulated expression occurs indirectly via changes in phytohormone levels, this seems to be only true for a very few genes. A much higher overlap was detected with the expression in P$_f$-fertilised roots and this is different to results obtained before with Glomus species. This is probably due to the higher P$_f$ concentration in the leaves of the plants colonised by G. rosea. It indicates that at least a number of these genes are not directly regulated by the AM fungus G. rosea, but rather indirectly by the better supply with the mineral nutrient. Variations in the expression patterns between the three AM fungi were also observed among the phosphate transporter gene family of M. truncatula. The fungi induce down-regulation of various number of PT-encoding genes with G. intraradices repressing most (MtPT1, 3, 5 and 6) and G. mosseae least genes (MtPT3 and 6), while G. rosea occupies a middle position down-regulating MtPT1, 3 and 6. Data for G. mosseae are not available, but the analysis of phosphate transport in tomato via the mycorrhizal or the direct uptake pathway showed that G. intraradices suppresses nearly totally the direct uptake of phosphate by the root itself and G. rosea does not (Smith et al. 2004). It has to be shown in future experiments, whether this is also true for M. truncatula and how the differential expression of the PT-encoding genes in M. truncatula is involved in such a phenomenon. The present results together with the finding that MtPT5 is expressed in root hairs and the epidermis and encodes a low affinity phosphate transporter (Liu et al. 2008) indicates that this gene might be important for the direct uptake pathway.

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