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Laser-microdissection unravels cell-type specific transcription in arbuscular mycorrhizal roots, including CAAT-box TF gene expression correlating with fungal contact and spread

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Abstract
Arbuscular mycorrhizae (AM) are the most widespread symbioses on Earth, promoting nutrient supply of most terrestrial plant species. To unravel gene expression in defined stages of *Medicago truncatula* root colonization by AM fungi, we here combined genome-wide transcriptome profiling based on whole mycorrhizal roots with real-time RT-PCR experiments that relied on characteristic cell-types obtained via laser-microdissection. Our genome-wide approach delivered a core set of 512 genes significantly activated by the two mycorrhizal fungi *Glomus intraradices* and *Glomus mossae*. Focussing on 62 of these genes being related to membrane transport, signaling, and transcriptional regulation, we distinguished whether they are activated in arbuscule-containing or the neighbouring cortical cells harbouring fungal hyphae. In addition, cortical cells from non-mycorrhizal roots served as a reference for gene expression under non-colonized conditions. Our analysis identified 25 novel arbuscule-specific genes and 37 genes expressed both in the arbuscule-containing and the adjacent cortical cells colonized by fungal hyphae. Amongst the AM-induced genes specifying transcriptional regulators were two members encoding CAAT-box binding transcription factors (CBF), designated *MtCbf1* and *MtCbf2*. Promoter analyses demonstrated that both genes were already activated by the first physical contact between the symbionts. Subsequently, and corresponding to our cell-type expression patterns, they were progressively up-regulated in those cortical areas colonized by fungal hyphae, including the arbuscule-containing cells. The encoded CBFs thus represent excellent candidates for regulators that mediate a sequential reprogramming of root tissues during the establishment of an AM symbiosis.
Introduction

Ecto- and endomycorrhizal symbioses between higher plants and soil fungi are the most widespread beneficial plant-microbe interactions on Earth. Mycorrhizae are characterized by the transfer of limiting nutrients, in particular phosphorus and nitrogen, from fungal hyphae to the plant. In return, plants deliver hexoses to the fungi (Nehls et al. 2007), leading to a strongly increased photosynthate allocation to mycorrhizal roots. Apart from direct advantageous effects resulting from improved nutrition, indirect benefits of mycorrhizal interactions are enhanced resistences against abiotic and biotic stress conditions (Smith and Read 2008).

Some 80% of all terrestrial plants enter an arbuscular mycorrhiza (AM) symbiosis with Glomeromycota fungi (Schüssler et al. 2001). During AM, extraradical hyphae emerging from germinating spores penetrate the rhizodermis via hyphopodia, pass through outer cortical cells and proliferate in the inner cortex (Parniske 2008). In Arum-type AM, these intraradical hyphae form highly branched intracellular structures termed arbuscules (Harrison 1999). It has been shown that arbuscules are transient structures that only operate for a couple of days, indicating a tight control of their development and function (Harrison 2005). Transfer of phosphorus and other minerals from fungal hyphae to the plant cytoplasm occurs at the periarbuscular interface that comprehends the fungal arbuscular membrane, the periarbuscular matrix, and the plant periarbuscular membrane (Parniske 2000, 2008). The uptake of phosphorus is energy-dependent and requires plant and fungal H⁺-ATPases (Requena et al. 2003) for an acidification of the periarbuscular space. Based on the localization of reporter proteins, Pumplin and Harrison (2009) demonstrated that different proteins are apparently targeted to specific domains of the periarbuscular membrane, indicating the presence of functional compartments at the symbiotic interface. In addition to AM-specific phosphate transporters essential for symbiotic P transfer (Harrison et al. 2002, Bucher 2007, Javot et al. 2007), two half-ABC transporters have recently been shown to be required for a functional AM symbiosis (Zhang et al. 2010).

An important benefit of legumes as AM models derives from the fact that this genus is able to enter a second beneficial plant-microbe interaction that leads to the development of nitrogen-fixing root nodules (Brewin et al. 1991). To initiate nodulation, secreted flavonoids first induce rhizobial genes required for the synthesis of lipochitooligosaccharide (LCO) nodulation (Nod)-factors. These Nod-factors, after perception by plant LysM-domain receptor kinases (Arrighi et al. 2006), activate downstream responses; that way inducing the formation of nodule primordia and mediating bacterial infection (Oldroyd and Downie 2008). During AM, a similar molecular dialogue is initiated by the host plant via strigolactones that promote the branching of fungal hyphae and activate fungal metabolism (Akiyama et al. 2005, Besserer et al. 2006). Subsequent recognition of AM fungi by the host involves the perception of diffusible Myc-signals (Kosuta et al. 2003) including LCOs structurally related to rhizobial Nod-factors (Maillet et al. 2011). Together, the diffusible Myc-signals prepare root infection by communicating the symbiotic nature of mycorrhizal fungi (Oldroyd et al. 2005, Ercolin and Reinhardt 2011). Following their entry via hyphopodia, fungal hyphae grow along a pre-penetration apparatus (PPA), a cytoplasmic channel formed after the establishment of hyphopodia (Genre et al. 2005, 2008). This coordinated cytological response of epidermal cells indicates that in addition to diffusible Myc-signals acting at a distance (Kuhn et al. 2010), there are others that require fungal contact (Kosuta et al. 2003, Kloppholz et al. 2011). It has to be pointed out that signaling also takes place in later stages
of the symbiosis; where genes specifically expressed in arbuscule-containing cells have to be activated (Harrison 2005). One such late signal triggering the induction of a phosphate transporter gene was identified as lyso-phosphatidylcholine (Drissner et al. 2007).

Targeted molecular research on AM suffers from two obstacles: an asynchronous development of the symbiosis leading to the concomitant presence of different stages, and an obligate biotrophy of AM fungi. In the last couple of years, untargeted high-throughput expression profiling was pursued to generate an inventory of AM-induced genes (Liu et al. 2003, Wulf et al. 2003, Frenzel et al. 2005, Hohnjec et al. 2005, Hohnjec et al. 2006, Küster et al. 2007b). These approaches benefited from the identification of two legumes that proved to be excellent AM models: *Medicago truncatula* (Barker et al. 1990, Rose 2008) and *Lotus japonicus* (Handberg and Stougaard 1992). In case of *M. truncatula*, current research relies on an advanced genome project (Cannon et al. 2009, Young and Udvardi 2009), more than 250,000 ESTs in the DFCI *Medicago* Gene Index (Quackenbush et al. 2001), and different microarray as well as GeneChip tools (Küster et al. 2004, Küster et al. 2007a, Hohnjec et al. 2005, Lohar et al. 2006, Benedito et al. 2008). Publicly available expression profiles based on EST, microarray, and GeneChip data can be queried using the DFCI Medicago Gene Index (Quackenbush et al. 2001), the Truncatulix (Henckel et al. 2009) and MediPlEx (Henckel et al. 2010) data warehouses as well as the Medicago truncatula Gene Expression Atlas, the latter exclusively integrating expression profiles generated via GeneChips (Benedito et al. 2008).

Although the transcriptome studies performed so far resulted in an identification of hundreds of AM-related genes (Balestrini and Lanfranco 2006, Hohnjec et al. 2006), there is limited information on the signaling components activated during the formation of arbuscules and in particular on the transcriptional regulators involved in the reprogramming of root cortical cells towards an accommodation of symbiotic fungi. One obvious reason for this can be seen in the fact that most AM-related expression profiles were based on pooled tissue samples containing a mixture of different cell-types and stages of arbuscule development. To overcome this problem, Balestrini et al. (2007) pioneered the use of laser-microdissection for the identification of arbuscule-specific phosphate transporter genes in AM roots of tomato via RT-PCR. A similar approach was later used in the studies of Gomez et al. (2009) and Guether et al. (2009) to track genes up-regulated in arbuscule-containing cells of *M. truncatula* and *L. japonicus*, respectively. On the level of individual genes, detailed in situ expression analyses were performed for selected genes (Küster et al. 2007b), including amongst others the phosphate transporter gene *MtPt4* (Liu et al. 2003), the *MtBcp1* gene encoding a blue-copper-binding protein (Hohnjec et al. 2005), and members of the AM-induced lectin gene family (Frenzel et al. 2005). Nevertheless, to our knowledge no promoter of a transcription factor gene specifically activated in AM roots was investigated so far.

In this study, we intended to sharpen our view on AM-related gene expression by a two-step approach. First, we performed a global transcriptome analysis of *M. truncatula* roots inoculated with the two widely studied AM fungi *Glomus intraradices* and *Glomus mosseae*. By using these two different microsymbionts, we were able to make use of the overlap of gene activation in both AM interactions, that way minimizing strain- or inoculum-related effects. To achieve a genome-wide identification of AM-activated genes, we relied on Medicago GeneChips; reported to cover more than 80% of the gene space in the model legume *M. truncatula* (Benedito et al. 2008). This marked
extension of gene-specific probes significantly advanced earlier microarray-based AM transcriptome studies (Liu et al. 2003, Manthey et al. 2004, Hohnjec et al. 2005), leading to the identification of a core set of 512 *M. truncatula* genes involved in both AM interactions. In a second step, we intended to shed light on the spatial expression of a subset of AM-related genes. To achieve this goal, we performed cellular expression studies via real-time RT-PCR, using RNA isolated from distinct pools of laser-microdissected cells. In our study, gene expression in *M. truncatula* arbuscule-containing cells was for the first time directly compared with transcription in the adjacent cortical cells colonized by fungal hyphae. As a control, cortical cells from non-mycorrhized roots were used to gain information on gene expression in the absence of a symbiotic interaction. With an emphasis on genes encoding membrane transporters, signaling-related proteins, and transcriptional regulators, this approach identified novel components of the cell-specific programme orchestrating AM symbiosis. In total, we identified 25 arbuscule-specific genes, while 37 genes were activated both in arbuscule-containing and in the neighbouring cells. Together, these results highlight general mechanisms underlying fungal colonization up to the formation of arbuscules. Among the transcriptional regulators, we identified two highly similar genes encoding CAAT-box binding transcription factors (CBF), which we analyzed in more detail via the expression of promotor-GUS fusions in transgenic roots. Remarkably, both genes are already activated by the initial physical contact between fungal hyphae and the plant epidermis and are expressed concomitantly with fungal colonization of the root cortex up to the formation of arbuscules, making the encoded CBFs excellent candidates for regulators mediating the sequential reprogramming of root tissues during the establishment of an AM symbiosis.

**Results and Discussion**

**Genome-wide transcriptome profiling identifies a core set of 512 *Medicago truncatula* genes related to root colonization by AM fungi**

*M. truncatula* roots colonized with either *G. intraradices* or *G. mosseae* under conditions of phosphate limitation (20 µM phosphate) were stained for fungal structures 28 days post inoculation (McGonigle et al. 1990). To minimize dilution by non-colonized regions, we selected roots with a similarly high root length colonization (60 to 80%) and high arbuscule frequencies (60 to 75%) for the isolation of total RNA. Samples were checked for AM (*MtPt4*) and nodule marker (*MtEnod18*) gene expression via RT-PCR, as described by Hohnjec et al. (2005). Since some AM-induced genes can be activated by phosphate, we incorporated a study of gene expression in nonmycorrhizal roots grown for 28 days in the presence of a high (2 mM) phosphate concentration. As a common control, nonmycorrhizal roots grown for 28 days under phosphate limitation (20 µM phosphate) were harvested. Total RNA isolated from three biological replicates of mycorrhizal and nonmycorrhizal roots was used for GeneChip hybridizations. In our study, each biological replicate is defined as a pool comprising five root systems. The complete dataset is available from the Gene Expression Omnibus (accession number GSE32208) and is also included as Supplemental Table S1. Applying a two-fold induction at a false discovery rate (FDR)-corrected p-value of p<0.05 as a cutoff, we found that 1214 and 888 genes were up-regulated in *G. intraradices*- and *G. mosseae*-colonized roots, respectively, whereas 576 genes were co-activated in both AM interactions (Figure 1, Supplemental Table S2).
Of the 576 AM-induced genes, 44 were also activated at the cutoffs mentioned above in roots treated with 2 mM phosphate, leaving a core set of 532 genes significantly induced by two different AM fungi but not via elevated phosphate levels (Figure 1, Supplemental Table S2). The low overlap between phosphate-induced gene expression and transcriptional activation by AM fungal colonization is in line with reports of Liu et al. (2003) and Hohnjec et al. (2005), illustrating that phosphate supply is unable to mimic AM-induced gene expression.

Our core set of 532 AM-related genes is supported by a search in the MediPlEx database (Henckel et al. 2010), combining in silico gene expression in AM EST collections from the DFCI Medicago Gene Index (libraries #ARE, #ARB, T1682, #GFS, 5520, #9CR) with the GeneChip hybridizations detailed above. This search returned 91 genes of the core set as AM-induced with an R-value larger than 2 (Stekel et al. 2000, Supplemental Table S2). Moreover, our core set contained 51 genes reported as co-induced in the same two AM interactions when relying on oligonucleotide microarrays (Hohnjec et al. 2005, Supplemental Table S2), whereas it encompassed 346 genes reported as activated at least twofold by G. intraradices under different experimental conditions on the basis of Medicago GeneChips (Gomez et al. 2009, Supplemental Table S2).

Amongst the core set of AM-related genes, 15 well-known AM expression markers (Hohnjec et al. 2006, Küster et al. 2007b) were identified as strongly induced in our conditions (Table 1). As an example, the MtPt4 gene encoding a phosphate transporter known to be required for an efficient AM symbiosis (Javot et al. 2007) was activated more than 1000 (log2=10.02)- and 400 (log2=8.65)-fold in G. intraradices- and G. mosseae-colonized roots, respectively. This very strong induction of AM-related marker genes (Table 1) indicates that our approach should allow an identification also of those genes expressed either transiently or activated less strongly during arbuscule development.

Via comparisons of Medicago GeneChip probes to currently available Glomus intraradices sequences (Tisserant et al. 2011), 20 Glomus genes were identified amongst the set of 532 genes co-induced in both AM interactions (Supplemental Table S2). The remaining 512 AM-related M. truncatula genes were grouped into functional categories, based on automated annotations of the encoded gene products via the SAMS software (Bekel et al. 2009) and Gene Ontology classifications (http://www.medicago.org). In addition, MapMan (Usadel et al. 2005) was used to visualize gene expression profiles and to identify functional categories expressed most significantly different from others.

In Figure 2, the distribution of AM-coinduced genes into functional categories is shown. Typical AM-related gene families of unknown biological function encoding annexins (e.g. MtAnn2, Manthey et al. 2004), blue-copper proteins (e.g. MtBcp1, Hohnjec et al. 2005), germin-like proteins (e.g. MtGlp1, Doll et al. 2003), and lectins (e.g. MtLec5 and MtLec7, Frenzel et al. 2005) are classified separately. For all these gene families except the one encoding annexins, comparisons to the Medicago truncatula Gene Expression Atlas (He et al. 2009) revealed that their members were expressed either specifically or were almost exclusively activated in AM roots in comparison to all other tissues analysed (data not shown), that way widening our knowledge on previously unknown AM-related family members.

In addition to these gene families, the following functional categories were most prominent (Figure 2, Supplemental Table S2): (1) cell wall biosynthesis and modification, including several enzymes related to a remodeling of the extracellular matrix; (2) chitin degradation, including several chitinases that
might be involved in the disassembly of fungal structures; (3) protein degradation, including a range of different proteinases and peptidases that can be connected to the dynamic turnover of mycorrhizal structures; (4) hormone metabolism, including several genes involved in gibberellin biosynthesis; and (5) secondary metabolism, including a high number of cytochromes, components of the carotenoid metabolism, and UDP-sugar transferases. The activation of these functions is in line with previous reports on gene expression in AM roots (Liu et al. 2003, Wulf et al. 2003, Hohnjec et al. 2006, Küster et al. 2007b, Guether et al. 2009).

Interestingly, the categories "membrane transporters", "signaling" and "transcription factors" were not only prominent but their members were also expressed most significantly different from others in AM roots (data not shown), based on statistical analyses implemented in MapMan (Usadel et al. 2005). In addition, many of these candidate genes either displayed a mycorrhiza-specific or mycorrhiza-enhanced expression in AM roots according to the Medicago Gene Expression Atlas (He et al. 2009), suggesting a relevance for AM interactions. Remarkably, these analyses also returned six TF genes co-induced in our conditions as being specifically expressed in AM tissues (Supplemental Figure S1).

Since the three cellular functions mentioned are particularly relevant for coordinating the reprogramming of root cortical cells towards an accomodation of fungal structures, we investigated the cell-type expression of 71 candidate genes selected from these categories via laser-microdissection.

**Longitudinal sections allow an accurate separation of specific cell-types from AM roots via laser-microdissection**

We used laser-microdissection to obtain specific pools of three different cell-types in four biological replicates: cortical cells from non-colonized control roots (CCR), cortical cells from mycorrhizal roots (CMR), and cortical cells containing arbuscules (ARB) (Figure 3). We found that embedding of roots in Steedman’s wax (Gomez et al. 2009) preserves root morphology, allowing an identification of vascular tissues, cortical cells and the epidermal cell layer including root hairs (Figure 3 A-C). In contrast to transverse sections (Balestrini et al. 2007, Gomez et al. 2009), our longitudinal sections offered the possibility to evaluate the colonization status of whole root regions. Chains of arbuscules in different developmental stages and fungal hyphae growing in the extracellular space of adjacent cortical cells were clearly visible (Figure 3 E-H). This facilitated cell harvest and allowed us to focus not only on mature arbuscules filling the complete cell lumen (ARB-samples; Figure 3 D,G,H), but also on cortical cells interspersed with fungal hyphae in the immediate neighbourhood of arbuscule-containing cells (CMR-samples; Figure 3 E,F). Since each cell pool from the CMR cell-type differs in the density and growth of fungal hyphae, the CMR samples are expected to display the strongest variation of transcriptional changes.

To assess the suitability of the collected samples and to check in particular for cross-contamination between CMR and ARB samples, the expression of the six AM marker genes *MtPt4*, *MtBcp1*, *MtScp1*, *MtLec5*, *MtGlp1* and *MtHa1* (Harrison et al. 2002, Hohnjec et al. 2005, Liu et al. 2003, Frenzel et al. 2005, Doll et al. 2003, Krajinski et al. 2002) was measured in all four biological replicates. Whereas transcripts of the *MtTef* gene used as a constitutive control were amplified from all three cell-types (Figure 4 A), transcripts of the six AM-specific genes tested were not detected in CCR samples, as could be expected (data not shown).
The phosphate transporter gene \( MtPt4 \) was used as an expression marker for cross-contamination between the CMR and ARB samples, since this gene is exclusively transcribed in cells containing arbuscules (Harrison et al. 2002). The detection of \( MtPt4 \) transcripts in all ARB in contrast to their absence in all CMR samples (Figure 4 B) indicates that no significant cross-contamination of CMR-samples with material from arbuscule-containing cells occurred. In contrast to \( MtPt4 \), transcripts of the other five genes tested could be amplified both in CMR and ARB samples (Figure 4 B). Of these, the serine carboxypeptidase gene \( MtScp1 \) was the only one expressed at similar levels in both cell-types, whereas the \( MtBcp1 \) gene encoding a blue copper protein, the lectin gene \( MtLec5 \), the \( MtGlp1 \) gene specifying a germin-like protein, and the H\(^+\)-ATPase gene \( MtHa1 \) displayed an up to 160-fold induction in ARB-cells (Figure 4 C). While the results obtained for \( MtScp1 \) and \( MtBcp1 \) are in accordance with promoter studies indicating expression outside of arbuscule-containing cells (Liu et al. 2003, Hohnjec et al. 2005, \( MtLec5 \), \( MtGlp1 \) and \( MtHa1 \) have so far been reported to be arbuscule-specific (Frenzel et al. 2005, Doll et al. 2003, Krajinski et al. 2002). This deviance might be due to a higher sensitivity of our PCR-based method in comparison to the in situ expression methods used in the cited studies.

Together, our results demonstrate that the total RNA prepared from all three cell-types was suitable for identifying arbuscule-specific as well as arbuscule-enhanced expression patterns, even if the harvested cell-types were collected in close proximity.

Laser-microdissection identifies novel arbuscule-specific genes and genes being generally expressed in cortical cells colonized by AM fungi

Expression of 71 candidate genes was measured by one-step real time RT-PCR in at least three biological replicates. In these experiments, nine genes could not be amplified reproducibly from any cell-type and were not considered further. Figure 5 gives an overview of the remaining 62 genes and their expression patterns in the laser-microdissected samples. Except of three genes encoding an ABC-transporter (Mtr.44070.1.S1_at), a calcium-binding protein (Mtr.4781.1.S1_at), and an AP2/ERF transcription factor (Mtr.11570.1.S1_at), none of these genes could be amplified reproducibly from CCR cells (data not shown), suggesting that they are either truly mycorrhiza-specific or are expressed in other cell-types than the ones investigated here, e.g. the vascular tissue.

Amongst the 62 genes analysed for cell-specific expression, 21 displayed a specific activation in the arbuscule-containing cells, including six membrane transporter genes, seven signaling-related genes, six genes encoding transcription factors, and two fungal genes (Figure 5, Supplemental Figure S2). In addition, four genes were detected in all ARB, but in only one out of three CMR samples, indicating that they are expressed at a very low level in the latter cell-type. Six genes specifically expressed in arbuscule-containing cells were described to be activated in these cells before (Gomez et al. 2009), but none of them was so far known to be restricted to this cell-type within mycorrhizal roots.

Interestingly, the majority of the genes investigated was found to be expressed both in arbuscule-containing cells (ARB) and in the adjacent cortical cells being colonized by fungal hyphae at different levels (CMR). Here, transcripts of 37 genes, including 14 membrane transporter genes, 8 signaling-related genes, 12 transcription factor genes, and three fungal genes were detected. Of these 37 genes, 28 were expressed at comparable levels in both cell-types, whereas 9 were significantly induced in either CMR or ARB (Figure 6). So far, only four genes of this group were known to be
expressed in arbuscule-containing cells (Gomez et al. 2009), but no information was available on their activity in the adjacent cells being connected to hyphal growth. In total, cell-specific expression patterns were identified for 62 AM-related genes. In the following, our results are discussed in detail with respect to the functional classification of the genes.

**Glomus intraradices genes**

Seven AM-related genes of fungal origin were selected for cell-specific transcriptome analysis. These genes encoded five membrane transporters, one calcium-binding protein, and two transcription factors (Figure 5). Interestingly, our analysis of the expression patterns in the specific cell pools revealed that four genes seemed to be specifically expressed in arbuscules, indicating that the distribution of fungal transcripts can be restricted to these, irrespective of the coenocytic nature of the microsymbiont. The genes investigated included three members specifying ATPases, two of which represented cytoplasmatic ATPases probably associated with proteasomes and thus being involved in protein degradation, whereas the third one was predicted to be membrane-localized and involved in the transport of phospholipids or cations (Figure 5). While two of the ATPase genes showed a specific expression in arbuscules, one was induced at comparable levels in both CMR and ARB samples. In addition, a gene specifying a voltage dependent channel protein was also activated in both cell-types, similar to a signaling-related gene encoding a calcium-binding protein (Figure 5). Interestingly, the two fungal genes related to transcriptional regulation, encoding a zinc finger and an RNA-binding protein, showed specific expression in ARB (Figure 5). These genes might therefore control the transcription of fungal genes specifically required during arbuscule formation or function.

**Medicago truncatula genes encoding membrane transporters**

A functional AM symbiosis is characterized by the exchange of nutrients between the micro- and macrosymbionts. It is thus not surprising that 37 *M. truncatula* genes encoding membrane transporters were co-activated in the two AM-interactions studied (Supplemental Table S3), including 13 genes that Benedito et al. (2010) identified previously on the basis of expression data reported by Gomez et al. (2009). Apart from the AM marker genes *MtPt4* and *MtHa1* already discussed above as well as the mycorrhiza-induced aquaporin gene *MtNip1* (Uehlein et al., 2007), cellular expression patterns were determined for 20 genes of this functional categorie (Figure 5).

The largest group comprised genes encoding oligopeptide transporters of the H+/oligopeptide symporter type. Eleven members of this class were coinduced in both AM interactions and seven of them were examined in the laser-microdissected cell pools (Figure 5). Proton-dependent oligopeptide transporters (POTs) are reported to be involved in the uptake of small peptides into eukaryotic cells (Paulsen and Skurray 1994). Possible functions are thus connected to an intake of degraded fungal proteins subsequent to the action of AM-activated proteases (Supplemental Table S2), alternatively the uptake of fungal effector peptides, such as the recently reported SP7 protein (Kloppholz et al. 2011, Plett et al. 2011), is an intriguing possibility. The wide range of possible POT functions is mirrored by cellular expression patterns, since we found two arbuscule-specific POT genes (Figure 5), three POT genes equally expressed in CMR and ARB cells (Figure 5), and one POT gene being induced in each cell-type (Figures 5, 6).
A second prominent group of AM-related genes encoded five ABC-transporters and three aquaporins. From this collection, we identified one ABC-transporter gene as ARB-specific (Figure 5) and three more as being expressed with no significant difference between CMR and ARB (Figure 5). With MtStr and MtStr2, two M. truncatula ABC transporter genes were recently reported to be important for AM (Zhang et al. 2010). Both genes belong to the G subfamily of ABC-transporters and are specifically expressed in arbuscule-containing cells. The ABC-transporter identified as arbuscule-specific in our study belongs to the same subfamily, suggesting a similar function. Zhang et al. (2010) speculated that strigolactones might be a substrate for these transporters, inducing the strong ramification of fungal hyphae that leads to arbuscule formation, which would explain the cell-specific expression. In contrast, the three ABC-transporter genes we found expressed in arbuscule-containing and adjacent cortical cells specify PGP MDR (P-glycoprotein multidrug resistance) transporters of the ABCB subfamily. Since several members of this group are known to transport auxins (Geisler and Murphy, 2006), a role for these might be the fine-tuning of auxin distribution in colonized root cells.

In the group of aquaporins, we investigated the expression pattern of two NIP (Nodulin 26-like intrinsic proteins) genes including MtNip1, which was described as AM-induced by Uehlein et al. (2007) and was also strongly AM-induced here. Due to the fact that this gene is slightly induced by phosphate, it was not included in our core set of 512 AM-related genes. In line with the results of Gomez et al. (2009), we found MtNip1 to be activated in ARB cells. In addition, we could show here that this gene is expressed exclusively in this cell-type, whereas the second NIP gene was expressed in the surrounding, hyphae-containing cortical cells as well (Figure 5). Interestingly, MtNip1 and other NIPs did not facilitate water uptake in heterologous expression systems, but acted as low-affinity transport system for ammonium (Uehlein et al. 2007), which is the main form in which nitrogen is supplied to the plant by AM fungi (Govindarajulu et al. 2005). These observations support the idea that nitrogen is provided by the microsymbiont via arbuscules as well as fungal hyphae growing in the apoplast of cortical cells, and the two NIPs might be involved in uptake of these nutrients into host cells.

Strikingly, the most strongly induced transporter genes in our experiment encoded three defensins. These seem to be of special importance for arbuscule-containing cells, since one defensin gene was exclusively expressed (Figure 5) and the other two were significantly upregulated in ARB (Figures 5, 6). Many members of the defensin gene family are activated in plants during defense reactions in response to diverse pathogens (Thomma et al. 2002), including harmful fungi. Although the activation of plant defence responses has been reported for the initial stages of mycorrhizal colonization, they seem to be effectively controlled and downregulated when the symbiosis is completely established (Harrison and Dixon 1993). Interestingly, some defensins are known to reduce the elongation of fungal hyphae and to cause strong hyphal branching (Broekaert et al. 1995). Therefore, those defensins specifically activated during a mature mycorrhiza may in fact influence hyphal ramification during early arbuscule development or control arbuscule lifespan.

The remaining membrane transporter genes we found to be upregulated during AM are mainly involved in the transfer of micro-nutrients, macro-nutrients such as nitrate, or carbohydrates. Concerning ion transporters of the MtZIP family (Burleigh et al. 2003), we identified a gene encoding the manganese transporter MtZip7 (López-Millán et al. 2004). We found the MtZip7 gene and a zinc transporter gene to be expressed in CMR and ARB cells alike (Figure 5), whereas a copper
transporter gene was found to be ARB-specific (Figure 5). Since AM fungi probably improve micro-nutrient supply of host plants (Clark and Zeto 2000, Parniske 2008), the three transporter genes might be activated by the plant to enhance uptake of these elements from the microsymbiont. The fact that we found two of the transporter genes mentioned to be expressed in both arbuscule-containing and surrounding, hyphae-containing cortical cells supports the assumption that an exchange of nutrients by the symbiotic partners or at least an uptake by the plant is not restricted to arbuscules.

With respect to sugar allocation in AM roots, we analyzed the expression patterns of two carbohydrate transporters. Due to the fact that hexoses derived from the plant metabolism are supplied to mycorrhizal fungi, AM roots represent strong carbon sinks. Whereas hexoses generated from sucrose via the activity of apoplastic invertases are directly available to the fungus (Schaarschmidt et al. 2007), hexoses provided by cytoplasmic invertases or sucrose synthases (Hohnjec et al. 1999) first have to be exported to the extracellular space (Baier et al. 2010). Up to now, little is known about the carbohydrate transporters involved in the translocation of symplastic hexoses to the apoplastic interface. The two genes investigated here are candidates for this function, but showed no consistent expression. Since only one was ARB-specific (Figure 5), sucrose transfer is probably not limited to arbuscules, which is consistent with the expression of genes encoding the cytoplasmatic sucrose-cleaving enzyme MtSucS1 (Hohnjec et al. 2003) and the hexose transporter Mtst1 (Harrison 1996).

**Medicago truncatula genes related to signaling**

In Supplemental Table S4, the expression characteristics of 41 genes encoding signaling-related components induced in both AM interactions are summarized. A large group consists of genes probably involved in intracellular signal transduction, including 13 genes encoding protein kinases, a phosphatase and a protein phosphatase inhibitor. Furthermore, genes specifying two inositol polyphosphate phosphatases, a phosphatidylinositol transfer protein, a calmodulin-binding protein, two Rop guanine nucleotide exchange factors, and a G-protein deserve attention. Together, they might represent those members of large gene families that mediate calcium, phosphoinosite, or G-protein signaling in AM roots. Seven genes from this group and an additional calmodulin gene only induced by *G. intraradices* were investigated for their cellular expression pattern (Figure 5). Interestingly, we found genes encoding three protein kinases (two serine/threonine kinases and a SNF1-related kinase) and one inositol triphosphate phosphatase to be exclusively expressed in ARB (Figure 5). To our knowledge, these are the first genes reported to be involved in internal signaling processes that are specifically up-regulated in arbuscule-containing in comparison to the neighbouring cells. Another serine/threonine protein kinase gene was strongly induced in ARB (Figures 5, 6), whereas a calmodulin gene, a G-protein gene and one additional serine/threonine protein kinase gene were expressed at equal levels in ARB and CMR (Figure 5), hinting that they are involved in signal processes occurring in cortical cells of colonized root areas in general.

A second group of AM-activated, signaling-related genes encodes components associated with vesicle-mediated transport, including a syntaxin, a clathrin assembly protein, a GTP-binding protein, and a basic secretory protein. It is tempting to speculate that these proteins are involved in the membrane biogenesis associated with an intracellular accomodation of fungal structures. The expression pattern of the two genes we analyzed in the specific cell pools is in line with the
assumption that in particular the arbuscule-containing cells are places of intensive membrane build-up and turnover, since a gene encoding a clathrin assembly protein was exclusively detected in ARB (Figure 5) and a syntaxin gene was found to be ARB-induced (Figures 5, 6).

Three chitinase genes strongly induced in both AM interactions were included in the analysis of signaling-related genes (Figure 5). Chitinases hydrolyze β-1,4-glycosidic bonds (Salzer et al. 2000) and are mostly regarded as part of plant defence mechanisms (Arlorio et al. 1992). Although several chitinase genes are up-regulated during early phases of nodulation and mycorrhization, this activation is transient and they are in general not regarded to play a role during later stages. Nevertheless, several members of the class III chitinase gene family are activated specifically during AM (Salzer et al. 2000), and for one of these genes an arbuscule-specific localization was found by in situ hybridization (Bonanomi et al. 2001). In this study, we investigated three members of the chitinase gene family and found two of them only in ARB (Figure 5), whereas the third one was active in CMR and ARB (Figure 5). The predominant ARB-expression of chitinase genes might thus support the formation of functional symbiotic interfaces by reducing the amount of chitinous elicitors.

The largest group of signaling-related genes encompassed 15 genes encoding receptor kinases with a predicted membrane localization that presumably perceive extracellular AM-related signals. From this group, we investigated genes encoding a leucin-rich repeat (LRR) receptor kinase and the LysM receptor kinase MtLyr1. In addition, a gene encoding a membrane-bound lectin with predicted kinase activity was analyzed (Figure 5). Whereas the lectin gene and the LRR receptor kinase gene were equally expressed in both cell-types studied, MtLyr1 was specifically expressed in ARB cells (Figure 5). In Medicago truncatula, the two LysM receptor kinases MtNfp, representing the Lyr-type, and MtLyk3, representing the Lyk-type of this family, were identified as Nod-factor receptors (Gough and Cullimore 2011). MtNfp also plays a role during early AM interactions, since it is essential for the perception of the recently characterized Myc-LCO signals (Maillet et al. 2011). Since MtNfp is not essential for the establishment of an AM (Amor et al. 2003), Maillet et al. (2011) speculated that another receptor active at higher Myc-LCO concentrations must exist. These could be achieved during arbuscule formation, as a result of a tighter contact between the two symbiotic partners during infection of cortical cells. Interestingly, it was shown recently that in the non-legume Parasponia, an orthologue of MtNfp named PaNfp is essential for the symbioses with nitrogen-fixing bacteria as well as AM fungi (Op den Camp et al. 2011). In contrast to legume plants, the Parasponia genome only contains a single Lyr-type receptor kinase gene, whereas legume plants contain two copies (Op den Camp et al. 2011). It was speculated, that during evolution of legume plants and their ability to form nodules, duplication of the initial Lyr-gene occurred and one copy became the receptor for rhizobial LCOs, whereas the other one is targeted by Myc-LCOs. The closest relative of MtNfp is MtLyr1, which could thus represent a potential receptor for Myc-LCOs (Op den Camp et al. 2011). We could show here that MtLyr1 is exclusively expressed in ARB and not in the surrounding cortical cells from mycorrhized roots. This fits well to the phenotype of RNAi-knockdowns of the more ancient PaNfp gene, since here the AM symbiosis is aborted at the point of arbuscule formation. Our finding that MtLyr1 is exclusively expressed in arbuscule-containing cells supports the fact that the encoded receptor is needed for this step, possibly to perceive Myc-LCOs secreted during later stages of AM.
**Medicago truncatula** genes encoding transcriptional regulators

In total, 25 genes encoding transcription factors (TFs) were identified as co-induced in AM roots (Table 2), comprising the families AP2/ERF, Z-C2H2, CAAT-box binding, GRAS, MYB, WRKY, and NAC. Of these, only five GRAS-TF genes and one gene encoding a Myb-TF (designated *MtMyb1*, designated in Table 1) were previously reported to be specifically activated in roots colonized with *Glomus* spec. (Liu et al. 2003, Gomez et al. 2009), while a few members of other gene families were reported to be of relevance in the root nodule symbiosis.

With six members each, genes encoding GRAS and AP2/ERF transcriptional regulators were most prominent amongst the TFs identified. This is particularly interesting, since analogous proteins are involved in early signaling in the root nodule symbiosis. Here, the GRAS-TFs *MtNsp1* and *MtNsp2* (Smit et al. 2005, Kalo et al. 2005) as well as the AP2/ERF-TF *MtErn1* (Middleton et al. 2007) are essential for the activation of symbiosis-related genes via Nod-factor signaling. Detailed studies on these transcription factors led to the identification of two further AP2/ERFs (*MtErn2*, *MtErn3*) and revealed that GRAS and AP2/ERF proteins interact with promoter sequences of early nodulin genes (Andriankaja et al. 2007, Hirsch et al. 2009). Since the TF proteins mentioned obviously represent an important control system in the regulation of symbiosis-specific genes, it is interesting that with *MtErn1* and *MtErn2*, two of these were found to be induced in AM roots in our study.

Additional prominent TF genes induced in AM roots specified CAAT-box binding factors (Cbf) of the HAP3 and HAP5 type, C2H2 zinc finger proteins, and Myb proteins, with three members each (Table 2). In the root nodule symbiosis, the Mthap2-1 CAAT-box binding factor was identified as a key developmental regulator by Combier et al. (2006), whereas the zinc-finger protein Mszpt2-1 was shown to be essential for the differentiation of the nitrogen-fixing zone of root nodules (Frugier et al. 2000). So far, both gene families were not reported to be related to *Glomus*-colonized roots.

Since we regarded transcriptional regulators as particularly interesting for our study, we investigated the cellular expression pattern of a large subset of 17 TF genes, including members of the five most prominent families. Additionally, an AP2/ERF induced by diffusible factors from AM fungi (N. Hohnjec, Leibniz Universität Hannover, Germany, unpublished data) was included in the analysis (Figure 5). Interestingly, we found arbuscule-specific or ARB-induced genes in four of the families investigated. Whereas a C2H2 zinc finger, two AP2/ERF, and three GRAS TF genes were ARB-specific (Figure 5), *MtMyb1* and a further GRAS gene were significantly activated in ARB cells (Figure 5). The encoded TFs thus represent candidates for regulators that control the expression of genes required for proper arbuscule development and function. On the other hand, all gene families containing ARB-specific members also included genes expressed at equal levels in CMR and ARB (Figures 5, 6), indicating that different members of a gene family control different steps of the symbiosis.

In contrast to the TF genes mentioned so far, all three Cbf genes were expressed at similar levels in CMR and ARB (Figure 5), indicating a more general role in the coordination of fungal colonization. Cbf proteins are known to regulate the expression of genes containing CCAAT motifs in their promoter sequences by forming heterotrimeric complexes that bind to the CAAT-box (Combier et al. 2008). Since CCAAT motifs are present in about 30% of eukaryotic promoters (Mantovani et al. 1998), Cbf proteins represent global regulators of gene expression that probably gain specificity via interactions with other transcription factors (Maity and Crombrugghe 1998). That way, Cbf activation during an AM
symbiosis can mediate an expression of whole sets of AM-related genes via the recognition of their promoter regions and a subsequent interaction with other transcriptional regulators. We therefore analysed the two genes Mtr.51511.1.S1_at (designated MtCbf1) and Mtr.16863.1.S1_at, (designated MtCbf2), both encoding CAAT-box binding TFs of the HAP5-type, in more detail.

**Expression of the MtCbf1 and MtCbf2 genes encoding CAAT-box binding transcription factors correlates with fungal contact and spread**

The two genes MtCbf1 and MtCbf2 are highly similar (96.3% identity on the level of nucleic acids, Supplemental Figure S3), indicating that they might be derived from a duplication event. They are located in close proximity on *M. truncatula* chromosome 2 (http://www.medicagohapmap.org/, BAC clone AC136138), being separated by 2 different *M. truncatula* genes. Both MtCbf promoter sequences display no marked similarities, except of the region immediately upstream of the start codons (Supplemental Figure S4). To obtain a comprehensive insight into the up-regulation of the MtCbf1 and MtCbf2 genes during successive stages of fungal colonization, their activity was analyzed both via real-time RT-PCR and via the expression of promoter-GUS-fusions in transgenic roots, using a four week time-course of mycorrhization. The results obtained confirmed the expression patterns of MtCbf1 and MtCbf2 detected in ARB and CMR cell-pools, and in addition revealed a striking activation of these genes already during very early stages of the AM interaction.

The analysis of reporter gene expression in transgenic roots showed that both promoters displayed no activity in roots grown in the absence of mycorrhizal fungi (data not shown). In mycorrhizal transgenic roots, first spots of blue staining in epidermal cell layers were observed as early as 5 dpi for MtCbf1 and MtCbf2 in those places where fungal hyphae just attached to the plant epidermis, but had not yet entered the host cells (Figures 7A to 7C; 8A and 8D; 8B and 8E). This activation was always dependent on direct physical contact between the two symbiotic partners and is therefore most probably not induced by diffusible signals from AM fungi. Once the fungus had entered the root cortex, promoter activity expanded to cortical and arbuscule-containing cells, always related to the progression of fungal hyphae (Figures 7D and 7G; 7E and 7H; 8C and 8F). Whereas epidermal staining became even more pronounced for MtCbf1 during these stages and was so intense that staining of underlying cell layers could hardly be distinguished (Figures 7F and 7I), epidermal staining for MtCbf2 remained at a lower level, and the activity of this promoter appeared stronger in the cortex (Figures 8G and 8H). During later AM stages, when fungal progress is mostly achieved by an intraradical spread of the hyphae, no activity in epidermal cells was observed any more for both promoters, and reporter gene activity became restricted to those cortical cells that were either in contact with fungal hyphae or that contained arbuscules (Figures 7K to 7P; 8I to 8P). No GUS-staining was observed in root regions which did not contain fungal infection units (data not shown).

In our laser-microdissection experiments, both genes were identified as expressed with no significant difference between CMR and ARB (Figure 9A and 9B), although MtCbf1 tended to be stronger expressed in CMR (3,5-fold induction) and MtCbf2 appeared activated in ARB (3,8-fold induction). This tendency was also mirrored in the reporter gene expression patterns driven by the two promoters, since GUS-staining mediated by the MtCbf2 promoter seemed to be more intense in the arbuscule-containing cells (Figure 8M), whereas GUS staining mediated by the MtCbf1 promoter was equal in
arbuscule-containing and surrounding cortical cells interspersed with fungal hyphae (Figure 7M). Taking into account that GUS-staining appears more intense in the densely filled cells containing arbuscules, these expression patterns largely confirm our results obtained via laser-microdissection. During the mycorrhizal time course performed from 7 to 28 dpi, the \textit{MtCbf1} and \textit{MtCbf2} promoters displayed a constantly rising activity, visualized by representative GUS staining patterns from different timepoints (Figure 9C to 9K). This increase closely correlated to the expression of the AM marker genes \textit{MtPt4}, representing an arbuscule-specific gene, and \textit{MtBcp1}, representing a gene expressed in arbuscule-containing as well as in the surrounding, hyphae-containing cortical cells (Figure 9L). In our time course, \textit{MtPt4} and \textit{MtBcp1} from 7 dpi to 28 dpi showed a constantly increased transcription, leading to an up-regulation of 836-fold and 341-fold, respectively, at 28 dpi in comparison to 7 dpi. Interestingly, the expression of both \textit{MtCbf1} and \textit{MtCbf2} was strongest already at 21 dpi, where 70-fold and 37-fold inductions in comparison to 7dpi was reached. At 28 dpi, the activity of both transcription factor genes already declined, probably due to a reduced number of epidermal infection events at this timepoint. Together with our promoter studies (Figures 9F and 9K), this observation underlines the importance of both genes already for early infection stages. It can be concluded that \textit{MtCbf1} and \textit{MtCbf2} seem to be relevant during all stages of the AM symbiosis being characterized by a direct physical contact between the two partners. Therefore, the encoded CAAT-box transcription factors represent excellent candidates for novel regulators not only during later AM stages, but especially for the first steps of the interaction, where knowledge both on regulators mediating infection as well as infection-related expression markers is scarce. Interestingly, after the first activation in the epidermal cell layer, activity of both promoters seemed to precede the actual fungal colonization, since GUS staining did not only spread into the surrounding epidermal cells, but was several cells ahead of the proceeding fungal hyphae in the cortex. It is tempting to speculate that this gene expression pattern is promoted by a short-distance signal that prepares cells for an arrival of the microsymbiont, possibly facilitating fungal entry in the proximity of the first infection site or the hyphal spread in the cortex. In this context, the ability of CBFs to interact with a large range of promoters becomes particularly interesting, since they might thus be able to activate large parts of the symbiotic programme in colonized root cortical cells, ultimately leading to the reprogramming of host cells towards an accommodation of symbiotic fungi.

**Conclusion**

An attribution of gene activity to defined developmental stages is an important step towards the understanding of complex biological processes. That way, our identification of specific cellular expression patterns for a large subset of AM-related genes in mycorrhizal roots allows an association of these genes with the different developmental stages of an AM symbiosis, as shown in Figure 10. The colonization of plant roots by AM fungi is characterized by at least four distinct steps. While the first step is characterized by an exchange of diffusible signals between the two partners, causing the activation of signal cascades in epidermal and cortical cells, the second step involves direct physical contact, leading to hyphopodium formation by the fungus and PPA formation on the plant side, followed by penetration of rhizodermal cells by fungal hyphae. In the third step, fungal hyphae spread in the apoplast, resulting in the fourth and most intimate step, the formation of arbuscules. It has to be
noted that step two, three, and four are achieved in a short time period, where AM development is characterized by a spread of infection units from initial entry points, being accompanied by a sequential build-up and decay of arbuscules.

Due to the fact that our AM expression profiles were recorded on the basis of pooled tissue samples of strongly mycorrhized roots, it is likely that processes occurring during later, functional stages of AM (step three and four) were preferentially identified, and these are also represented by our specific cell pools derived by laser-microdissection. We propose that the genes we identified to be active in cortical cells colonized by fungal hyphae and arbuscule-containing cells are related to the general progression of fungal hyphae in the root cortex, whereas genes only active in arbuscule-containing cells account for specific functions of this highly specialized symbiotic interface. Our identification of transporters, signaling-related genes and transcription factors with distinct expression patterns provide insight into the genetic processes accompanying and allowing progression towards a functional symbiosis (Figure 10). Interestingly, only two genes with enhanced expression in cortical cells colonized by fungal hyphae in comparison to arbuscule-containing cells were identified, suggesting that processes exclusively related to the growth of fungal hyphae in the cortex, which are not needed for arbuscule formation and function, are rare. It remains unclear, whether the mechanisms allowing fungal penetration of roots also guide intraradical growth of fungal hyphae. Regarding the fact that the primary infection is guided intracellularly via the PPA, whereas the growth of fungal hyphae in the cortex is mostly apoplastic and keeping in mind that fungal hyphae in contact to cortical cells may also be involved in nutrient transfer to the plant, there are differences between these two steps. It therefore seems feasible that there will be a set of genes whose activity is required during all stages, like the two CAAT-box transcription factor genes identified in this study (Figure 10). In addition, some genes specifically needed for the initial infection likely exist, analogous to the arbuscule-specific genes we identified here. Both steps are tightly controlled by the plant, mirrored in two types of mutants identified for the AM symbiosis (Harrison 2005). These either mediate no entry of fungal hyphae at all, or allow entry and spread but no arbuscule-formation, suggesting the existence of specific fungal signal molecules and transduction pathways required in these stages.

The identification of two CBF genes specifically activated during all stages of an AM infection delivers two candidates mediating a high-level control of gene expression during the colonization of roots by AM fungi. It will be interesting to see, whether other CAAT-box TF genes identified as AM-induced show a similar expression pattern and how many genes active during apoplastic growth of fungal hyphae are already activated during earlier stages. These questions can only be solved via an analysis of cell-types from early stages of AM interaction such as epidermal regions challenged with AM fungal signals or cortical regions harbouring PPAs.

Materials and Methods

Plant growth, AM fungal inoculation, and visualisation of AM fungal structures

*Medicago truncatula* Gaertn cv Jemalong genotype A17 seeds were surface-sterilized and scarified as reported by Hohnjec *et al.* (2003). Plants were grown in the climate chamber (humidity: 70 %; photosynthetic photon flux: 150 μmol m⁻² s⁻¹) at a 16 h light (23 °C) and 8 h dark (18 °C) regime. For subsequent *Medicago* GeneChip hybridizations using whole roots, plants were mycorrhized with AM
fungi under conditions of phosphate limitation (20 µM phosphate). In addition, nonmycorrhizal roots grown at 20 µM phosphate as well as nonmycorrhizal roots grown at 2 mM phosphate were generated as described previously (Hohnjec et al. 2005). Two different AM fungal inocula were used: *Glomus mosseae* granular AMF inoculum BEG 12 (Biorize R&D, Dijon, France), and *Glomus intraradices* Schenck and Smith DAOM197198 inoculum (Premier Tech Biotechnologies, Rivière-de-Loup, Québec, Canada), the latter having recently been reassigned to *Rhizophagus irregularis* (Blaszk., Wubet, Renker, and Buscot) C. Walker & A. Schüßler comb. nov. (Stockinger et al. 2009). At 28 days post inoculation (dpi) with AM fungi or at 28 days growth under nonmycorrhizal conditions, roots were harvested and frozen in liquid nitrogen. Randomly selected areas of mycorrhizal roots were stained for fungal colonization using the gridline intersection method according to McGonigle et al. (1990). Here, the percentage of root length colonization (RLC; scoring hyphae, spores, vesicles, or arbuscules) ranged from 60% to 80%, while the relative arbuscule frequency in colonized fragments varied between 60% and 75%.

To obtain mycorrhizal roots for embedding in Steedman’s wax, two-weeks old seedlings were mycorrhized by adding 15% (v/v) inoculum *Glomus intraradices* isolate 49 (Maier et al. 1995) produced in leek cultures (*Allium porrum* cv. Elefant) to the substrate. Mycorrhizal and non-mycorrhizal plants were fertilized with half-strength Hoagland’s solution containing 20 µM phosphate and an additional 2mM NH₄NO₃. RLC was checked regularly via ink-staining according to the protocol of Vierheilig et al. (1998) and gridline intersection counting according to McGonigle et al. (1990). 70-80% RLC turned out to be most convenient for the selection of arbuscule-containing and adjacent cortical cells. This level of colonization was usually reached at 21 dpi.

**Tissue embedding, tissue sectioning, and laser-microdissection**

Roots were embedded using the Steedman’s wax protocol (Gomez et al. 2009) with the following modifications: Eosin was already added in the first step of the ethanol series (75% (v/v) ethanol with 0.1% (v/v) Eosin y), the overnight fixation step in Farmer’s fixative and the overnight incubation step in 1:1 ethanol:wax were extended from 12 to 14 h, and root pieces were embedded in TurbOflow®II molds as well as cassettes (McCormick Scientific, Richmond, USA). Blocks were stored in vacuum-sealed plastic bags containing desiccant bags at 4°C.

Longitudinal root sections of 12 µm were obtained using a Hyrax M55 rotary microtome (Zeiss, München, Germany). Ribbons were spread on heat-sterilized glass slides and were stretched with DEPC-treated autoclaved water. Slides were dried for one hour in a hybridization oven at 32°C. Slides with sections were used on the same or the following two days and stored in vacuum-sealed plastic bags with desiccation bags at 4°C if necessary. Sections were de-waxed immediately before cell harvest by washing the slides with absolute ethanol several times at 38°C on a heating plate, until the wax was not visible anymore. Subsequently, slides were dried on the heating plate.

The P.A.L.M. Microbeam system with a Capmover (Zeiss, München, Germany) was used for laser-microdissection and pressure catapulting (LMPC). To collect cells, the “CloseCut and Auto-LPC” function was used according to the manufacturer’s instructions. Cells were collected into 500 µl adhesive caps (Zeiss, München, Germany) and stored directly at -80°C after the harvest was completed.
For each cell-type, four biological replicates were produced, based on distinct rounds of plant
cultivation and root embedding. Biological replicates consisted of three technical replicates of
approximately 1000 cells each, which were pooled after RNA isolation and amplification.

**RNA isolation and amplification**

Whole-root mycorrhizal samples were taken from frozen stocks, pooled, and ground using lysing
matrix D tubes (MP Biomedicals, Illkirch, France) in a FastPrep (MP Biomedicals, Illkirch, France) prior
to RNA extractions. Total RNA isolation and DNase I on-column digestion was performed via RNCasey
kits (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. RNA preparations were
quality-checked both via spectrophotometry (NanoDrop ND-1000, Peqlab, Erlangen, Germany) and
via capillary electrophoresis in RNA Nano chips (Agilent Bioanalyzer, Agilent, Böblingen, Germany),
as recommended by the manufacturers.

Total RNA was isolated from laser-microdissected cells using the RNaseasy Micro kit (Qiagen, Hilden,
Germany). 350 µl RLT buffer containing β-mercaptoethanol were added to each sample followed by a
30 min incubation at room temperature. The lysate was spun down for 5 min at 13.400 g, mixed 1:1
with ethanol absolute and transferred to the clean-up column. On-column DNase I digestion was
peformed according to the manufacturer’s instructions. RNA from laser-microdissected cells was
amplified using the TargetAmp 2-round aRNA amplification kit (Epicentre Biotechnologies, Madison,
USA), as specified by the manufacturer. Quantity and quality of total RNA as well as T7-amplified
aRNA was checked via capillary electrophoresis in RNA Pico and Nano Chips, respectively, using an
Agilent 2100 Bioanalyzer (Agilent, Böblingen, Germany).

**Medicago GeneChip hybridizations**

RNA was processed for use on Affymetrix (Santa Clara, CA, USA) GeneChip *Medicago* Genome
Arrays, according to the manufacturer’s GeneChip 3’ IVT Express kit user manual. Briefly, 100 ng of
total RNA with a RIN number (Agilent 2100 Bioanalyzer, Agilent, Böblingen, Germany) of at least
8.5 containing spiked-in poly-A⁺ RNA controls was used in a reverse transcription reaction (GeneChip
3’ IVT Express Kit; Affymetrix, Santa Clara, CA, USA) to generate first-strand cDNA. After second-
strand synthesis, double-stranded cDNA was used in a 16 h *in vitro* transcription (IVT) reaction to
generate aRNA (GeneChip 3’ IVT Express Kit; Affymetrix, Santa Clara, CA, USA). Size distribution of
in vitro transcribed aRNA and fragmented aRNA, respectively, was assessed via an Agilent 2100
Bioanalyzer (Agilent, Böblingen, Germany), using an RNA 6000 Nano Assay. 10 µg of fragmented
aRNA was added to a 300-µl hybridization cocktail also containing hybridization controls. 200 µl of the
mixture was hybridized on GeneChips for 16 h at 45°C. Standard post hybridization wash and double-
stain protocols (FS450_0001; GeneChip HWS kit; Affymetrix, Santa Clara, CA, USA) were used on an
Affymetrix GeneChip Fluidics Station 450. GeneChips were scanned on an Affymetrix GeneChip
scanner 3000 7G.

**Evaluation of data from Medicago GeneChip hybridizations**

Cel files obtained from *Medicago* GeneChip hybridizations were analysed using the Robin software
(http://mapman.gabipd.org/web/guest/robin). Normalization was performed across all GeneChips
using the Robust Multichip Average (RMA) algorithm. Intensity values calculated for each probe set were log2-transformed and averaged across all three biological replicates. Log2 differences between the conditions studied were evaluated statistically by applying a false discovery rate (FDR) correction for p-values implemented in Robin. Original annotations of probes from *Medicago* GeneChips were replaced by automated annotations as well as functional classifications generated via SAMS (Bekel *et al.* 2009) and Gene Ontology (GO) classifications (http://www.medicago.org/GeneChip). To visualize gene expression profiles, MapMan (Usadel *et al.* 2005) was used. Data from *Medicago* GeneChip hybridizations were related to *in silico* expression profiles using MediPlEx (Henckel *et al.* 2010), applying the "arbuscular mycorrhizal root libraries" preselection. Since *Medicago* GeneChips are based on gene models from EST and genomic sequences, the number of probe sets exceeds the number of genes represented to a certain extent. Nevertheless, we refer to genes instead of probe sets in this work for reasons of simplicity.

**real-time RT-PCR**

Primers were designed using the Primer3 web interface (Rozen and Skaletsky, 2000) according to the following criteria: product size range 100-150 bp; primer size min: 18 bp, opt: 21 bp, max: 24 bp; primer Tm min: 52°C, opt: 53°C, max: 55°C. If the position of the coding region was known, the amplicon was preferentially positioned in the 3' region of the gene or the 3' UTR. The primer pairs suggested by the primer3 program were blasted against the DFCI Medicago Gene Index (Quackenbush *et al.* 2001) to check for mispriming in known *M. truncatula* transcript sequences. The number of matching base pairs in off-target genes and especially matches at the 3' end of the primer were taken into account and the most suitable primer pair according to these parameters was chosen. All primer pairs were tested for performance and specificity with RNA from whole root tissue prior to using them for RNA from laser-microdissected samples. For some of the genes, primer pairs published by Gomez *et al.* (2009) were used. These are indicated in Supplemental Table S5, where all primer pairs are listed.

For the laser-microdissected samples, 50 ng of T7-amplified RNA (aRNA) were used for real-time RT-PCR, using the SensiMix™ SYBR one-Step kit (Bioline, Luckenwalde, Germany). RT-PCR conditions (Realplex cycler, Eppendorf, Germany) were as follows: 10 min 42°C, 10 min 95°C, 50 cycles (15 sec 95°C, 30 sec 55 °C, 30 sec 72°C), 15 sec 95°C, melting curve from 40-95°C. In addition to a melting curve analysis, all real-time PCR-products were separated on 2 % (w/v) agarose gels to check both for specificity and a correct amplification size. Resequencing of selected PCR-products was used to additionally confirm the specific amplification of the target gene. Expression results were averaged over 3 biological replicates. In case one replicate delivered a significantly deviant result from the other two, the measurement was repeated. If the difference persisted, the fourth replicate was included into the analysis and replaced the replicate in question, if this resulted in a more consistent expression pattern. The constitutive translation elongation factor gene *MtTef1α* (TC178258 in the DFCI *Medicago* Gene Index) was used for normalization across different conditions. *MtTef1α* expression was analyzed in 4 technical replicates, and the average value was used to calculate relative gene expression levels using the 2^(-ΔΔCT) value with ΔΔCT=CTgene-CTMtTef1α. Expression differences were analyzed for significance using the Student’s t-test incorporated in MS® Excel® 2007 (Microsoft® Corp., Seattle, USA).
For the whole root samples, 50 ng of total RNA were used for real-time RT-PCR, using the SensiMix™ SYBR one-Step kit (Bioline, Luckenwalde, Germany) according to the manufacturer’s instructions. The constitutive translation elongation factor gene \( MtTef1\alpha \) (TC178258 in the DFCI \textit{Medicago} Gene Index) was used for normalization across different conditions. All expression results were averaged over 3 biological replicates. Relative gene expression levels were calculated as described above.

**Construction and Histological Analysis of Transgenic Hairy Roots**

The promoters of two CAAT-box transcription factor genes represented by probe sets Mtr.51511.1.S1_at (referred to as \( MtCbf1 \)) and Mtr.16863.1.S1_at (referred to as \( MtCbf2 \)) were PCR-amplified with Phusion Hot Start DNA polymerase (Finnzymes, Biozym, Hessisch Oldendorf, Germany) from \textit{Medicago truncatula} Gaertn cv Jemalong genotype A17 genomic DNA, using gene-specific primers containing appropriate restriction sites. The amplified products covered the -1513/-3 region for \( MtCbf1 \) and the -1484/-6 region for \( MtCbf2 \) relative to the start codon. The promoter regions were cloned in front of the gusAint gene of pGUSint (Hohnjec \textit{et al.} 2003) using \textit{SphI}/\textit{SmaI} restriction sites for \( MtCbf1 \) and \textit{SphI}/\textit{EcoRI} restriction sites for \( MtCbf2 \). Whereas the promoter-GUS-fusion of \( MtCbf1 \) was excised using \textit{SpeI}, the promoter-GUS-fusion of \( MtCbf2 \) was excised using \textit{SalI}/\textit{StuI}. Both fragments were filled in with Klenow Polymerase, and cloned into the \textit{SmaI} site of the binary plasmid pRedRoot (Limpens \textit{et al.} 2004). The plasmids obtained were electroporated into \textit{Agrobacterium rhizogenes} ARqua1, and the resulting strains were used to generate hairy roots on \textit{M. truncatula} cv Jemalong A17 according to Vieweg \textit{et al.} (2004). Transgenic roots were identified using dsRed fluorescence (Leica MZ 10F, Leica Microsystems, Wetzlar, Germany) and were mycorrhized by adding 15% (v/v) inoculum \textit{Glomus intraradices} isolate 49 (Maier \textit{et al.} 1995) produced in leek cultures (\textit{Allium porrum} cv. Elefant) to the substrate. GUS assays were performed as described by Hohnjec \textit{et al.} (2003), without preheating of the samples. To obtain thin sections, GUS-stained roots were embedded in 5% agarose and cut using a Leica VT1000S vibratome (Leica, Wetzlar, Germany). Counterstaining of fungal structures was performed with Alexa Fluor® 488 WGA conjugate (Invitrogen, Darmstadt, Germany). Roots were bleached for 3 min in 10% KOH at 95°C, washed three times with water and stained in PBS-buffer containing 20 µg/ml Alexa Fluor® 488 WGA conjugate overnight. For staining of fungal structures in thin sections and staining of extraradical hyphae on the root surface, samples were directly transferred into PBS-buffer containing 20 µg/ml Alexa Fluor® 488 WGA conjugate and stained overnight. Photo documentation was performed with a Leica MZ 10F stereomicroscope (Leica Microsystems, Wetzlar, Germany), equipped with an Olympus XC50 camera (Olympus, Hamburg, Germany) and with a Zeiss Axio Observer Z1 microscope equipped with an AxioCam ICc1 (Zeiss, München, Germany).

**Accession number**

GeneChip data: Gene Expression Omnibus (GEO) accession number GSE32208.
Supplemental Material

Supplemental Figure S1: Identification of AM-specific and AM-enhanced transcription factor genes.
The expression of 19 genes encoding transcriptional regulators (including 2 genes from *Glomus intraradices*) presented in Fig. 5 was analysed in the *Medicago* Gene Expression Atlas (He *et al.* 2009). While six genes were specifically expressed in AM tissues (A), 13 were activated stronger in mycorrhizal vs non-mycorrhizal roots (B). In some cases, marked expression in additional symbiotic or non-symbiotic conditions was evident for the genes displayed in (B).

Supplemental Figure S2: Genes exclusively expressed in arbuscule-containing cells.
Gel-electrophoresis of the final real-time RT-PCR amplification products representing those genes classified as arbuscule-specific in Figure 5. Gene expression was measured in three biological replicates of two different cell-types: cortical cells from mycorrhized roots (CMR), and arbuscule-containing cells (ARB). All amplified fragments had the correct sizes. For Mtr.17343.1.S1_at, Mtr.1591.1.S1_at, and Mtr.31910.1.S1_at, an unspecific amplification product is visible in CMR_2, CMR_1, and CMR_3, respectively, which had a different size and melting temperature than the specific product. Footnotes are as indicated in Figure 5.

Supplemental Figure S3: Alignment of the coding sequences of the two CAAT-box binding transcription factor genes *MtCbf1* and *MtCbf2*.
Identical nucleotides are marked by asterisks. The start and stop codons are depicted in bold type.

Supplemental Figure S4: Alignment of promoter sequences of the two CAAT-box binding transcription factor genes *MtCbf1* and *MtCbf2*.
Sequences shown represent the regions from -1513 to +3 for *MtCbf1* and from -1486 to +3 for *MtCbf2*. Identical nucleotides are marked by asterisks. The start codon is depicted in bold type.

Supplemental Table S1
Gene expression in *Medicago truncatula* roots in response to *Glomus intraradices* colonization (28 dpi, at 20 µM phosphate), *Glomus mosseae* colonization (28 dpi, at 20 µM phosphate), and a 28 day treatment with 2 mM phosphate. Roots grown for 28 days in the presence of 20 µM phosphate were used as common controls.

Supplemental Table S2
*Medicago truncatula* genes co-activated in response to *Glomus intraradices* and *Glomus mosseae* colonization. The individual sheets contain selected subsets of genes, as explained in the Supplemental file.

Supplemental Table S3
AM-activated *Medicago truncatula* genes encoding membrane transporters.
Supplemental Table S4
AM-activated *Medicago truncatula* genes encoding signaling-related proteins.

Supplemental Table S5
Real time RT-PCR primers used in this study and size of predicted PCR-products.

Acknowledgements
We are grateful to Karen Gomez and Maria Harrison (Boyce Thompson Institute for Plant Research, Tower Road, Ithaca, USA) for sharing protocols and for helpful discussions on laser-microdissection. Excellent bioinformatics support was provided by Kolja Henckel (Bioinformatics Resource Facility, Center for Biotechnology, Bielefeld University).

Literature Cited


the infected cells of root nodules and in the arbuscule-containing cells of mycorrhizal roots from different legume and nonlegume plants. *Mol Plant Microbe Interact* **17**: 62-69


**Figure Legends**

**Fig. 1**: Transcriptional response of *M. truncatula* roots to a colonization with different AM fungi and to a treatment with 2 mM phosphate.

*M. truncatula* roots were inoculated with *G. intraradices* and *G. mosseae* for 28 days under conditions of phosphate limitation (20 µM phosphate). Alternatively, roots were grown for 28 days in the presence of 2 mM phosphate. Genes significantly upregulated 2-fold at an FDR-corrected p-value of p<0.05 in relation to control roots grown under conditions of phosphate limitation were compared to identify co-regulation of expression. Numbers indicate genes activated in different conditions. Diagrams were drawn using Venny (http://bioinfogp.cnb.csic.es/tools/venny/index.html).

**Fig. 2**: Cellular functions of *M. truncatula* genes activated in mycorrhizal roots.

All 512 *M. truncatula* genes co-induced at least two-fold at an FDR-corrected p-value of p<0.05 in response to the AM fungi *G. intraradices* and *G. mosseae* that were not induced by a treatment with 2 mM phosphate (Supplemental Table S2) were grouped into functional categories. The number of genes allocated to each functional category is indicated. The bars are coloured as follows: black, functional categories studied by laser-microdissection; dark grey, AM-related gene families; light grey, other functional categories.

**Figure 3**: Laser-microdissection of three specific cell-types from *M. truncatula* roots.

Root areas designated for cell harvest are marked with a green line and blue dots. Along the line the laser dissects the cells from the surrounding tissue, while dots represent single catapulting events. A to C: Longitudinal section of a non-mycorrhized root used for the collection of cortical cells from control roots (CCR). A and B: Section before and after selection of CCR for laser-microdissection. C: Section after laser-microdissection of CCR. D: View into the collection-tube showing typical flakes of harvested cells (in this case arbuscule-containing cells). E to H: Longitudinal sections of mycorrhized roots displaying chains of arbuscules at different developmental stages and fungal hyphae growing in the apoplast of outer cortical cells. E and F: Section before and after selection of cortical cells from mycorrhized roots (CMR). The harvested area was extended to inner cortical cells in case no arbuscules were visible in these cells. Fungal hyphae are present in the apoplast (blue arrows). G and
Section before and after selection of arbuscule-containing cells (ARB). Only cells harbouring mature arbuscules filling up the whole lumen were harvested. These cells could be easily distinguished from those containing young or severely degraded arbuscules (yellow arrows in E). Scale bars represent 300 µm for D and 150 µm for all other panels.

**Figure 4: Detection of AM marker gene transcripts in laser-microdissected cell-types.**
Marker gene expression was measured by real-time RT-PCR in four biological replicates of three different cell-types: cortical cells from non-mycorrhizal control roots (CCR), cortical cells from mycorrhizal roots containing fungal hyphae (CMR), and arbuscule-containing cells (ARB). A and B: Gel-electrophoresis of the final real-time RT-PCR amplification products representing the control gene *MtTef*α and six AM marker genes. All amplified fragments had the correct sizes. Note that *MtPT4* transcripts were not detected in CMR. C: Real-time RT-PCR measurement of five AM marker genes induced in ARB in comparison to CMR. Expression values are displayed as log2 mean values of all four biological replicates. Numbers and bars represent fold-induction and standard errors, respectively. Asterisks indicate significance levels of a Student’s t-test on the expression values in the two different cell-types: *=p<0,1; **=p<0,05; ***=p<0,005. Abbreviations: CCR, cortical cells from non-mycorrhizal control roots; CMR, cortical cells from mycorrhizal roots; ARB, arbuscule-containing cells; *MtTef*α, transcriptional elongation factor α; *MtPT4*, phosphate transporter 4; *MtBcp1*, blue copper protein 1, *MtScp1*, serine carboxypeptidase 1; *MtLec5*, lectin 5, *MtGlp1*: germin-like protein 1; *MtHa1*, H+-ATPase 1.

**Figure 5: Cell-type specific expression of genes activated in mycorrhizal roots.**
Real-time RT-PCR measurement of gene expression in three biological replicates of the laser-microdissected cell-types CMR (cortical cells from mycorrhizal roots) and ARB (arbuscule-containing cells). Differences in transcription are indicated by different shades of grey (legend see below). p-values represent significance levels of a Student’s t-test on the expression values in the two different cell-types. In addition, the log2 expression ratios of gene expression in roots mycorrhizized with *Glomus intraradices* vs. non-mycorrhizal roots (Supplemental Table S1) are shown. Footnotes: (1): One of the three biological replicates was replaced by replicate four, to obtain three gene-specific PCR-products of the correct size or a consistent expression pattern. (2): A gene-specific PCR product could only be obtained for two out of three biological replicates of ARB. (3): A gene-specific PCR product could only be obtained for two out of three biological replicates of CMR. (4): A gene-specific PCR product was obtained for only one out of three biological replicates of CMR.

**Figure 6: Genes differentially expressed in cortical and arbuscule-containing cells.**
Real-time RT-PCR measurement of the expression of selected genes classified as CMR- or ARB-induced in Figure 5. Gene expression is displayed as the log2 mean value of three biological replicates. Numbers and bars represent fold-induction and standard errors, respectively. Different genes are labeled with abbreviated GeneChip probe IDs (compare Figure 5). Asterisks indicate significance levels of a Student’s t-test on expression values in the two different cell-types: *=p<0,1; **=p<0,05; ***=p<0,005. Abbreviations: CMR: cortical cells from mycorrhizal roots; ARB: arbuscule-containing cells.
Figure 7: Activity of the MtCbfl promoter in M. truncatula mycorrhizal roots.

Figure 8: Activity of the MtCbfl2 promoter in M. truncatula mycorrhizal roots.
A, B, C, G, I, K, L and M: Light micrographs of M. truncatula mycorrhizal roots expressing the gusA/int gene under the control of the MtCbfl2 promoter. D, E, F, H, N, O, and P: Corresponding fluorescence micrographs showing counterstaining of fungal structures with Alexa Fluor® 488 WGA conjugate at exactly the same root position. In A to H, whole roots are shown, whereas I to P show 60 µm thin sections. B and E represent enlarged regions of the root shown in A and D. K, L, N and O represent enlarged regions of the root shown in I. A, B, D and E: Promoter activity during early stages, with fungal hyphae being just attached to the root epidermis. In this case, the hyphae in contact to the root surface (indicated by arrows) emerged from a highly mycorrhized leek root attached to the M. truncatula root. C and F: Promoter activity in a young infection unit. G and H: Promoter activity in a region with an expanding infection unit. In contrast to MtCbfl, no strong MtCbfl2 activity can be observed in epidermal cell layers in this stage, hence the strong GUS-staining in the cortex is visible from the outside. I, K, L, N and O: Promoter activity in a densely colonized root. M and P: Enlargement of a group of arbuscules. Scale bars represent 200 µm for A, D, G, H, and I; 100 µm for C and F; 50 µm for B, E, K, L, N, and O; 20 µm for M and P.

Figure 9: Expression of MtCbfl1 and MtCbfl2 in specific cell-types and during a time course of mycorrhization.
A and B: Real-time RT-PCR measurement of MtCbfl1 and MtCbfl2 expression in arbuscule-containing cells (ARB) and adjacent cortical cells colonized by fungal hyphae (CMR). Gene expression is displayed as the log2 mean value of three biological replicates. Bars represent standard errors. C to F: Promoter activity of MtCbfl1 during a time course covering 7 to 28 dpi of mycorrhization, represented by roots displaying typical GUS staining patterns for each time point. G to K: Promoter activity of MtCbfl2 during a time course covering 7 to 28 dpi of mycorrhization, represented by roots displaying typical GUS staining patterns for each time point. L: Real-time RT-PCR measurement of MtCbfl1, MtCbfl2, MtPt4, and MtBcp1 in M. truncatula roots during the 7 to 28 dpi time course of mycorrhization. Bars represent standard errors.
Figure 10: Schematic summary of fungal and plant gene expression patterns during four different stages of the AM symbiosis.

Proteins encoded by the genes identified are grouped according to functional categories. Fungal gene products are listed in orange, plant gene products in green boxes. The total number of genes with identical annotations is indicated in brackets. Note that it remains to be elucidated to what extent genes identified as expressed in arbuscule-containing as well as in the adjacent cortical cells colonized by fungal hyphae are already active during stages I and II.
Tables

Table 1

*M. truncatula* AM marker genes activated in roots colonized with *G. intraradices* and *G. mosseae*.

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<th>FDR-p</th>
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Probe IDs of *Medicago* GeneChips are referenced to the corresponding DFCI *Medicago truncatula* Gene Index IDs (release 10) and to *M. truncatula* gene names from the literature. Log2 ratios of gene expression for *G. intraradices*-colonized (Gi-Myc), *G. mosseae*-colonized (Gm-Myc), and 2 mM phosphate-treated (2 mM-P) roots, all measured against roots grown at 20 µM phosphate, are given. Whereas all 15 AM marker genes are significantly upregulated in the AM roots used for expression profiling, none of them is activated in roots treated with 2 mM phosphate. FDR-corrected p-values (FDR-p) are indicated. References for the AM-induced genes identified are as follows: *MtPt4* (Javot et al. 2007), *MtMyb1* (Liu et al. 2003), *MtTi1* (Grunwald et al. 2004), *MtLec7* (Frenzel et al. 2005), *MtGst1* (Wulf et al. 2003), *MtBcp1* (Hohnjec et al. 2005), *MtLec5* (Frenzel et al. 2004), *MtGlp1* (Doll et al. 2003), *MtHa1* (Krajinski et al. 2002), *MtScp1* (Liu et al. 2003), *MtVapyrin* (Pumplin et al. 2009), *MtSbtM1* (Takeda et al. 2011), *MtZip7* (Burleigh et al. 2003), *MtAnn2* (Manthey et al. 2004), and *MtDxs2* (Floss et al. 2008).
Table 2
Overview of 25 *M. truncatula* AM-induced genes encoding transcriptional regulators.

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Probe IDs of *Medicago* GeneChips are referenced to *M. truncatula* gene names, where applicable. Log2 ratios of gene expression for *G. intraradices*-colonized (Gi-Myc), *G. mosseae*-colonized (Gm-Myc), and 2 mM phosphate-treated (2 mM-P) roots, all measured against roots grown at 20 µM phosphate, are given. FDR-corrected p-values (FDR-p) are indicated. References for the AM-induced transcription factor genes identified are as follows: *MtErn1* (Middleton et al. 2007), *MtErn2* (Adriankaja et al. 2007), *MtMyb1* (Liu et al. 2003), *MtCbf1* (this work), and *MtCbf2* (this work).
Fig. 1: Transcriptional response of *M. truncatula* roots to a colonization with different AM fungi and to a treatment with 2 mM phosphate.

*M. truncatula* roots were inoculated with *G. intraradices* and *G. mosseae* for 28 days under conditions of phosphate limitation (20 μM phosphate). Alternatively, roots were grown for 28 days in the presence of 2 mM phosphate. Genes significantly upregulated 2-fold at an FDR-corrected p-value of p<0.05 in relation to control roots grown under conditions of phosphate limitation were compared to identify co-regulation of expression. Numbers indicate genes activated in different conditions. Diagrams were drawn using Venny (http://bioinfogp.cnb.csic.es/tools/venny/index.html).
Fig. 2: Cellular functions of *M. truncatula* genes activated in mycorrhizal roots.

All 512 *M. truncatula* genes co-induced at least two-fold at an FDR-corrected p-value of p<0.05 in response to the AM fungi *G. intraradices* and *G. mosseae* that were not induced by a treatment with 2 mM phosphate (Supplemental Table S2) were grouped into functional categories. The number of genes allocated to each functional category is indicated. The bars are coloured as follows: black, functional categories studied by laser microdissection; dark grey, AM-related gene families; light grey, other functional categories.
Figure 3: Laser-microdissection of three specific cell-types from *M. truncatula* roots.

Root areas designated for cell harvest are marked with a green line and blue dots. Along the line the laser dissect the cells from the surrounding tissue, while dots represent single catapulting events. **A-C:** Longitudinal section of a non-mycorrhized root used for the collection of cortical cells from control roots (CCR). **A&B:** Section before and after selection of CCR for laser-microdissection. **C:** Section after laser-microdissection of CCR. **D:** View into the collection-tube showing typical flakes of harvested cells (in this case arbuscule-containing cells). **E-H:** Longitudinal sections of mycorrhized roots displaying chains of arbuscules at different developmental stages and fungal hyphae growing in the apoplast of outer cortical cells. **E&F:** Section before and after selection of cortical cells from mycorrhized roots (CMR). The harvested area was extended to inner cortical cells in case no arbuscules were visible in these cells. Fungal hyphae are present in the apoplast (blue arrows). **G&H:** Section before and after selection of arbuscule-containing cells (ARB). Only cells harbouring mature arbuscules filling up the whole lumen were harvested. These cells could be easily distinguished from those containing young or severely degraded arbuscules (yellow arrows in E). Scale bars represent 300 μm for D and 150 μm for all other panels.
Figure 4: Detection of AM marker gene transcripts in laser-microdissected cell-types. Marker gene expression was measured by real-time RT-PCR in four biological replicates of three different cell-types: cortical cells from non-mycorrhizal control roots (CCR), cortical cells from mycorrhizal roots containing fungal hyphae (CMR), and arbuscule-containing cells (ARB). A and B. Gel-electrophoresis of the final real-time RT-PCR amplification products representing the control gene MtTefa and six AM marker genes. All amplified fragments had the correct sizes. Note that MtPT4 transcripts were not detected in CMR. C. Real-time RT-PCR measurement of five AM marker genes induced in ARB in comparison to CMR. Expression values are displayed as log2 mean values of all four biological replicates. Numbers and bars represent fold-induction and standard errors, respectively. Asterisks indicate significance levels of a Student’s t-test on the expression values in the two different cell-types: *p<0.1; **p<0.05; ***p<0.005. Abbreviations: CCR, cortical cells from non-mycorrhizal control roots; CMR, cortical cells from mycorrhizal roots; ARB, arbuscule-containing cells; MtTefa, transcriptional elongation factor α; MtPT4, phosphate transporter 4; MtBcp1, blue copper protein 1; MtScp1; secrete carboxypeptidase 1; MtLec5, lectin 5; MtGlp1: germin-like protein 1; MtHa1; H+-ATPase 1.
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**Figure 5:** Cell-type specific expression of genes activated in mycorrhizal roots. Real-time RT-PCR measurement of gene expression in three biological replicates of the laser-microdissected cell-types CMR (carpellary cortex), ARB (archeocyte containing cells). Differences in expression are indicated by different shades of green (Student’s t-test). p-values represent significance levels of a Student’s t-test on the expression values in two different cell types. In addition, the log2 ratio expresses the ratio of gene expression in roots mycorrhizated with Glomus intraradices vs. non-mycorrhizal roots (Supplemental Table S1). No significant difference was found. One of the three biological replicates was replaced by replicate four to obtain three gene-specific PCR products for each condition of the correct size or a consistent expression pattern. (1): A gene-specific PCR product could only be obtained in two of the three biological replicates of CMR. (2): A gene-specific PCR product could only be obtained for one of the three biological replicates of CMR. (3): A gene-specific PCR product could only be obtained for two of three biological replicates of CMR. (4): A gene-specific PCR product was obtained for only one of three biological replicates of CMR. (5): No significant difference was found.
Figure 6: Genes differentially expressed in cortical and arbuscule-containing cells.

Real-time RT-PCR measurement of the expression of selected genes classified as CMR- or ARB-induced in Figure 5. Gene expression is displayed as the log2 mean value of three biological replicates. Numbers and bars represent fold-induction and standard errors, respectively. Different genes are labeled with abbreviated GeneChip probe IDs (compare Figure 5). Asterisks indicate significance levels of a Student’s t-test on expression values in the two different cell-types: *=p<0.1; **=p<0.05; ***=p<0.005. Abbreviations: CMR: cortical cells from mycorrhizal roots; ARB: arbuscule-containing cells.
Figure 7: Activity of the MtCbf1 promoter in *M. truncatula* mycorrhizal roots. A, B, D, E, F, K, L, and M: Light micrographs of *M. truncatula* mycorrhizal roots expressing the gusA/mg2 gene under the control of the MtCbf1 promoter. C, G, H, I, N, O, P: Corresponding fluorescence micrographs showing counterstaining of fungal structures with Alexa Fluor® 488 WGA conjugate at exactly the same root position. In A, B, C, F, and I, whole roots are shown; whereas D, E, G, H, and K to P show 60 μm thin sections. E, H, L, and O represent enlarged regions of the roots shown in D, G, K, and N, respectively. A to C: Promoter activity during early AM stages, with fungal hyphae being just attached to the root epidermis. D, E, G, and H: Promoter activity in a young infection unit. F and I: Strong epidermal promoter activity in a region with an expanding infection unit. K, L, N, and O: Promoter activity in a densely colonized root. M and P: Enlargement of a single arbuscule. Scale bars represent 500 μm for A; 200 μm for F, I, K and N; 100 μm for B, C, D and G; 50 μm for L and O; 20 μm for E, H, M and P.
Figure 8: Activity of the MtCbf2 promoter in *M. truncatula* mycorrhizal roots.

A, B, C, G, I, K, L, and M: Light micrographs of *M. truncatula* mycorrhizal roots expressing the gusAint gene under the control of the MtCbf2 promoter. D, E, F, H, N, O, and P: Corresponding fluorescence micrographs showing counterstaining of fungal structures with Alexa Fluor® 488 WGA conjugate at exactly the same root position. In A to H, whole roots are shown, whereas I to P show 60 µm thin sections. B and E represent enlarged regions of the root shown in A and D. K, L, N and O represent enlarged regions of the root shown in I. A, B, D and E: Promoter activity during early stages, with fungal hyphae being just attached to the root epidermis. In this case, the hyphae in contact to the root surface (indicated by arrows) emerged from a highly mycorrhizized leek root attached to the *M. truncatula* root. C and F: Promoter activity in a young infection unit. G and H: Promoter activity in a region with an expanding infection unit. In contrast to MtCbf1, no strong MtCbf2 activity can be observed in epidermal cell layers in this stage, hence the strong GUS-staining in the cortex is visible from the outside. I, K, L, N and O: Promoter activity in a densely colonized root. M and P: Enlargement of a group of arbuscules. Scale bars represent 200 µm for A, D, G, H, and I; 100 µm for C and F; 50 µm for B, E, K, L, N, and O; 20 µm for M and P.
Figure 9: Expression of MtCbf1 and MtCbf2 in specific cell-types and during a time course of mycorrhization.
A and B: Real-time RT-PCR measurement of MtCbf1 and MtCbf2 expression in arbuscule-containing cells (ARB) and adjacent cortical cells colonized by fungal hyphae (CMR). Gene expression is displayed as the log2 mean value of three biological replicates. Bars represent standard errors. C to F: Promoter activity of MtCbf1 during a time course covering 7 to 28 dpi of mycorrhization, represented by roots displaying typical GUS staining patterns for each time point. G to K: Promoter activity of MtCbf2 during a time course covering 7 to 28 dpi of mycorrhization, represented by roots displaying typical GUS staining patterns for each time point. L: Real-time RT-PCR measurement of MtCbf1, MtCbf2, MtPt4, and MtBcp1 in M. truncatula roots during the 7 to 28 dpi time course of mycorrhization. Bars represent standard errors.
Figure 10: Schematic summary of fungal and plant gene expression patterns during four different stages of the AM symbiosis.

Proteins encoded by the genes identified are grouped according to functional categories. Fungal gene products are listed in orange, plant gene products in green boxes. The total number of genes with identical annotations is indicated in brackets. Note that it remains to be elucidated to what extent genes identified as expressed in arbuscule-containing as well as in the adjacent cortical cells colonized by fungal hyphae are already active during stages I and II.