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Cell-specific pathways recruited for symbiotic nodulation in the *Medicago truncatula* legume

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Abstract

Medicago truncatula is a model legume species that has been studied for decades to understand the symbiotic relationship between legumes and soil bacteria collectively named rhizobia. This symbiosis called nodulation is initiated in roots with the infection of root hair cells by the bacteria, as well as the initiation of nodule primordia from root cortical, endodermal, and pericycle cells, leading to the development

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of a new root organ, the nodule, where bacteria fix and assimilate the atmospheric dinitrogen for the benefit of the plant. Here, we report the isolation and use of the nuclei from mock and rhizobia-inoculated roots for the single nuclei RNA-seq (sNucRNA-seq) profiling to gain a deeper understanding of early responses to rhizobial infection in *Medicago* roots. A gene expression map of the *Medicago* root was generated, comprising 25 clusters, which were annotated as specific cell types using 119 *Medicago* marker genes and orthologs to *Arabidopsis* cell-type marker genes. A focus on root hair, cortex, endodermis, and pericycle cell types, showing the strongest differential regulation in response to a short-term (48 h) rhizobium inoculation, revealed not only known genes and functional pathways, validating the sNucRNA-seq approach, but also numerous novel genes and pathways, allowing a comprehensive analysis of early root symbiotic responses at a cell type-specific level.

Keywords: *Medicago* root, single-cell transcriptomic, rhizobium, nodule initiation, root hair cells, cortical cells

Introduction

Legumes symbiotically interact with nitrogen-fixing soil bacteria collectively named rhizobia. The molecular, physiological, and cellular responses of this symbiosis named nodulation have been extensively studied over the past decades (Roy et al., 2019). Legume nodulation is a complex biological process that requires the activation of temporally and spatially coordinated programs in a limited number of root cells. Briefly, legume nodulation is initiated by the perception of the rhizobial lipochitooligosaccharide Nod factors (NFs) and the subsequent infection of plant root hair cells by rhizobia. Concomitantly, a nodule primordium emerges. In *Medicago truncatula*, a legume species generating indeterminate nodules, these primordia are initiated from cell divisions within the root inner cortex, endodermis, and pericycle layers, and an apical meristem is then established and maintained during the entire life of the nodule. Rhizobia infect the developing root nodule primordia, differentiate into bacteroids, and fix and assimilate the atmospheric dinitrogen allowing a steady supply of nitrogen for the plant.

Functional genomic studies revealed the role of many legume genes controlling the early stages of nodulation, notably in root hair cells (trichoblasts) where the initial microsymbiont perception and the rhizobial infection take place, but also to a lower extent in pericycle and cortex inner root cell layers where nodule organogenesis initiates

(Roy et al., 2019). Several *M. truncatula* genes have been functionally characterized to control the infection of the root epidermis. Among them, Nodule Pectate Lyase (*MtNPL*) (Xie et al., 2012), a gene encoding a cell wall degrading enzyme required for the initiation of infection threads in curled root hairs, Lumpy Infections (*MtLIN*) (Kiss et al., 2009; Liu et al., 2019a), which encodes a putative E3 ligase, Rhizobium Directed Polar Growth (*MtRPG*) (Arrighi et al., 2008), a gene encoding a protein with a coiled-coiled domain, *MtFLOT4* (Haney and Long, 2010), *MtVPY* (Murray et al., 2011), NADPH oxidase/respiratory burst oxidase homologs (e.g., Rboh) (Montiel et al., 2016), and *MtCBS1*, a gene encoding a Cystathionine- β -Synthase-like Domain-Containing Protein (Sinharoy et al., 2016), are all upregulated in response to rhizobium inoculation. In addition, other rhizobium-upregulated genes participate in the NF signaling pathway, such as LysM receptors that perceive NF bacterial signals. These genes belong to the LysM receptor kinase and LYK-related gene families (*MtLYK* and *MtLYR*, respectively; e.g., *MtNFP* [NF Perception]) (Gough et al., 2018), Does not Make Infections (*MtDMI1*, 2 and 3) genes (Endre et al., 2002; Anéet al., 2004; Gleason et al., 2006), Plant U-box protein 1 and 2 (*MtPUB1* and 2) involved in protein degradation (Mbengue et al., 2010; Liu et al., 2018) NF Hydrolase 1 (*MtNFH1*) (Cai et al., 2018), which controls NF degradation, Interacting with DMI3 (*MtIPD3*) (Messinese et al., 2007), and various transcription factor (TF) genes (e.g., Nodulation Signaling Pathway 1/2 [*MtNSP1*] [Kaló et al., 2005; Smit et al., 2005], Nuclear factor-YA1 [*MtNF-YA1*] [Combier et al., 2006], Ethylene Response Factor [*MtERN1*] Required for Nodulation [Andriankaja et al., 2007; Middleton et al., 2007], and *MtNIN* [Nodule Inception] [Schauser et al., 1999]).

To date, a limited number of studies have highlighted the regulatory mechanisms controlling the response of inner root cell layers to rhizobial inoculation (i.e., pericycle, endodermis, and cortex). Among them, cytokinin signaling is necessary and sufficient to promote the initiation of nodule primordia and inhibit the response of epidermal root cells to rhizobia and NFs (Gonzalez-Rizzo et al., 2006; Murray et al., 2007; Plet et al., 2011; Boivin et al., 2016; Jardinaud et al., 2016; Gamas et al., 2017; Lin et al., 2021). Genes controlling root development were also shown to be recruited for nodule development, such as *MtPLT* (Plethora) (Franssen et al., 2015), Knotted homeobox (*Mt-KNOX*) (Di Giacomo et al., 2017), LOB Binding Domain 16 (*MtLBD16*)

(Schiessl et al., 2019; Soyano et al., 2019), ShortRoot (*MtSHR*) (Dong et al., 2021), Scarecrow (*MtSCR*) (Dong et al., 2021), and *MtNOOT1* (Shen et al., 2019). On some occasions, genes were shown to have a dual function to promote root epidermal infection and nodule initiation at the level of the pericycle and cortical cells, such as *MtNIN* (Liu et al., 2019b).

To gain a more accurate picture of the symbiotic transcriptional programs controlling the root hair signal perception and infection, transcriptomic studies were conducted on populations of isolated legume root hair cells using root hair shaving or laser dissection (Libault et al., 2009; Breakspear et al., 2014). Similarly, -omics analyses targeting the zone of emerging nodules revealed the transcriptomic programs controlling the initiation and development of nodule primordia (Lohar et al., 2005; Larrainzar et al., 2015; van Zeijl et al., 2015; Schiessl et al., 2019). While valuable, these approaches suffer from the cellular heterogeneity of the isolated root hair populations (i.e., a mixture of unresponsive, responsive but uninfected, and infected root hair cells) (Bhuvaneswari et al., 1981), and from the cellular complexity of the root. For instance, as a reflection of the cellular heterogeneity of the root hair population, it has been estimated that only 1%–5% of the root hair cells are infected by rhizobia (Nutman, 1959).

While these approaches increased the resolution of plant transcriptomic analyses, they have been recently superseded by the emergence of single-cell RNA sequencing (scRNA-seq) and single-nucleus RNA-seq (sNucRNA-seq) transcriptomic technologies. Here, we report the use of the sNucRNA-seq technology on *M. truncatula* roots mock-inoculated or inoculated with its symbiont, *Ensifer* (*Sinorhizobium*) *meliloti*, to precisely capture the transcriptomic programs induced during the early stages of the nodulation process in each cell type composing the *M. truncatula* root. The establishment of a single-cell resolution transcriptomic map of the *M. truncatula* root allowed for the characterizing of the transcriptomic response of the *M. truncatula* root hair, cortical, endodermal, and pericycle cells at an early stage (48 h) after *E. meliloti* inoculation. The transcriptomic analysis of these different cell types revealed the dynamic regulation of *M. truncatula* genes in response to rhizobial infection, including many novel genes and functional pathways, as well as the differential recruitment of previously known nodulation-related and hormonal genes depending on cell types.

Results

Establishment of a transcriptional map of the rhizobium-inoculated *M. truncatula* root at a single-cell resolution

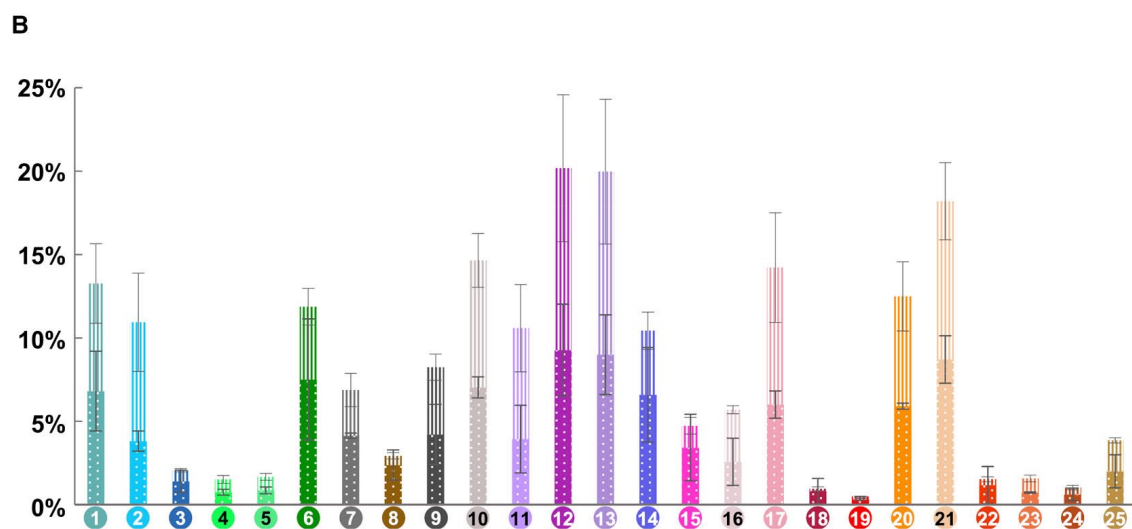
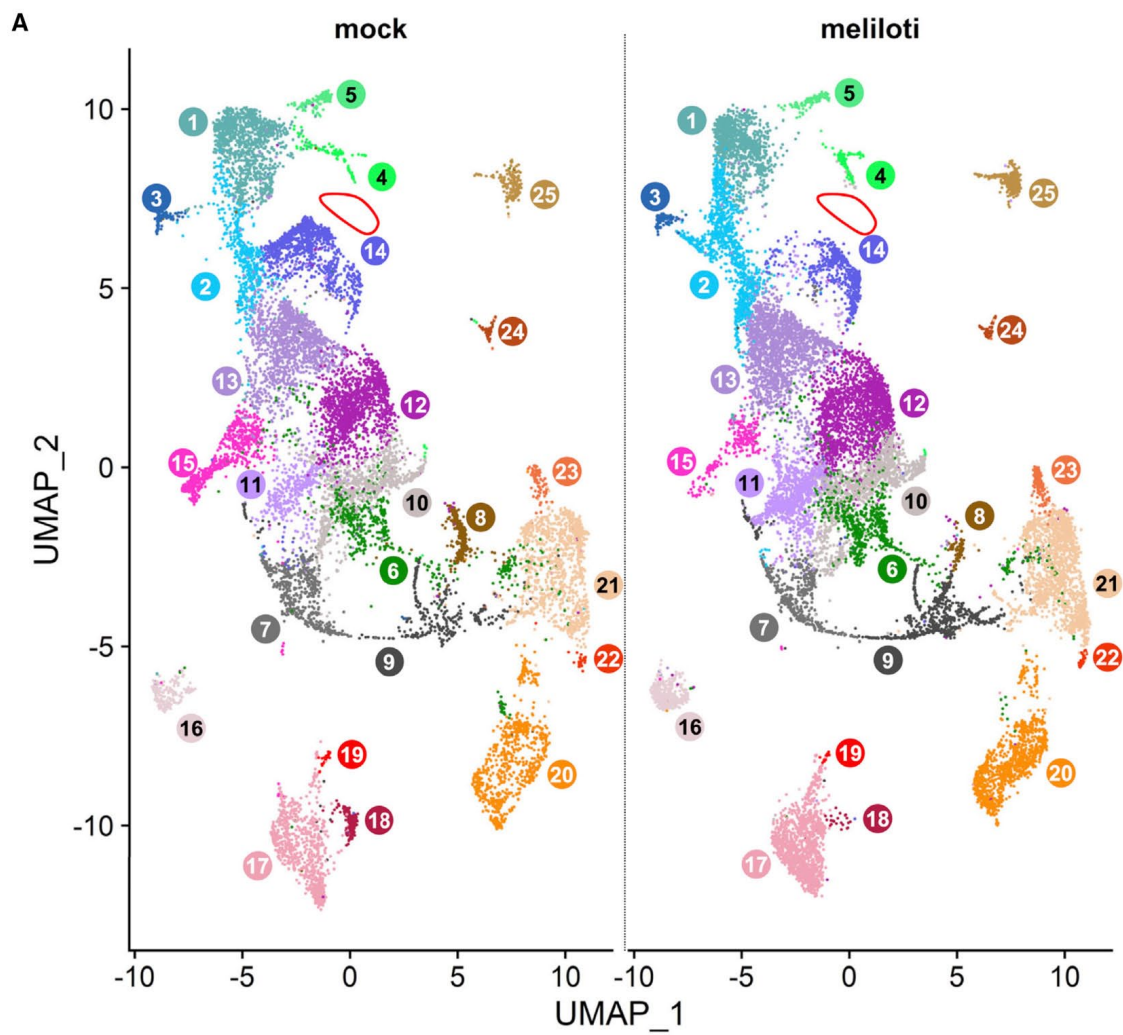
Isolated plant protoplasts and nuclei have been successfully used to establish single-cell resolution transcriptomes notably from *Arabidopsis* root cells (Denyer et al., 2019; Jean-Baptiste et al., 2019; Ryu et al., 2019; Shulze et al., 2019; Zhang et al., 2019; Farmer et al., 2021). The high correlation between cellular, nuclear, and whole root transcriptomes (Farmer et al., 2021) supports the biological relevance of both scRNA-seq and sNucRNA-seq approaches. However, the nuclear-based transcriptomic technology has unique advantages compared with the protoplast-based transcriptomic technology, including the ease of nuclei isolation from various plant species and organs, and the limited induction of stress-related genes (i.e., in contrast, protoplastization leads to the induction of hundreds of stress-responsive genes) (Birnbaum et al., 2003; Denyer et al., 2019). Therefore, we conducted sNucRNA-seq experiments to characterize the transcriptomic profiles of the different cell-types composing the *Medicago* root apex and their early response to rhizobial inoculation.

Shortly, *Medicago* seedlings were inoculated 4 days after germination with water or a bacterial suspension of *E. meliloti* (OD_{600nm} = 0.1). Forty-eight hours after rhizobium inoculation, root tips including fully elongated root hair cells were collected and committed to nuclei isolation (see methods for details). Three independent *E. meliloti*-inoculated and three independent mock-inoculated sNucRNA-seq libraries were generated using the 103 Genomics Chromium platform. To establish transcriptomic profiles of *Medicago* root cells, we applied a pre-mRNA strategy using Cell Ranger (103 Genomics) to map sequencing reads against transcripts and introns of the v1.8 annotation of the MtrunA17r5.0-ANR genome (Pecrix et al., 2018). This first step was followed by the removal of the ambient transcriptomic noise and the detection and removal of doublets (i.e., two nuclei encapsulated into the same reaction volume) (see methods). To support the quality of the libraries, we analyzed the distribution of the number of expressed genes per nucleus and observed an expected normal distribution (Supplemental Figure 1). To remove outliers and low-quality

nuclei, we applied a 95% confidence interval to the normal distribution of the six integrated libraries (Supplemental Figure 1). Taken together, the transcriptomes of 15 854 *E. meliloti*-inoculated and 12 521 mock-inoculated *M. truncatula* root nuclei were further analyzed (Supplemental Table 1), with a median value of 1,053 expressed genes per nucleus and a total of 31,307 expressed protein-coding genes detected (70.2% of the 44 615 Medicago -protein-coding genes) (Pecrix et al., 2018). Considering that the number of Arabidopsis and Medicago expressed genes per nucleus are similar (i.e., 1,124 expressed genes per Arabidopsis nucleus), the percentage of Medicago protein-coding genes expressed is less compared with Arabidopsis (i.e., 89.4% of 27,420 protein-coding genes) (Farmer et al., 2021). Such a limited percentage might be a consequence of the neo- or subfunctionalization of Medicago genes following the whole-genome duplication that occurred 58 million years ago, or the result of an over-estimation of the number of protein-coding genes in Medicago compared with the reference Arabidopsis genome.

Using the Seurat package, we normalized and integrated 28,375 nuclei transcriptomes before applying the uniform manifold approximation and projection (UMAP) technique to cluster the nuclei according to their transcriptomic profiles. Twenty-five clusters were identified (**Figure 1A**) the dataset can be interrogated using the https://shinycell.legumeinfo.org/medtr.A17.gnm5.ann1_6.expr.Cervantes-Perez_Thibivilliers_2022/ web interface) (Ouyang et al., 2021), for which the percentage of nuclei per cluster was not statistically different between inoculated and mock-inoculated conditions (Student t-test > 0.05; Figure 1B; Supplemental Table 1). While *E. meliloti* inoculation did not lead to changes in the number of clusters or the overall topography of the UMAP projection, we repetitively noticed a local modification in the distribution of a subpopulation of cluster #2 nuclei (Figure 1A, red circle). These results highlight that the *E. meliloti* inoculation induces significant changes in the transcriptome of these Medicago root cells.

Figure 1. Single-nuclei RNA-seq of the *M. truncatula* roots reveals 25 different root clusters. (A) UMAP clustering of *M. truncatula*/*E. meliloti*- and mock-inoculated root nuclei according to their transcriptomic profiles. While the overall topography of these two UMAPs is well conserved, subtle differences are observed (e.g., highlighted in red for cluster #2). (B) Percentage of *E. meliloti* (dash bars) and mock-inoculated (solid bars) nuclei allocated in the 25 clusters composing the *M. truncatula* root UMAP.

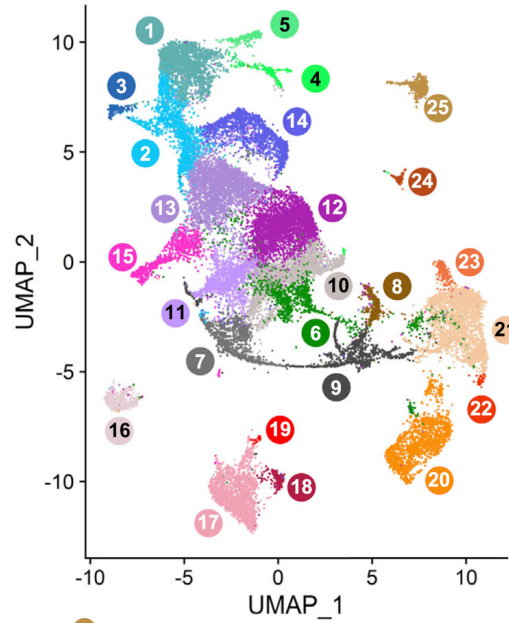
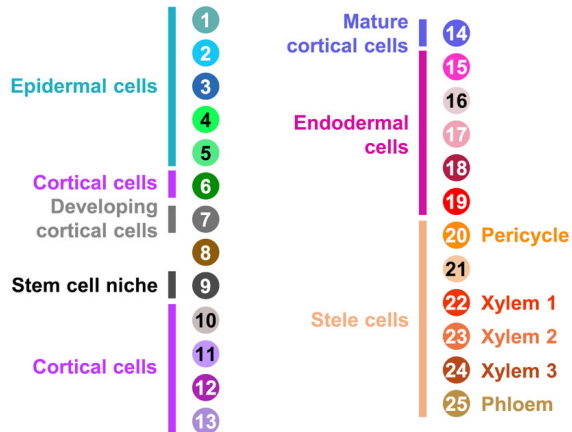


Functional annotation of *Medicago* root nuclei clusters

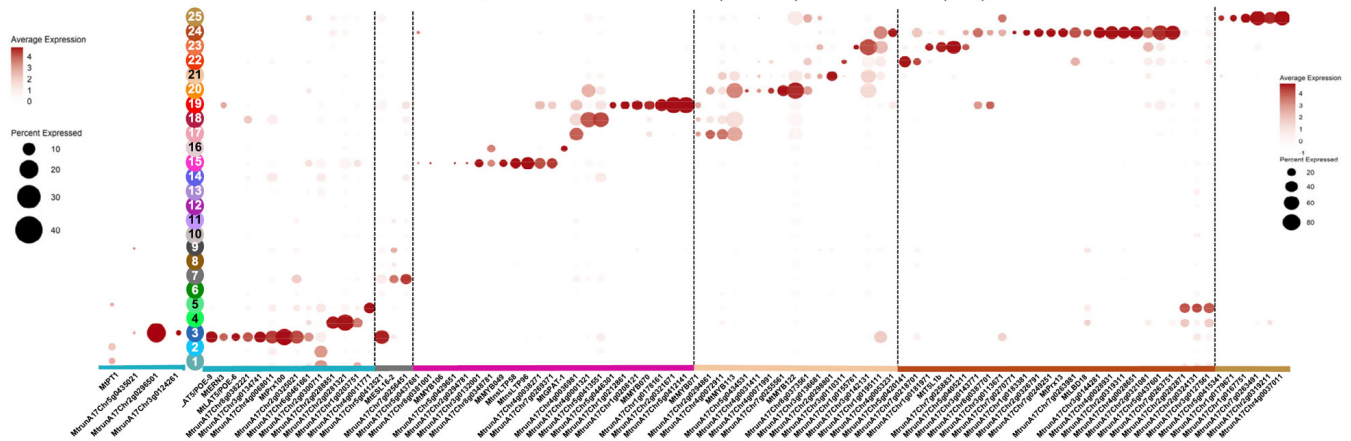
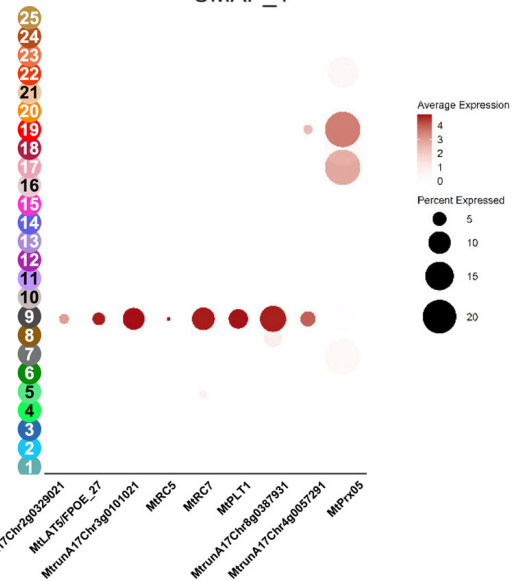
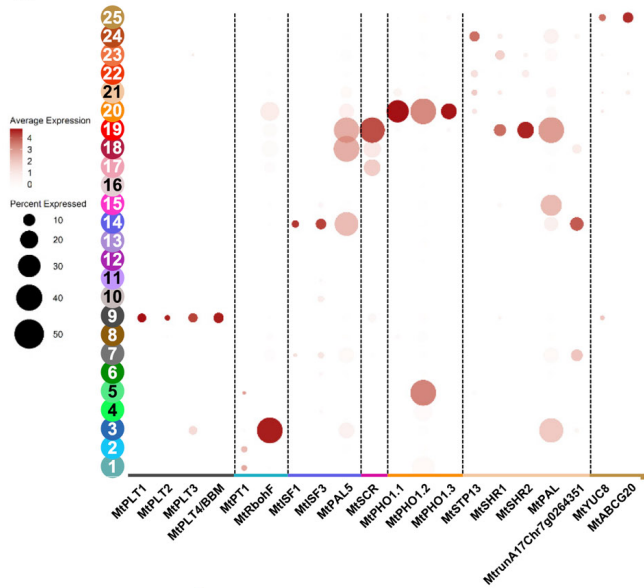
The functional annotation of root cells/nuclei according to their transcriptomic profile was previously successfully achieved in the model species *A. thaliana* by exploring the transcriptional pattern of a large number of functionally characterized cell type-specific marker genes (Denyer et al., 2019; Jean-Baptiste et al., 2019; Ryu et al., 2019; Shulze et al., 2019; Zhang et al., 2019; Farmer et al., 2021). To annotate the 25 nuclei clusters of the UMAP, we first analyzed the transcriptional pattern of the few available functionally characterized *Medicago* root cell type-specific markers (Supplemental Table 2). The *MtPLT1-4* genes are specifically expressed in the quiescent center of the root and nodule primordia (Franssen et al., 2015). Looking at their expression pattern in the *Medicago* UMAP, they are all preferentially expressed in the central star-shaped cluster #9, supporting its annotation as the stem cell niche cluster (**Figure 2A** and **Figure 2B**). To annotate *Medicago* epidermal cells, the expression of the phosphate transporter *MtPT1* gene, which is specifically expressed in *Medicago* root hairs and epidermal cells (Chiou et al., 2001), was analyzed. *MtPT1* was mostly expressed in clusters #1 and #2, and to a lesser extent, in cluster #5 (Figure 2B). The root hair-specific *MtRbohF* gene (Marino et al., 2011) was mostly expressed in cluster #3 (Figure 2B), suggesting its annotation as root hair cells. To support the annotation of the root epidermal cells, we also analyzed the expression of 45 genes previously identified as specifically expressed in the root hair (Breakspear et al., 2014) and 25 *Medicago* genes orthologous to the 168 root hair-specific *Arabidopsis* genes (Cvrčková et al., 2010). Taken together, most of these genes are preferentially or specifically expressed in the *Medicago* root epidermal cells (Supplemental Figure 2, red rectangles),

Figure 2. Functional annotation of the 25 *M. truncatula* root clusters. (A) UMAP clustering and functional annotation of the *Medicago* root cell-type clusters based on the expression of *Medicago* marker genes and of genes orthologous to *Arabidopsis* root marker genes. (B) Normalized expression levels of cell-type marker genes functionally characterized in *Medicago* (detailed in Supplemental Table 2), or orthologous to *Arabidopsis* root cell type-specific marker genes (detailed in Supplemental Table 4) across the 25 *Medicago* root clusters, shown on the y axis. The percentage of nuclei expressing the gene of interest (circle size), and the mean expression (circle color) of genes, are shown for each sub-panel.

A



B



especially in cluster #3. Based on the expression pattern of these different *Medicago* markers, clusters #1 and #2 could be thus confidently annotated as root epidermal cell clusters, and cluster #3 as a root hair cell cluster (Figure 2A). To identify the *Medicago* cortical cells on the UMAP projection, we analyzed the transcriptional activity of the cortical cell-specific genes *MtIFS1*, *MtIFS3*, and *MtPAL5* (Biala et al., 2017). *MtIFS1* was almost specifically expressed, and *MtIFS3* and *MtPAL5* were preferentially expressed in cluster #14 (Figure 2B). Besides, *MtIFS3* and *MtPAL5* are also detected in clusters #7 and #10, and in clusters #18 and #19, respectively. Taken together, these results support the annotation of cluster #14 as a cortical cell cluster. *MtSCR* is mostly expressed in the endodermis as well as in cortical and epidermal cells (Dong et al., 2021) and was thus used for annotating the endodermis (Figure 2A). *MtSCR* was mostly detected in clusters #17, #18, and #19 that are colocalized on the UMAP projection (Figure 2B), which were thus annotated as endodermal cell clusters. *MtPHO1.1*, *MtPHO1.3*, and to a lesser extent, *MtPHO1.2*, are preferentially expressed in the root stele and more specifically in pericycle cells (Nguyen et al., 2020). *MtPHO1.1* and *MtPHO1.3* were most expressed in cluster #20, where *MtPHO1.2* is also highly expressed, as well as in cluster #5. The transcriptional activity of these three *PHO1* genes supports that cluster #20 is associated with the root pericycle and/or stele (Figure 2A). Other genes expressed in the root stele, such as *MtHext1/STP13*, *MtSHR1*, *MtSHR2*, and *MtPAL* (Gaude et al., 2012; Biala et al., 2017; Dong et al., 2021), were co-expressed in clusters #19–#24 (Figure 2B). Considering that *MtSCR*, an endodermal cell marker gene, is highly expressed in cluster #19, this suggests that clusters #20–#24 correspond with root stele cell types (Figure 2A). Besides, *MtYUC8* and *MtABCG20* genes that are active in the vascular bundle of the *Medicago* root (Pawela et al., 2019; Schiessl et al., 2019) were mostly expressed in cluster #25 (Figure 2A). Taken together, this supports the annotation of cluster #20 as the *Medicago* pericycle cell cluster, and of clusters #21–#25 as *Medicago* root vasculature cell-type clusters. Finally, the expression pattern of *MtSUNN*, a receptor-like kinase acting in the autoregulation of nodulation (AON) pathway specifically expressed in the phloem (Schnabel et al., 2012), precisely maps to cluster #25, thus refining its annotation as the phloem cells cluster, and the *MtRDN1* AON-related enzyme modifying CLE signaling peptides

(Kassaw et al., 2017) to cluster #24, thus refining its annotation as a xylem cells cluster (Figure 2A).

Use of Medicago orthologs of Arabidopsis root cell-type markers for functional annotation of clusters

To further support the functional annotation of these Medicago root clusters, we additionally analyzed the transcriptional activity of Medicago genes orthologous to 1,086 Arabidopsis root cell-type marker genes (Supplemental Table 3), assuming the conservation of their cell-type-specific/-enriched transcriptional patterns. Among these genes, 101 were previously validated markers (Böhme et al., 2004; Fendrych et al., 2014; Olvera-Carrillo et al., 2015; Denyer et al., 2019; Jean-Baptiste et al., 2019; Ryu et al., 2019; Shulze et al., 2019; Turco et al., 2019; Zhang et al., 2019; see Farmer et al., 2021 for an exhaustive list), 324 were identified in at least 2 independent Arabidopsis root single-cell RNA-seq studies (Denyer et al., 2019; Jean-Baptiste et al., 2019; Ryu et al., 2019; Shulze et al., 2019; Zhang et al., 2019), and 868 genes were identified as the most specifically expressed genes within the 21 sNucRNA-seq clusters of the Arabidopsis root (Farmer et al., 2021). We found that 196 Medicago orthologs share microsyntenic relationships with the 1,086 Arabidopsis root cell type-specific marker genes thanks to the Comparative Genomic database (CoGe; <https://genomevolution.org/coge/>) (Lyons et al., 2008; Lyons and Freeling, 2008). Among these 196 genes, 38 (19.4%) and 58 (29.6%) genes were very low/not expressed, or ubiquitously expressed across all Medicago root clusters, respectively. In the end, we considered 100 Medicago genes orthologous to root cell type-specific Arabidopsis marker genes to annotate Medicago root clusters (Supplemental Table 4).

Among these 100 genes, 10, 2, and 4 are orthologous to Arabidopsis trichoblast-, atrichoblast-, and root cap-specific marker genes, respectively, including the *MtPT1* gene (Chiou et al., 2001) (Supplemental Table 2). Nine trichoblast-specific genes are mostly expressed in cluster #3, whereas *MtPT1* and the remaining six atrichoblast markers are most expressed in clusters #1, #2, #4, and #5 (Figure 2B). This conclusion is further supported by the transcriptional activity in cluster #3 of the *MtLAT52/POE_9* and *MtERN3* genes orthologous to the soybean root hair-specific *Glyma.18G025200* and *Glyma.05G157400/*

Glyma.08G115000 genes (Qiao et al., 2017), respectively (Figure 2B). Using the same approach, we found that the expression pattern of other Medicago genes orthologous to Arabidopsis cell-type marker genes similarly supported the annotation of the different Medicago root cell-type clusters. The stem cell niche annotation was confirmed for cluster #9 based on the transcriptional activity of nine Medicago genes orthologous to Arabidopsis root meristematic genes (Franssen et al., 2015) (Figure 2B, dark grey). Confirming the expression of the endodermis-specific *MtSCR* gene in clusters #17–#19 (Figure 2B, pink), we additionally identified 23 Medicago genes orthologous to Arabidopsis endodermal cell-specific genes strongly expressed in clusters #15–#19 (Figure 2B, pink). The annotation of clusters #20–#25 as stele cells was also confirmed based on the transcriptional activity of 49 orthologous genes in these clusters. Among them, xylem (i.e., clusters #22, #23, and #24) and phloem cells (i.e., cluster #25) were identified based on the expression of 23 and 6 Medicago genes orthologous to Arabidopsis xylem- and phloem-specific genes, respectively (Figure 2B, brown and light brown). As a note, three genes orthologous to Arabidopsis xylem marker genes (i.e., *MtrunA17Chr2g0282871*, *MtrunA17Chr2g0324131*, and *MtrunA17Chr3g0127561*) were also expressed in clusters #4 and #5. We assume that their activity in these two root epidermal cell clusters could reflect the induction of the cell death program, which was previously reported to be shared between xylem and root cap cells (Kumpf and Nowack, 2015; Heo et al., 2017; Farmer et al., 2021), refining the annotation of clusters #4 and #5 as containing root cap cell types. Finally, the transcriptional activity of three Medicago genes orthologous to Arabidopsis cortical cell-specific genes in cluster #7 (Figure 2B, grey) suggests that it is composed of developing cortical cells. This result is further supported by the activity of the cortical cell-specific *MtIFS3* and *MtPAL5* genes in cluster #7 (Figure 2B, purple).

Taken together, the combined use of previously characterized Medicago root cell-type marker genes with the analysis of the transcriptional activity of Medicago genes orthologous to Arabidopsis root cell-type marker genes led to the functional annotation of 19 out of the 25 Medicago root cell clusters of the UMAP (Figure 2A). To annotate the remaining six clusters (#6, #8, #10, #11, #12, and #13), all located at the center of the UMAP, we conducted a correlation analysis

between the different *Medicago* root cell clusters, hypothesizing that two clusters sharing highly correlated transcriptomic profiles would relate to the same cell type. As expected, this analysis revealed several high correlation scores, such as between the root epidermal clusters #1 and #2, and for stele/pericycle clusters #20, #21, and #23 (Supplemental Figure 3, highlighted in orange and red squares, respectively). As a note, the remaining non-annotated cluster #8 did not share a correlation with any other cluster, suggesting that cells composing this cluster have a very different transcriptomic profile compared with all other clusters. However, we were able to associate cluster #8 with an intense activity of mitochondrial and ribosomal genes (Supplemental Table 5; Supplemental Figure 4). Previous studies linked the high expression of ribosomal genes with plant developmental processes in maize, *Arabidopsis*, and tobacco plants (Ponnala et al., 2014; Makabe et al., 2017). These results thus suggest that cluster #8 is composed of cells with high biological activity. Interestingly, we found high correlation scores between clusters #6, 7, 10, 11, 12, 13, and 14 (Supplemental Figure 3, yellow squares). As the cell type marker-based analyses identified clusters #7 and #14 as cortical cells, we assume that clusters #6, #10, #11, #12, and #13 are also composed of cortical cells (Figure 2A), potentially at different stages in their differentiation process and/ or corresponding to different layers of the cortex, knowing that there are 4–5 layers of cortical cells in *M. truncatula* roots.

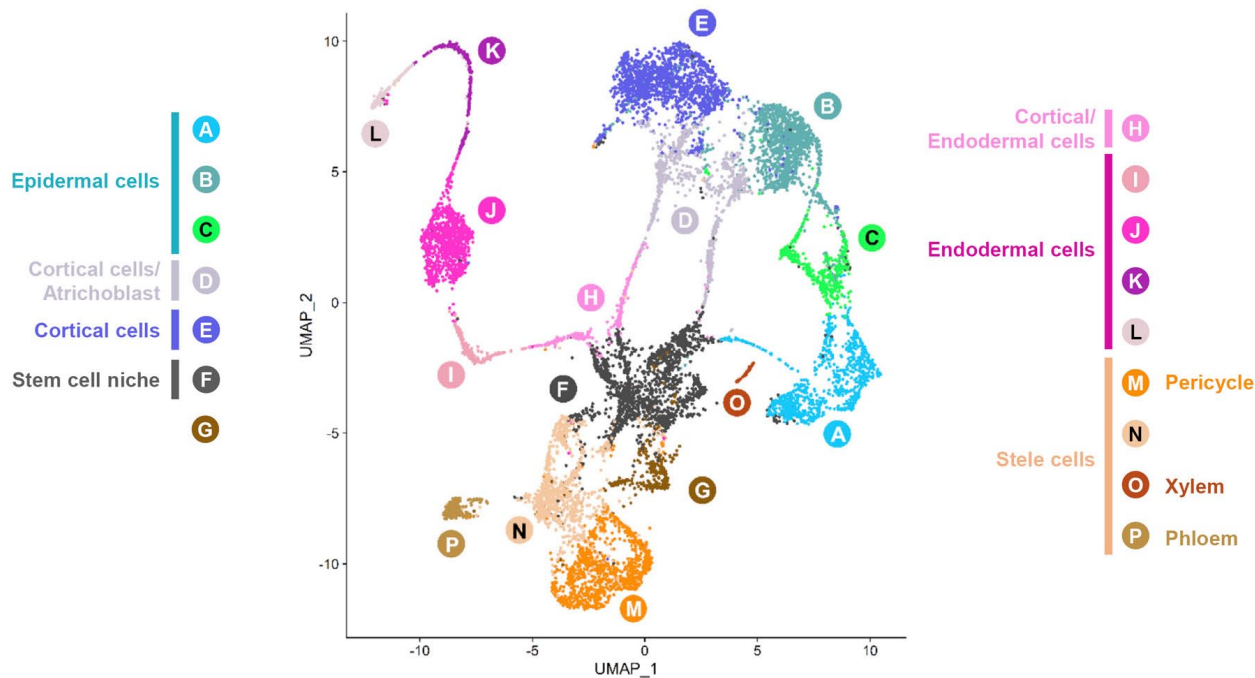
Conservation of expression patterns between orthologous Arabidopsis and Medicago genes at the single-cell level

The previous *Arabidopsis*-*Medicago* comparative genomic and transcriptomic analysis used to annotate *Medicago* root clusters (Figure 2) suggested that transcriptional patterns of orthologous genes could be largely conserved upon speciation of the 2 plants 108 million years ago (Zeng et al., 2017). To further explore the extent of this conservation at the single-cell level, a correlation analysis was conducted on 3,921 pairs of orthologous genes sharing microsyntenic relationships based on the CoGe database (Lyons et al., 2008; Lyons and Freeling, 2008) (Supplemental Table 6). To maximize the biological significance of the analysis, we processed the previously generated *Arabidopsis* sNucRNA-seq datasets (Farmer et al., 2021),

similarly to the *Medicago* sNucRNA-seq datasets (see methods). This updated analysis led to the identification and re-annotation of 16 *Arabidopsis* root clusters (Farmer et al., 2021) (**Figure 3A**). We observed a similar topology between the *Medicago* and *Arabidopsis* sNucRNA-seq UMAPs (Figures 2A and 3A), with the stem cell niche (#9) located in a star-shaped cluster at the center of the UMAP projection, whereas the most differentiated cell types (e.g., epidermal [#1, #2, and #3], phloem [#25], xylem cells [#22, #23, and #24]) were retrieved in the periphery. To evaluate the conservation of the transcriptional profiles between *Arabidopsis* and *Medicago* orthologs, the average gene transcriptional activity of one-to-one orthologs was similarly calculated for each of the 25 mock-inoculated *Medicago* (Figure 1A) and of the 16 *Arabidopsis* root clusters (Figure 3A), and a correlation analysis was then conducted (Figure 3B). Stem cell niche (#9) (Figure 3B, black square) and stele cells (#20–#25) (Figure 3B, peach-orange square) shared the highest correlation between the two species, suggesting that the transcriptional activities of orthologous genes were most conserved across these root cell-types. To a lesser extent, the transcriptomic profiles of orthologous genes were also conserved between the two species in trichoblasts/epidermal root hair cells (#1–#3) (Figure 3B, blue square). Similar conservation was recently reported for root hair, xylem, and phloem cells between the more distantly related rice and *Arabidopsis* roots (Liu et al., 2021; Zhang et al., 2021). We hypothesize that the unique functions of vascular tissues (xylem and phloem) and trichoblast cells for plant nutrition, as well as the role of the stem cell niche in root development, favored the evolutionary conservation of a core transcriptome between these plant species. In contrast, the transcriptomes of endodermal (#15–#19) and cortical cell types (#6, #7, and #10–#14) were most divergent between *Arabidopsis* and *Medicago* (Figure 3B, purple and violet squares), suggesting either a difference

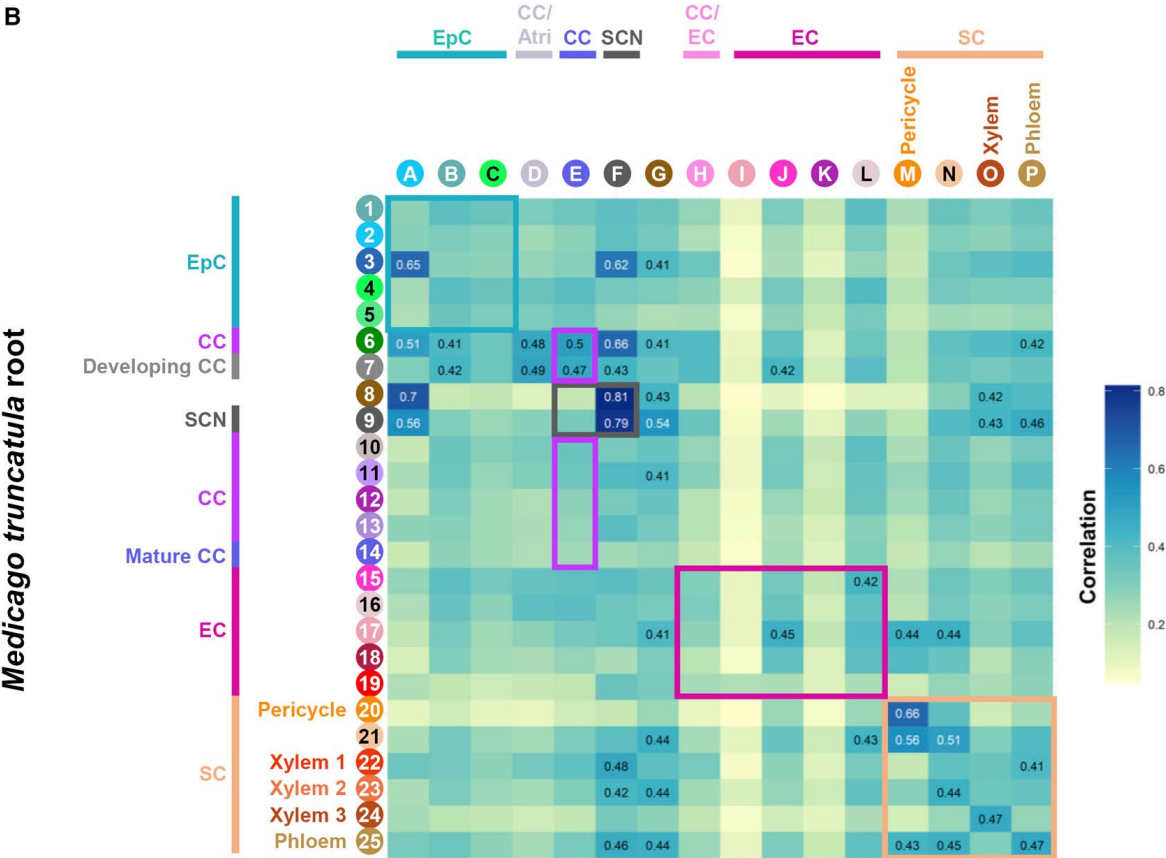
Figure 3. Comparative transcriptomic analysis of the *M. truncatula* and *A. thaliana* root cell clusters. (A) Functional annotation of *Arabidopsis* root nuclei clusters based on the expression profile of cell-type marker genes defined from (Farmer et al., 2021). (B) Pairwise correlations of *Arabidopsis* (x axis) and *Medicago* (y axis) root cell clusters. Only correlation numbers greater than 0.4 (black numbers) or 0.5 (white numbers) are shown in the heatmap. CC, cortical cells; EC, endodermal cells; EpC, epidermal cells; SC, stele cells; SCN, stem cell niche.

A



Arabidopsis thaliana root

B



in nutrient provision when growing the *Arabidopsis* and *Medicago* plants, a lower pressure to maintain the transcriptomic signature of these cell-types between the two species, and/or diverging biological functions of these cell types between Fabaceae (legumes) and Brassicaceae, such as their differential capacity to interact with soil beneficial microbes through endosymbiosis.

A differential transcriptional regulation in response to E. meliloti of nodulation and hormonal genes depending on cell types

To date, transcriptomic responses of legume roots to rhizobial inoculation were investigated mainly at the whole root level (Mergaert et al., 2019) or on isolated populations of root hair and epidermal cells (Libault et al., 2009; Breakspear et al., 2014). These bulk analyses, however, lacked resolution, considering that only a subset of plant cells respond to and are infected by rhizobia and that the transcriptional response of cell types located deeper within roots cannot be easily assessed. Here, we had a unique opportunity to explore the differential regulation of gene expression across the different cell types in response to *E. meliloti*.

To estimate the percentage of individual *Medicago* root epidermal cells responding to *E. meliloti*, we quantified the number of epidermal root cells (i.e., clusters #1, #2, and #3, excluding the epidermal/root cap clusters #4 and #5) expressing typical early rhizobial infection marker genes, namely *MtRPG*, *MtFLOT4*, and *MtVPY* (Roy et al., 2019). In *E. meliloti*-inoculated roots, 19.5% (433/2225 epidermal nuclei) of epidermal nuclei expressed at least one of these symbiotic marker genes, whereas only 2.3% (39/1694 epidermal nuclei) were detected in mock-inoculated roots. The increase in the number of rhizobia-responsive epidermal cells upon *E. meliloti* inoculation identified is, thus, significantly larger than the previously estimated size of the epidermal cells population infected by rhizobia, which was approximately 1%–5% of root hair cells (Nutman, 1959). These results suggest that only a subset of the transcriptionally responsive root epidermal cells from these three clusters is effectively infected by rhizobia.

To identify the set of differentially expressed genes (DEGs) in response to rhizobium inoculation from each *Medicago* root cell cluster,

we used DEsingle, a bioinformatics package designed to identify DEGs from single-cell RNA-seq datasets (Miao et al., 2018). Using a p value threshold of less than 0.05 and a | fold change (FC)| of more than 1.5, we identified a total of 8513 DEGs (Supplemental Table 7). Focusing on the most transcriptionally responsive clusters (i.e., >500 DEGs) (**Figure 4A**; Supplemental Table 7, bold characters), the root hair cells cluster #2 was retrieved, as well as the cortical clusters #7 and #11, the endodermal clusters #15, #16, and #18, and the pericycle cluster #20. This result nicely fits with knowledge previously gained using microscopy during early nodule ontogeny (Xiao et al., 2014), which showed that cellular symbiotic processes mainly affected epidermal and cortical cells, and to a lesser extent, endodermis and pericycle cells. We thus further focused our analysis on these clusters, as well as on genes previously known to be acting in nodulation and/or previously annotated as related to hormonal pathways regulating early nodulation stages (Roy et al., 2019).

In the trichoblast cluster #2, several known infection-related genes were retrieved as upregulated, namely *MtCBS1*, *MtRbohG*, and *MtRbohH* (Montiel et al., 2018), *MtRPG*, *MtVPY*, and *MtAnn1* (de Carvalho Niebel et al., 1998), and *MtNMN1* (i.e., an ortholog of *GmNMN1*) (Libault et al., 2011). Other genes that belong to the NF signaling pathway were also upregulated in root hair and epidermal cells in response to *E. meliloti*, namely *MtNFH1* and *MtLYK10* (Larrainzar et al., 2015), *MtPUB1* (noting that *MtPUB2* is repressed in this same cluster), *MtIPD3*, *MtDMI1*, *MtDMI2*, *MtDMI3*, *MtNSP1*, *MtNSP2*, *MtERN1*, and *MtERN2* (Cerri et al., 2016), and *MtNIN* (Supplemental Tables 8 and 9, **Figure 5**). Unexpectedly, *MtKNOX3* and *MtKNOX5* genes, previously proposed to control nodule development (Di Giacomo et al., 2017; Dolgikh et al., 2020), and *Does Not fix Nitrogen 2* (*MtDNF2*) (Bourcy et al., 2013)] and *Nodule Cysteine Rich peptide 112* (*MtNCR112*) (Alunni et al., 2007) genes, regulating later stages of the nodulation process, were also upregulated in cluster #2 48 h after rhizobium inoculation. In addition, we noticed that the *MtNFYA2* and *MtNF-YA6* genes, phylogenetically related to the early nodulation *MtNF-YA1* gene (Baudin et al., 2015), were also upregulated in cluster #2. Genes belonging to hormonal regulatory pathways were also induced in this cluster in response to rhizobial inoculation. These include notably cytokinin (e.g., *MtIPT1*, *CYP735A1*-like, *MtHPT1*, *MtRRB6*, *MtRRA2*, and *MtRRA5*)

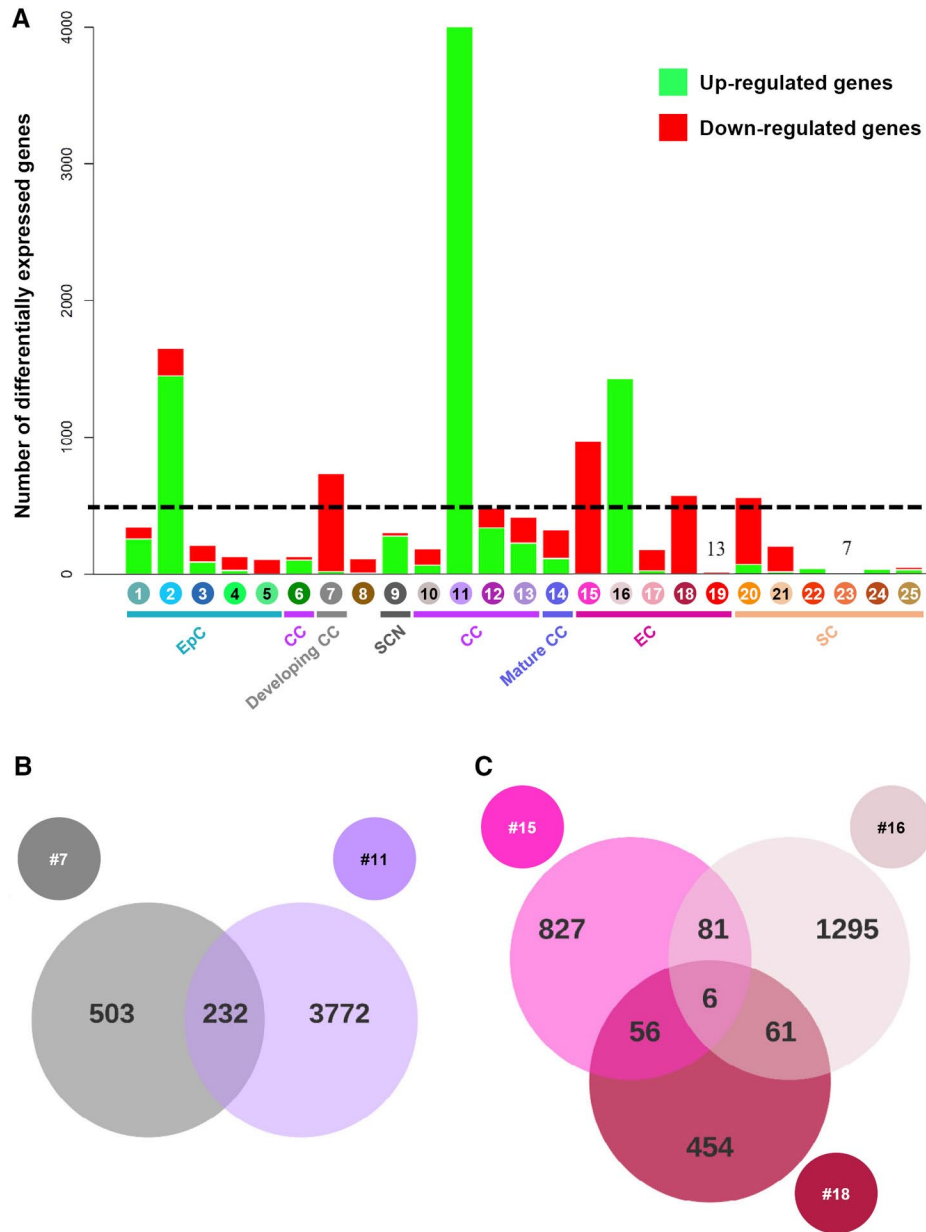


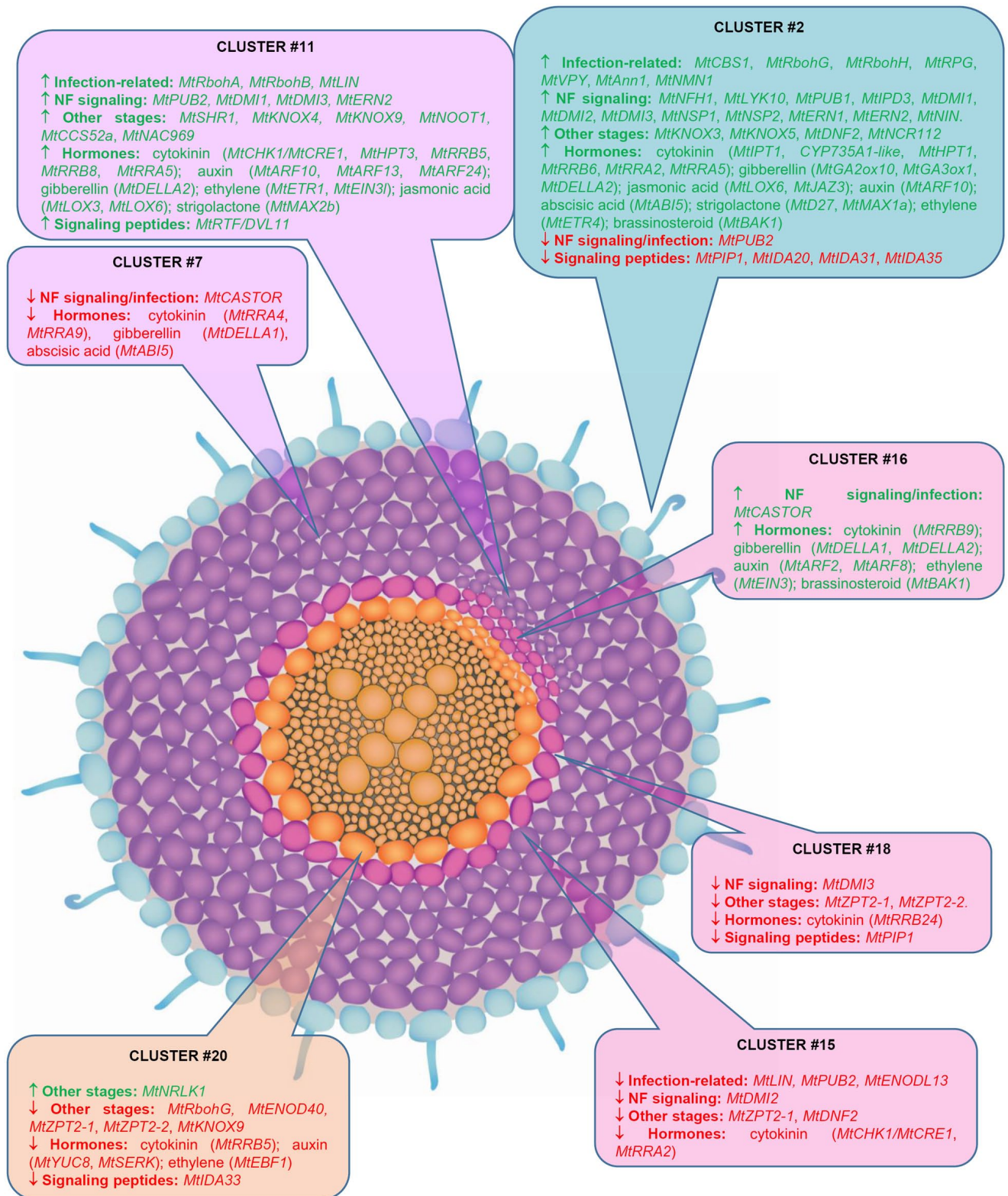
Figure 4. Differential expression of the *M. truncatula* genes in response to *E. meliloti* inoculation across the 25 root cell clusters. (A) The transcriptional response of Medicago root cells to *E. meliloti* inoculation differs between celltype clusters. The number of up- and downregulated genes are highlighted in green and red bars, respectively. The dashed bar reflects the 500 DEGs thresholds. (B and C) Comparison of the number of DEGs between the cortical cell clusters #7 and #11 (B) and between the endodermal cell clusters #15, #16, and #18 (C). CC, cortical cells; EC, endodermal cells; EpC, epidermal cells; SC, stele cells; SCN, stem cell niche.

(Azarakhsh et al., 2018; Tan et al., 2019) and gibberellin (*MtCPS1*, *MtGA2ox10*, *MtGA3ox1*, and *MtDELLA2*) (Fonouni-Farde et al., 2016; Kim et al., 2019) biosynthesis and signaling genes. In addition, the expression of genes related to the jasmonic acid (*MtLOX6* [Gao et al., 2007] and *MtJAZ3* [Ge et al., 2016]), auxin (*MtARF10*) (Shen et al., 2015), abscisic acid (*MtABI5*) (Verdier et al., 2013), strigolactone (*MtD27*, *MtMAX1a*) (Liu et al., 2011; Müller et al., 2019), ethylene (*MtETR4*) (Tan et al., 2019), and brassinosteroid (*MtBAK1*) (Tavormina et al., 2015) pathways were also upregulated upon rhizobium inoculation in cluster #2. Conversely, the expression of several genes encoding signaling peptides was repressed by rhizobium in cluster #2, such as *MtPIP1*, *MtIDA20*, *MtIDA31*, and *MtIDA35* (Inflorescence Deficient in Abscission) (de Bang et al., 2017) (Supplemental Tables 8 and 9, Figure 5). To further estimate the relevance of the trichoblast DEGs identified, we conducted a comparative analysis with the list of 267 DEGs previously reported in the Medicago root hair cells in response to rhizobium inoculation (Breakspear et al., 2014). We found that 99 of these genes (37%) were differentially expressed in the sNucRNAseq dataset and, among them, 68 genes (69%) were significantly differentially expressed in at least one of the epidermal clusters #1, #2, or #3 (Supplemental Figure 5). Considering the difference in sensitivity between the technologies used, these results overall support the identification of genes differentially expressed in Medicago trichoblasts in response to rhizobia inoculation.

The cortical cell cluster #7 DEGs are all repressed upon rhizobium inoculation, including one gene, *MtCASTOR*, homologous to an *L. japonicus* nodulation gene (Charpentier et al., 2008), several cytokinin-related genes (two *LOG-like* genes and two *RRA* signaling genes, *MtRRA4* and *MtRRA9*), the gibberellin signaling gene *MtDELLA1*, and the abscisic acid signaling gene *MtABI5* (Supplemental Tables 8 and 9, Figure 5). Conversely, the cortical cell cluster #11 DEGs are all upregulated and include several genes controlling the rhizobia-infection process (*MtRbohA* [Marino et al., 2011], *MtRbohB*, and *MtLIN*), NF signaling (*MtPUB2*, *MtDMI1*, *MtDMI3*, and *MtERN2*), and nodule development (*MtSHR1*, *MtKNOX4*, and *MtKNOX9* [Di Giacomo et al., 2017], *MtNOOT1* and *MtCCS52a* [Cebolla et al., 1999]) and function (*MtNAC969*) (de Zélicourt et al., 2012)] (Supplemental Tables 8 and 9, Figure 5). In addition, hormonal genes related to cytokinin (*MtCHK1/*

Figure 5. Summary of the *M. truncatula* root cell type-specific transcriptional response to rhizobial inoculation. Selected genes previously known as related to nodulation and hormonal pathways and identified as differentially expressed in the clusters showing more than 500 DEGs are listed. Besides the expected induced expression pattern of numerous nodulation-related genes such as in the root hair cells cluster #2, more unexpected expression profiles were also highlighted notably for some late nodulation genes, and for repressed early nodulation and hormone-related genes in cortical and endodermal clusters #7 and #15–#18. Genes are listed in the following categories: rhizobial infection-related, NF signaling, other nodulation stages, hormones, and signaling peptides. Upward arrows indicate gene inductions by rhizobia, and downward arrows, repressions.

MtCRE1 [Gonzalez-Rizzo et al., 2006], *MtHPT3*, *MtRRB5*, *MtRRB8*, and *MtRRA5*), auxin (*MtARF10*, *MtARF13*, and *MtARF24*), gibberellin (*MtDELLA2*), ethylene (*MtETR1*, *MtEIN3-like*), jasmonic acid (*MtLOX3* and *MtLOX6*), and strigolactone (*MtMAX2b*), were also upregulated upon rhizobium inoculation in cluster #11. Finally, two nitrate signaling-related TFs were upregulated (*MtNLP1* and *MtNLP4*) (Luo et al., 2021) as well as a specific signaling peptide (*MtRTF/DVL11*), in agreement with the symbiotic function previously reported for *MtDVL1* (Combie et al., 2008) (Supplemental Tables 8 and 9, Figure 5). Of note, despite the opposite transcriptional responses between clusters #7 and #11 to rhizobial inoculation, 232 genes out of the 736 DEGs in cluster #7 (31.5%) were shared as significantly differentially expressed with cluster #11 (Figure 4B). This indicates that an unexpectedly high number of genes show opposite transcriptional regulation in response to rhizobium within different cortical cell clusters. This exemplifies that reaching a cell type-specific level allows identifying that strong up- or downregulations can occur simultaneously in different clusters at the same 48 h after rhizobium inoculation time-point, which would be thus likely missed if whole roots would be used. In addition, the differential transcriptional responses observed between cortical cell clusters for several hormonal pathways and root/nodule developmental genes upon rhizobial inoculation might mark cells that are activated for nodule organogenesis from those that are not, and may relate to the different layers of cortex (i.e., inner vs. outer), or correspond with cortical cells opposite to proto-phloem versus proto-xylem poles where nodule organogenesis is differentially initiated (Heidstra et al., 1997).



The endodermal clusters #15, #16, and #18, showing more than 500 DEGs, once more comprise either only downregulated genes (#15 and #18), or upregulated genes for cluster #16 (Figure 4A; Supplemental Table 7). In contrast with the cortical cell clusters, these endodermal clusters shared a limited number of DEGs (Figure 4C). Surprisingly, among clusters #15 and #18 downregulated genes, many early nodulation genes were observed, including for cluster #15 the infection-related genes *MtLIN*, *MtPUB2*, *MtENODL13*, the NF signaling-related gene *MtDMI2*, and the cytokinin signaling genes *MtCHK1/MtCRE1* and *MtRRA2*, as well as the late nodulation genes *MtZPT2-1* (Frugier et al., 2000) and *MtDNF2*; and for cluster #18, the NF signaling gene *MtDMI3*, the cytokinin signaling gene *MtRRB24*, the late nodulation genes *MtZPT2-1* and *MtZPT2-2*, and a signaling peptide, *MtPIP1* (Supplemental Tables 8 and 9, Figure 5). Concerning endodermis cluster #16 upregulated genes, the symbiotic-related gene *MtCASTOR* was retrieved, as well as several hormone-related genes including *MtARF2* and *MtARF8* (auxin), *MtDELLA1* and *MtDELLA2* (gibberellin), *MtRRB9* (cytokinin), *MtEIN3* (ethylene), and the co-receptor *MtBAK1*. In addition, the expression of the nitrate signaling-related *MtNLP1* gene was induced (Supplemental Tables 8 and 9, Figure 5).

Finally, in the pericycle (cluster #20), whereas the rhizobium-induced genes do not include any previously studied early nodulation genes, the *MtNRLK1* receptor-like kinase gene (Laffont et al., 2018) was retrieved as slightly induced in response to rhizobia. The *MtRbohG*, *MtENOD40*, *MtZPT2-1*, and *MtZPT2-2* nodulation genes were unexpectedly repressed by rhizobium, as well as the *MtKNOX9* developmental gene. Regarding hormone-related genes, the cytokinin *MtRRB5*, the auxin *MtYUC8*, and the ethylene *MtEBF1* gene, as well as the *MtIDA33* signaling peptide and the *MtSERK* coreceptor, were also repressed (Supplemental Tables 8 and 9, Figure 5).

Taken together, our unique dataset allowed a refined expression pattern analysis, which revealed unexpected celltype specificity/enrichment for some of the already well-known early nodulation genes (e.g., a differential expression across cortical cell clusters and repression in endodermal cell clusters) or for the *MtCASTOR* gene which currently has no symbiotic function reported in *M. truncatula* unlike its homolog from *L. japonicus* (Venkateshwaran et al., 2012). In addition, some nodulation genes that were previously linked to later

symbiotic stages (e.g., *MtNCR112*, *MtZPT2-1*, *MtZPT2-2*, *MtDNF2*, and *MtNAC969*) also showed a differential expression in response to rhizobium in specific root cell-type clusters. Surprisingly, a few anticipated early nodulation genes were missing from the DEG dataset (e.g., *MtLIN*, *MtNF-YA1*, and *MtNPL*), but a manual inspection of their expression profiles revealed differential regulations by rhizobia that were below the statistical threshold used. This indicates that our statistical analysis is conservative, allowing providing a robust dataset of DEGs, but also likely missing other genes of interest. More refined statistical analyses could be, however, performed in the future on this dataset, focusing only on a subset of specific clusters to extract such additional information that is currently lost because of the high variance between all nuclei analyzed.

Cell type-specific expression of nodulation and cytokinin-signaling related genes

As many nodulation-related genes can be already expressed in cells before bacterial inoculation, we additionally conducted a comprehensive analysis of their expression patterns independent of their response to rhizobial inoculation, focusing notably on genes showing cell type-enriched or -specific patterns that were not previously identified as DEGs (see above). Epidermal cells (#1–#5) most specifically expressed the *MtROP5* (Riely et al., 2011), *MtLIN*, and *MtPT5* (Wang et al., 2022) genes associated with rhizobial infections (clusters #2 and #3) (Damiani et al., 2016); *MtNFP*, *MtLYK3* (Smit et al., 2007), *MtLYK6*, *MtCNGC15c* (Charpentier et al., 2016), and *MtNF-YA1* genes, related to NF signaling (clusters #2 and #3); *MtCHIT5a* that is linked to NF degradation (clusters #4 and #5) (Tian et al., 2013), as well as genes related to late nodulation stages, namely *MtRab7A1* (Limpens et al., 2009), *MtSYP132* (Pan et al., 2016), *MtVPE* and *MtSPK1* (Andrio et al., 2013), *MtZIP6* (Abreu et al., 2017) (clusters #2 and #3), *MtDNF2* (cluster #4), *MtDGD1* (Si et al., 2019) (clusters #4 and #5), and *MtNAC969* (cluster #5); and *MtNLP1*, a gene linked to nitrate signaling (clusters #2 and #3) (**Figure 6A**).

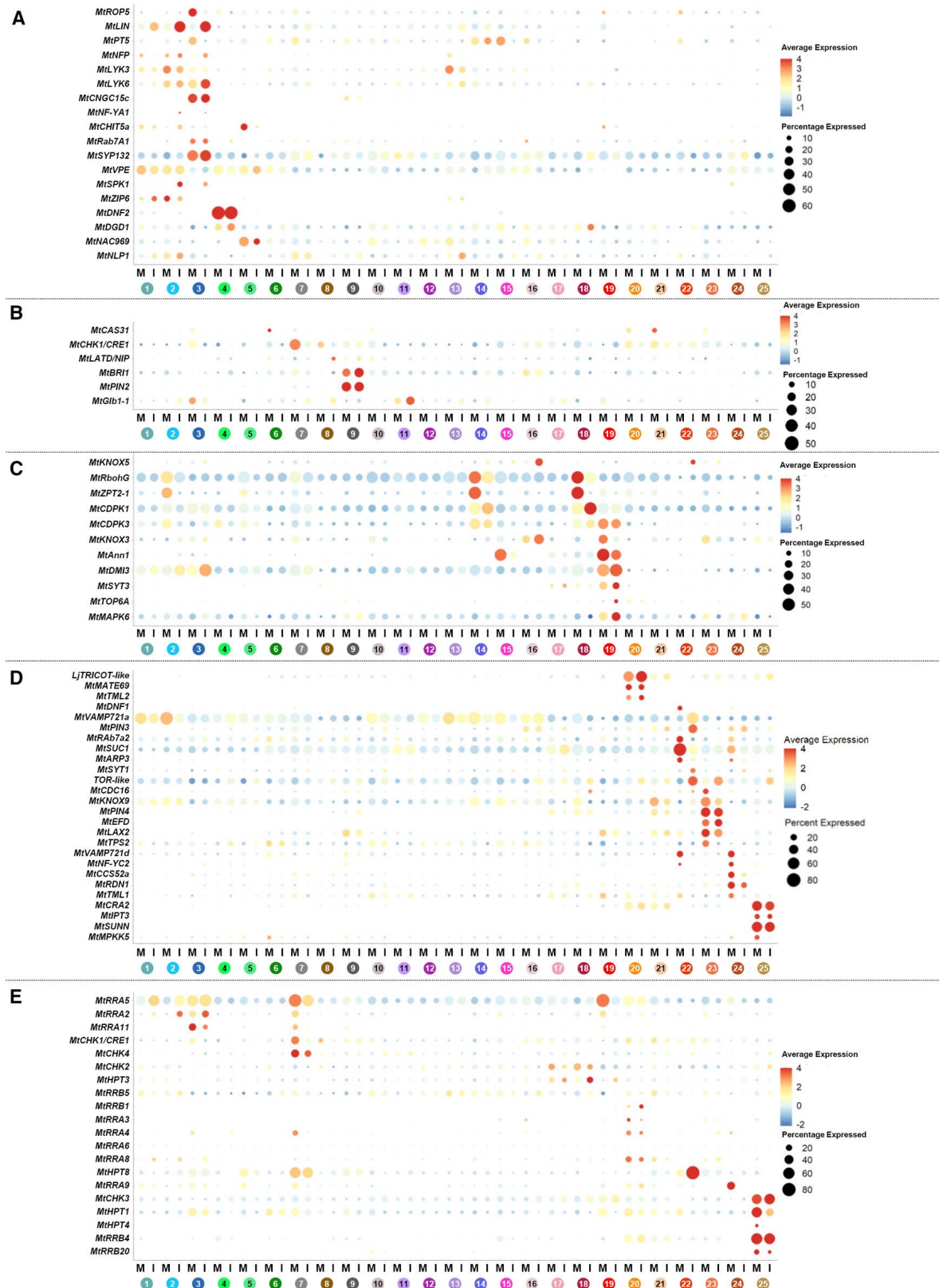
In contrast, only a small number of previously characterized nodulation-related genes were specifically expressed/enriched in the non-annotated cluster #8, in the stem cell niche cluster #9, and in the

cortical clusters #6, #7, and #10–#14 (i.e., *MtCHK1/CRE1*, a cytokinin receptor required for nodule organogenesis in cluster #7; *MtLATD/NIP*, an abscisic acid transporter linked to early nodulation in cluster #8 [Bagchi et al., 2012]; *MtBRI1*, a brassinosteroid receptor linked to nodulation [Cheng et al., 2017,] and *MtPIN2*, an auxin efflux carrier linked to nodule organogenesis in cluster #9 [Huo et al., 2006]; and *MtGlb1-1*, a gene involved in later nodulation stages in cluster #11 [Berger et al., 2020]) (Figure 6B).

In the endodermis (#15–#19), we identified several nodulation-related genes specifically expressed in one or several clusters (i.e., *MtKNOX5* [cluster #16], *MtRbohG*, *MtZPT2-1*, and *MtCDPK1* [cluster #18, as well as in the cortical cell cluster #14] [Ivashuta et al., 2005]), *MtCDPK3* (clusters #18 and #19), *MtKNOX3* (clusters #16 and #19), *MtANN1* (clusters #15 and #19), *MtDMI3* (cluster #19 in addition to the epidermal cell clusters #1, #2, and #3), *MtSYT3* (Gavrin et al., 2017), *MtTOP6A* (an ortholog to *LjSUNERGOS1* [Yoon et al., 2014]), and *MtMAPK6* (cluster #19 [Chen et al., 2017]) (Figure 6C).

In the stele (#20–#25), a *LjTRICOT-like* gene (Suzaki et al., 2013), *MtMATE69* (Wang et al., 2017), and *MtTML2* (Gautrat et al., 2019) were specifically expressed in cluster #20 (pericycle cells), while *MtDNF1* (Van de Velde et al., 2010; Wang et al., 2010), *MtVAMP721a* (Sinha-roy et al., 2013), *MtPIN3* (Huo et al., 2006), *MtRAB7a2* (Limpens et al., 2009), *MtSUC1* (Hohnjec et al., 2003), *MtARP3* (Gavrin et al., 2015), *MtSYT1* (Gavrin et al., 2017), TOR-like, *MtCDC16* (Kuppusamy et al., 2009), *MtKNOX9* and *MtPIN4* (Huo et al., 2006), *MtEFD* (Verniéet al., 2008), *MtLAX2* (Roy et al., 2017), *MtTPS2* and *MtVAMP721d* (Ivanov et al., 2012), *MtNF-YC2* (Baudin et al., 2015), *MtCCS52a*, and *MtRDN1* (Sch-nabel et al., 2011) were enriched in at least one of the xylem clusters (i.e., #22–#24). Finally, in cluster #25 (phloem), we identified several genes associated with the systemic regulation of nodulation (Gautrat

Figure 6. Cell-type enrichment of known *M. truncatula* nodulation and cytokinin signaling genes. (A–E) Normalized expression levels of *Medicago* nodulation-related genes specifically expressed/enriched in the epidermal (A), cortical (B), endodermal (C), and steles cells (D), as well as of *Medicago* cytokinin signaling-related genes (E). The 25 *Medicago* root clusters identified are shown on the x axis. The percentage of nuclei expressing the gene of interest (circle size), and the mean expression (circle color) of genes, are shown for each sub-panel. I, rhizobia-inoculated condition; M, mock-inoculated condition.



et al., 2021), namely, *MtCRA2* (Huault et al., 2014; Mohd-Radzman et al., 2016), *MtSUNN*, *MtTML1* (Gautrat et al., 2019), *MtIPT3*, and *MtMPKK5* (Figure 6D). These observations are well supported by previous studies reporting the cell-type specificity of these genes.

Considering the cytokinin signaling pathway which is key for early nodulation (Gamas et al., 2017), we found *MtRRA5* most expressed in epidermal clusters #1–#3, as well as in clusters #7 (cortex) and #19 (endodermis), *MtRRA2* in clusters #2 and #3 (epidermis/root hairs), *MtRRA11* in clusters #3 (epidermis) and #7 (cortex), *MtCHK1/CRE1*, *MtCHK4*, *MtHPT8*, and *MtRRA4* in cluster #7 (cortex), *MtCHK2* and *MtHPT3* in clusters #17 and #18 (endodermis), *MtRRB5* in cluster #18 (endodermis), *MtRRB1*, *MtRRA3*, *MtRRA4*, *MtRRA6* and *MtRRA8* in cluster #20 (pericycle), *MtHPT8* and *MtRRA9* in clusters #22, #23, and #24 (xylem), and *MtCHK3*, *MtHPT1*, *MtHPT4*, *MtRRB4*, and *MtRRB20* in cluster #25 (phloem) (Figure 6E).

These results overall support that many early nodulation genes, including those involved in cytokinin signaling, have a cell-type specific/enriched expression pattern, indicating a coordinated activity between cell types to successfully promote nodule initiation. Interestingly, the co-regulation in a specific cluster of different genes belonging to a large family, or even to the same functional pathway, combined with phylogenetic analyses, now provides critical information to develop more efficient functional analyses to overcome functional redundancy. As an example, the *RRB* family related to cytokinin signaling contains 12 genes for which expression was detected in at least one cluster of the UMAP. Noteworthy, *MtRRB4* and *MtRRB20* have overlapping expression patterns in the phloem (cluster #25), suggesting a likely functional redundancy, also knowing their close phylogenetic relationship in the same clade (Tan et al., 2019). When combining this information with the expression pattern of genes from other cytokinin signaling families, a specific cytokinin signaling pathway preferentially acting in the phloem cells can now be identified, involving the *MtCHK3* receptor, the *MtHPT1* and *MtHPT4* phosphotransfer proteins, and the *MtRRB4* and *MtRRB20* TFs. Similarly, when considering systemic pathways regulating nodulation (Gautrat et al., 2021), our analysis strikingly demonstrates that most known genes (i.e., *MtCRA2*, *MtSUNN*, *MtIPT3*, and *MtTML1*) are specifically expressed, and even induced by rhizobium in phloem cells (cluster #25). Importantly, getting access

to such very detailed and clear-cut spatial expression information allows generating innovative working hypotheses to be further tested functionally.

Root cell type-specific versus shared functional pathways enriched in response to a short-term rhizobium inoculation

To reveal new biological functions potentially controlling the response of Medicago root cell types to *E. meliloti*, we performed a gene ontology analysis using the Mapman software (Schwacke et al., 2019; Tellstrom et al., 2007) on the DEGs from clusters #2, #7, #11, #15, #16, #18, and #20 containing more than 500 DEGs (Figure 4A). The full list of enriched functional pathways is shown in Supplemental Table 10 and Supplemental Figure 6.

In cluster #2, functional pathways corresponding with changes in cellular organization, including modification of the cell wall (pectin esterases) and vesicle transport, were enriched for genes upregulated in response to bacterial inoculation (Wilcoxon rank-sum test, $p < 0.05$). This likely reflects the curling of root hairs associated with rhizobial infections. Among metabolic pathways, the flavonoid metabolism was enriched, notably the biosynthesis of dihydroflavonols, as well as the cytokinin and gibberellin biosynthesis pathways. This result strikingly fits with knowledge gained during the last decades where these three plant signaling pathways were shown as crucial for the regulation of early stages of rhizobial infections in the Medicago root epidermis (Gonzalez- Rizzo et al., 2006; Plet et al., 2011; Fonouni-Farde et al., 2016, 2017; Roy et al., 2019). In addition, both lysine motif and leucine-rich repeats receptor kinases were enriched in this trichoblast cluster #2 in response to rhizobium inoculation, which includes already known receptors required for rhizobial recognition and infection, as well as the phosphoinositide metabolism and PHD finger or GRAS TFs, the latter family including also already known early nodulation genes. Concerning genes downregulated by rhizobia, the brassinosteroid and ethylene pathways were enriched, the latter hormone being previously extensively characterized as an inhibitor of rhizobial infections, notably in Medicago (Penmetsa et al., 2008).

Cortical cell clusters #7 downregulated DEGs showed enrichment for cell wall modifications and lipid metabolism, several hormonal

pathways namely abscisic acid metabolism, ethylene signaling, as well as AP2/EREBP and Trihelix TF families; and cluster #11 upregulated DEGs for biotic stress responses, lipid, terpenoid, phenylpropanoid, and glucosinolate metabolic functions, as well as C₂H₂ zinc finger, CCAAT box binding factors, G2-like (GARP), and MYB-related TF families, the protein targeting secretory and protein degradation pathways (subtilases and autophagy), and cytoskeleton reorganization.

Endodermal cells clusters #15 and #18 (downregulated genes) were enriched for cell wall degradation and lipid metabolism functions, as well as ethylene and jasmonate metabolism, GRAS TF families, leucine-rich repeat receptor kinases, and protein degradation via ubiquitination. Regarding cluster #16 (upregulated genes), the biotic stress response and glucosinolate degradation functions were enriched, as well as the jasmonate hormone and the ARR TF family related to cytokinins, cell division and cell cycle, and sugar transport.

Finally, in pericycle cells (cluster #20), rhizobium downregulated genes were once more enriched for ethylene signaling pathway genes and AP2/EREBP TFs, as previously observed for the root hair cluster #2, the cortex cluster #7, and the endodermal clusters #15 and #18, highlighting one of the few shared biological responses across different cell types.

Overall, as previously noticed for the analysis of known nodulation-related genes, the novel cell-type specific information gained allows for generating new hypotheses, such as the existence of a tight interaction in inner root tissues of symbiotic responses with defense pathways through the modulation of specific specialized metabolite production. It also highlights specific hormonal pathways and TF families for which functional studies remain still limited, or even lacking, and that could thus be targeted in the future in relation to cell-type specific phenotypes (e.g., rhizobial infections or nodule organogenesis).

Plant root development requires tightly coordinated regulation of transcriptomic programs. We and others revealed root transcriptomic profiles at a single-cell level notably in the model plant *A. thaliana* (Denyer et al., 2019; Jean-Baptiste et al., 2019; Ryu et al., 2019; Shulze et al., 2019; Zhang et al., 2019; Farmer et al., 2021). In this study, we provide a comprehensive annotation of the *Medicago* root cell types according to their transcriptomic profiles, as well as an analysis of the transcriptomic response of *Medicago* root cells to rhizobial infection.

Our study largely confirmed knowledge gained during the last decades in legume nodulation (i.e., the regulation of the expression of nodulation-related and hormonal genes known to regulate rhizobial infection and/or nodule organogenesis), but also nicely illustrates the gain of knowledge obtained using such single nuclei transcriptomic approaches to better understand the cell type specifically restricted responses of plants to microbial infection. The robust and high-quality dataset generated is also a resource to enable the discovery of new genes of interest not previously highlighted by bulk transcriptomic analyses. In particular, accessing cell-type information allows for generating more precise hypotheses regarding the symbiotic processes potentially affected by these novel candidate DEGs, and thus facilitates planning more appropriately experimental designs, notably by using tissue-specific promoters for which single-cell datasets are a key resource, and for performing refined focused phenotyping of rhizobial infections versus nodule organogenesis. The application of single-cell -omics technologies to other symbiotic and pathogenic plant-microbe interactions would help better understand the intimate complexity of the relationships between plants and microbes.

Methods

Plant materials, root nucleus isolation, library preparation, and sequencing

Medicago seedlings were sterilized as described in Pingault et al. (2018). Eight seeds were then placed on agar B&D medium (Broughton and Dilworth, 1971) without nitrogen, and placed in a growth chamber in the dark for 4 days (26°C for 16 h and 20°C for 8 h). On the fourth day, 4 mL *E. meliloti* suspension (OD_{600nm} = 0.1), or water for the mock-inoculated samples, were applied to the seedlings' roots. The plates were placed back into the growth chamber for 48 h in the dark. On the sixth day, a subset of the *E. meliloti*-inoculated plants was transferred in vermiculite:perlite (3:1) and grown in the growth chamber (16 h daylight) for 3 weeks to confirm rhizobial infection and the formation of nodules. The remaining roots were used to collect the nuclei as described in Thibivilliers et al. (2020). The root

samples used for these experiments were around 3–4 cm long, starting from the tip and ending in the zone where root hairs are fully differentiated, thus including the zone susceptible to rhizobial infection. Briefly, roots were then chopped and passed through a 30-mm cell strainer. The filtered nuclei were purified by cell sorting using FACSaria II cell sorter (BD Biosciences). An average of 80–100,000 nuclei were collected for each sample, centrifuged, and re-suspended in phosphate buffered saline-bovine serum albumin 0.5% RNA inhibitor solution. The six sNucRNA-seq libraries (i.e., three *E. meliloti*- and three mock-inoculated root libraries) were constructed following the Chromium Single Cell 30 Library & Gel Bead Kit v3.1 protocol (103 Genomics). The sequencing of single-indexed paired-end libraries was performed on an Illumina NovaSeq 6000 platform according to the 103 Genomics recommendations.

Pre-processing of raw data, integration, clustering, and annotation

The six *Medicago* sNucRNA-seq libraries were preprocessed individually using the 103 Genomics Cell Ranger software v6.1.1.0, and then aligned against the latest version of the *Medicago truncatula* reference genome and genome annotation (at <https://medicago.toulouse.inra.fr/MtrunA17r5.0-ANR/>) (Pecrix et al., 2018). Upon removal of background contamination using the SoupX software (Young and Behjati, 2020), filtration of doublets using the DoubletDetection prediction method (Adam Gayoso, 2022), and applying a statistical threshold on the data distribution (i.e., an interval of confidence of 95% to remove outliers) (see Supplemental Figure 1 to access the parameters for each sNucRNA-seq library), the normalization of individual sNucRNA-seq datasets and their respective integration to generate UMAPs was performed using Seurat V4 (Hao et al., 2021), selecting the top 2,000 variable genes for feature selection. Integration anchors were defined for the combined set of six sNucRNA-seq datasets based on the first 20 dimensions of the canonical correlation analysis method. After integration, the dimensionality reduction was performed with the first 40 principal components to generate the UMAP projection. Besides, the clustering was generated with the method FindClusters from Seurat with a resolution of 0.6. For downstream analyses, the expression

values of each gene were calculated for each cluster using the `AverageExpression` function from Seurat.

For the annotation of cell types, the cluster-specific genes were identified with the `FindAllMarkers` function in Seurat. In addition, the expression patterns of known cell type-specific gene markers from *M. truncatula* (Supplemental Table 2 and Medicago genes orthologous to *A. thaliana* root cell-type marker genes were further analyzed (Farmer et al., 2021) (Supplemental Table 4).

UMAP visualization

For visualization purposes, all sNucRNA-seq libraries were combined using the `Cell Ranger aggr` function from 10X Genomics to combine all counts in a single `cloupe` file, and to show the UMAP coordinates projections and cell cluster assignments obtained from the Seurat analysis. We use ShinyCell (available at <https://github.com/SGDDNB/ShinyCell>), a web application allowing the visualization of single-cell data, to allow direct inquiries of the Medicago root single-cell transcriptome atlas (available from the https://shinycell.legumeinfo.org/medtr.A17.gnm5.ann1_6.expr.Cervantes-Perez_Thibivilliers_2022/ web interface).

Differential gene expression analysis

To identify DEGs, raw read counts were extracted to calculate a normalized average expression for each gene, in each cluster, and for each condition (Supplemental Table 9) before applying the `DEsingle` package (Miao et al., 2018) using a p value of less than 0.05 and FC of greater than 1.5 thresholds. This package allows the identification of DEGs between *E. meliloti*- and mock-inoculated nuclei in a raw read count matrix employing the zero-inflated negative binomial model (Wang et al., 2019).

Correlation analysis between plant root single nuclei transcriptomes

To support the functional annotation in the *M. truncatula* root clusters, we compared the transcriptomes of the different Medicago root clusters upon extracting the pseudo-bulk expression of each gene

among all clusters and then conducted Pearson's correlation analyses to reveal the most similar transcriptomes among all nuclei clusters.

To compare the transcriptome of *Medicago* and *Arabidopsis* root clusters, *A. thaliana* sNucRNA-seq datasets were obtained from previously published data (Farmer et al., 2021), based on the following SRA files: GSM4698755, GSM4698756, GSM4698757, GSM4698758, and GSM4698759. The five replicates were processed individually using the 103 Genomics Cell Ranger v6.1.1.0 pipeline, and then mapped against a reference genome constructed with TAIR10.26 genome and Araport11 annotations. The same parameters were then used for pre-processing the *Arabidopsis* datasets as previously used for the *Medicago* sNucRNA-seq datasets (see above). To correlate the *A. thaliana* and *M. truncatula* root sNuc transcriptomes, we extracted pseudo-bulk information for the one-to-one orthologs between the two plant species (CoGe; <https://genomevolution.org/coge/>) (Lyons et al., 2008; Lyons and Freeling, 2008) and conducted a Pearson's correlation analysis between the expression of these orthologous genes for each cell cluster.

Genes of interest and functional classification analyses

The Legoo knowledge base (at <https://lipm-browsers.toulouse.inra.fr/k/legoo/>) was used to identify genes of interest (Carrère et al., 2019), as well as the Mapman software (at <https://mapman.gabipd.org/>) for the analysis of gene functions (Thimm et al., 2004).

Accession Numbers Expression data are available at the Gene Expression Omnibus (GEO: GSE210881). The *Medicago* root single-cell transcriptome atlas can be accessed through https://shinycell.legumeinfo.org/medtr.A17.gnm5.ann1_6.expr.Cervantes-Perez_Thibivilliers_2022/.

Supplemental Information Supplemental information is attached to the archive record for this article.

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Author Contributions S.T. performed experiments. S.A.C.P., S.T., C.L., A.D.F., F.F., and M.L. carried out data analysis. M.L. coordinated the study. S.A.C.P., S.T., F.F., and M.L. drafted the manuscript. All authors contributed to the preparation of the manuscript.

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