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# Ectopic activation of cortical cell division during the accommodation of arbuscular mycorrhizal fungi

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#### 1 Summary

Arbuscular mycorrhizas (AM) between plants and soil fungi are widespread symbioses with a
major role in soil nutrient uptake.

In this study we investigated the induction of root cortical cell division during AM colonization
by combining morphometric and gene expression analyses with promoter activation and protein
localization studies of the cell plate-associated exocytic marker TPLATE.

7 Our results show that TPLATE promoter is activated in colonized cells of the root cortex where 8 we also observed the appearance of cells that are half the size of the surrounding cells. Furthermore, 9 TPLATE-GFP recruitment to developing cell plates highlighted ectopic cell division events in the inner root cortex during early AM colonization. Lastly, transcripts of TPLATE, KNOLLE, and 10 CYC1 are all upregulated in the same context, alongside endocytic markers AP2A1 and CHC2, 11 12 known to be active during cell plate formation. This pattern of gene expression was recorded in wild-13 type Medicago truncatula roots but not in a CSSP mutant where fungal colonization is blocked at the 14 epidermal level.

Altogether, these results suggest the activation of cell division-related mechanisms by AM hosts
during the accommodation of the symbiotic fungus.

#### 1 Introduction

2 The majority of extant plants, including both wild and domesticated crop species, is supported in its 3 uptake of soil nutrients by Glomeromycotina fungi (Lin et al., 2014) - in a beneficial symbiosis called arbuscular mycorrhiza (AM). The broad diffusion of this interaction among plant species is 4 5 considered a consequence of its ancient evolutionary origin (Bonfante & Genre, 2008). Indeed, 400-6 million-year old plant fossils already hosted hyphal structures that closely resemble the arbuscules 7 produced by living AM fungi (Remy et al., 1994; Strullu-Derrien et al., 2018). Arbuscules are highly 8 branched hyphae that develop inside living cells of the root cortex (Gutjahr & Parniske, 2013). In 9 fact, arbuscules are accommodated inside a novel cell compartment enveloped by the perifungal (or 10 periarbuscular) membrane, a specialized extension of the host plasmalemma (Pumplin & Harrison, 11 2009). This symbiotic interface compartment assembles within the pre-penetration apparatus (PPA), 12 a broad, nucleus-associated cytoplasmic aggregation, where the secretory process is focused and 13 coordinated (Genre et al., 2005; 2008).

14 Both PPA organization and fungal accommodation depend on the activation of a conserved signal 15 transduction pathway (Delaux et al., 2015), the so-called Common Symbiotic Signaling Pathway, or 16 CSSP (see Oldroyd, 2013 for a comprehensive review) and represent the earliest visible response of 17 plant cells to AM fungi (Genre et al., 2005). The CSSP is believed to transduce diffusible AM fungal 18 signals through the sequential activation of a number of proteins, starting from receptor-like kinases 19 in the plasma membrane and culminating with the nuclear localized kinase DOESN'T MAKE 20 INFECTIONS 3 (DMI3; Lévy et al., 2004), which mediates the regulation of gene expression by 21 activating a set of transcription factors (Oldroyd, 2013; Zipfel and Oldroyd, 2017; Floss et al., 2017). 22 Altogether, CSSP activation is essential for root colonization (Novero et al., 2002; Lévy et al 2004, 23 Kistner et al., 2005; Parniske, 2008), as demonstrated by the block of PPA formation (Genre et al., 24 2005) and epidermal penetration in *dmi3* mutants (Morandi et al., 2005).

Biogenesis of the perifungal membrane has been described as a massive exocytic process that involves cytoskeletal rearrangements as well as the preemptive accumulation of an extensive quantity of secretory membranes, such as trans-Golgi network (TGN) compartments and vesicles (Genre *et al.*, 2012). Alongside secretion-related endomembrane compartments, transmission electron microscopy imaging of PPA aggregates has shown the abundant presence of multivesicular bodies or late endosomes - indicating that endocytic processes are likely also in place during interface biogenesis (Genre *et al.*, 2012), even if their role has not been investigated further.

32 Such intense and targeted membrane dynamics have few analogs in plant cells and possibly the most 33 notable similarity is with cell division. This process initiates with vacuole partitioning by fusion of 34 multiple trans-vacuolar cytoplasmic bridges (phragmosome) and nuclear positioning in the cell center

1 (Smith, 2001; Lloyd & Buschmann, 2007). Phragmosome formation is followed by regular mitosis 2 and cell plate assembly through the progressive fusion of TGN-derived cytokinetic vesicles (Segui-3 Simarro et al., 2004; Smertenko et al., 2017). Intriguingly, Siciliano et al., (2007) reported that 4 histone H2B1 gene was induced in root segments of Medicago on which hyphopodium formation had 5 occurred, hinting at the possibility that cell cycle determinants are active during early AM 6 colonization. The involvement of cell division-related processes has also been proposed for the 7 symbiotic nitrogen fixation, where infection thread growth is coupled to cell cycle reactivation and 8 cell division events (Timmers et al., 1999; Fournier et al., 2008; Breakspear et al., 2014; Downie, 9 2014).

10 During plant cell division, the massive exocytic process directed by cytoskeletal fibers in the 11 phragmoplast (Lee & Liu, 2013; Boruc & van Damme, 2015) is associated with endocytic recycling 12 of surplus membrane. In fact, next to cell plate specific markers such as the syntaxin KNOLLE, other 13 plasma membrane proteins also strongly accumulate at the forming cell plate (Van Damme et al., 14 2006; Richter et al., 2014), among which, the adaptin-related protein TPLATE has been shown to 15 accumulate on both the cell plate membranes and plasmalemma at the cortical division zone, where 16 the cell plate will eventually fuse (Van Damme et al., 2006; Van Damme et al., 2011; Gadeyne et al., 17 2014). TPLATE is the founding member of an octameric protein complex, the TPLATE complex 18 (TPC), which acts largely together with Adaptor Protein complex 2 (AP-2) and other components 19 (such as Dynamin-related proteins) and the clathrin scaffolding proteins for the assembly of the 20 clathrin endocytic machinery at the plasma membrane (Gadeyne et al., 2014). The accumulation of 21 TPC members at the cortical division zone has been proposed to facilitate membrane reorganization 22 during cell plate fusion with the peripheral plasmalemma (Boutté et al., 2010; Van Damme et al., 23 2011). In short, TPLATE appears to be a promising marker to investigate the occurrence of both cell 24 division- and endocytosis-related processes during the accommodation of AM fungi.

25 Here, we report that *TPLATE* expression is activated in cortical root cells colonized by arbuscular 26 mycorrhizal fungi, where we also observed the appearance of arbusculated cells with half the size of 27 the surrounding cortical cells. Combined with TPLATE-GFP recruitment to forming cell plates, this 28 allowed us to identify ectopic cell division events that are induced in the inner root cortex during AM 29 colonization of the outer root tissues. Lastly, TPLATE transcripts, alongside other cell division and 30 endocytic markers, are upregulated in the colonized area of wild-type roots but not in a CSSP mutant 31 where fungal colonization is blocked at the epidermal level. Taken together, these results suggest that 32 extant plants activate cell division-related mechanisms during AM fungal accommodation.

#### 1 Materials and Methods

2

#### 3 **Plant materials**

4 The generation of composite L. japonicus plants was performed using Agrobacterium rhizogenes

5 strain Ar1193 carrying the*pLjTPLATE:GUS* vector, according to the protocol by Stougaard (1995)

6 on wild-type seedlings of MG20 genotype (see below).

*Medicago truncatula* Jemalong A17 and *Lotus japonicus* MG20 plants were grown from surface
 sterilized seeds in pots containing sterile quartz sand and respectively inoculated with *Gigaspora margarita* or *Funneliformis mosseae* and watered with Long-Ashton solution (Hewitt, 1966).

10 Agrobacterium rhizogenes-transformed root organ cultures (ROCs) expressing the 35S::AtTPLATE-

11 GFP vector (Van Damme et al., 2004) were obtained from Medicago truncatula Jemalong A17 wild-

- 12 type and *dmi3-1* plants (Sagan et al., 1995; kindly provided by M. Chabaud, LIPM, INRA, Toulouse, 13 France), according to Boisson-Dernier et al. (2001). ROC generation was repeated in two independent 14 experiments for both lines, with overlapping results in terms of GFP fluorescence pattern. ROCs from 15 Daucus carota var Sativus expressing the same vector were obtained according to Bécard & Fortin 16 (1988). For both species, transformed roots with a high level of fluorescence were selected 21 days 17 after transformation, decontaminated and subcultured on M medium (Bécard & Fortin, 1988) at 25°C 18 in the dark for subsequent use as hairy roots. In each case, a single representative clone was chosen 19 for further studies. Transformation efficiency and expression of 35S::AtTPLATE-GFP was checked 20 with GFP specific primers (Supporting Information Table S1) on both genomic DNA and cDNA 21 obtained from all the selected lines. Wild type and *dmi3-1* mutant *M. truncatula* ROCs expressing a 22 nuclear cameleon YC2.1 probe were already available in the lab (Chabaud et al., 2011; Genre et al.,
- 23 2013) and were used for all gene expression analyses.
- 24

## 25 **Fungal materials**

26 Different AM fungi were chosen depending on how their characteristics suited the experimental 27 setup. The AM fungus Funnelliformis mosseae (strain BEG 12), supplied by MycAgroLab 28 (http://www.mycagrolab.com/), was used to inoculate L. japonicus composite plants with a 10% ratio 29 of inoculum to total substrate volume; Gigaspora margarita (BEG34, International Bank for 30 Glomeromycota, University of Kent, Canterbury, UK) was used to inoculate M. truncatula whole 31 plants (50 spores per pot) and ROCs (Chabaud et al., 2002) for gene expression analyses. Gigaspora 32 gigantea (isolate HC/FE30, Herbarium Cryptogamicum Fungi, University of Torino, Italy), which is 33 characterized by strong cytoplasmic autofluorescence (Genre *et al.*, 2005), was used to inoculate 34 ROCs for live confocal imaging. Spores of both G. gigantea and G. margarita were collected from 1 pot cultures in sand (with leek and clover, respectively), surface-sterilized and stored at 4°C according

2 to Bécard & Fortin (1988).

3

## 4 GUS assay

5 A *LjTPLATE* promoter fragment of 2073 bp (Supporting Information Fig. S1) was PCR-amplified 6 from genomic DNA by using specific primers (Supporting Information Table S1). The resulting 7 promoter fragment was cloned into pDONR221 (Invitrogen) and recombined upstream of the GUS 8 gene into pKGWFS7.0 (Gateway vector), modified to contain the red fluorescent marker DsRed 9 under the control of the constitutive *Arabidopsis* UBIQUITIN10 promoter (pUBQ10; Limpens *et al.*, 10 2005).

11 Half of the 30 L. japonicus composite plants selected for DsRed fluorescence were inoculated with

*F. mosseae* and the remaining were used as controls. Root sampling was repeated at 14, 21 and 42days post inoculation.

14 Hairy roots expressing DsRed fluorescence were selected using a stereomicroscope (Leica M205 FA) 15 from both inoculated and control plants; excised roots were immediately submerged in freshly 16 prepared GUS buffer (0.1 M potassium phosphate buffer pH 7, 5 mM K4Fe(CN)6, 5 mM K3Fe(CN)6, 17 0.3% Triton X-100, 0.3% X-GlcA; Duchefa Biochemie), incubated overnight at 37°C in the dark, 18 destained with 70% ethanol and washed twice in distilled water. Part of the roots were then counter-19 stained with 0.01% (w/v) acid fuchsin in lactoglycerol (lactic acid:glycerol: water, 14:1:1; Kormanik 20 & McGraw, 1982) to highlight intraradical fungal structures. Samples were then either directly 21 mounted on a microscope slide for observation, or previously embedded in agarose (5%) and cut into 22 100 µm thick vibratome slices. Observations were done using a bright field microscope (Nikon 23 Eclipse E400).

24

## 25 Confocal microscopy imaging

26 The targeted AM inoculation technique for studying early stages of the symbiotic association between 27 Gigaspora species and transformed root cultures, developed by Chabaud et al. (2002) and adapted 28 for confocal observation by Genre et al. (2005), was applied to both Medicago truncatula and Daucus 29 carota ROCs expressing 35S::AtTPLATE-GFP. An upright Leica TCS SP2 confocal microscope 30 fitted with a long distance 40X water-immersion objective (HCX Apo 0.80) was used for imaging 31 living ROCs directly in the Petri dishes. The argon laser band of 488 nm was used to excite both GFP 32 and G. gigantea autofluorescence. The two signals were distinguished using specific emission 33 windows: 500 to 525 nm for GFP and 590 to 630 nm for fungal autofluorescence. The latter channel 34 was then false-colored in red to maximize the contrast in overlapping images.

#### 1 Morphometric analyses

2 In order to visualize intraradical fungal structures and perform a quantitative analysis of cortical cell 3 morphology, root samples from 8 weeks old pot-grown plants were harvested and fixed in phosphate-4 buffered saline (PBS), pH 7.2, containing 1% formaldehyde and 10% dimethyl sulfoxide (DMSO) 5 for 24 hours at 4°C. Fixed roots were sliced using a Vibratome (Oxford Vibratome® sectioning 6 system) and stained with wheat germ agglutinin and propidium iodide (Kojima et al., 2014) or acid 7 fuchsin (Kormanik & McGraw, 1982) to highlight intraradical fungal structures and plant cell walls, 8 respectively. The argon laser line at 488nm was used to excite all fluorochromes and emission 9 windows were set at 500-530nm to record FITC, 600-700nm for propidium iodide and fuchsin 10 fluorescence. A total of 42 optical sections (375 x 375 µm) were recorded from 13 independent root 11 segments sampled from 3 different uninoculated plants of *M. truncatula*; 47 optical sections from 8 12 independent root segments (3 plants) were used for colonized M. truncatula; 22 optical sections (6 13 root segments from 3 plants) were used for uninoculated L. japonicus and 23 optical sections (6 root 14 segments from 3 plants) for colonized L. japonicus. For all our analyses, 0.5cm-long segments from 15 lateral roots of comparable size, length and position in the root system were sampled from both 16 inoculated and uninoculated plants.

In addition, the number of split cells (Supporting Information Table S2) was counted in the inner
cortical layer of live uninoculated or inoculated ROCs of *D. carota* and wild type or *dmi3-1 M. truncatula* expressing *35S::AtTPLATE-GFP*. A minimum of 22 optical sections (375 x 375µm) from
at least 5 different roots of comparable size, length and position, belonging to 3 independent ROCs
was used for each condition.

22

#### 23 Quantitative gene expression analyses

24 Wild type and *dmi3-1 Medicago truncatula* ROCs were inoculated with *Gigaspora margarita*; fungal 25 development and root colonization were followed under a stereomicroscope to identify hyphopodium 26 development. A few of these samples were further observed in fluorescence microscopy (Leica TCS-27 SP2) to verify the repositioning of YC2.1 fluorescent nuclei under the hyphopodia - a hallmark of 28 epidermal cell prepenetration responses. After 48/60 hours - required for the development of young 29 colonization units - the corresponding root segments were excised and immediately frozen in liquid 30 nitrogen. All root tips were removed before sampling. Four biological replicates - each consisting of 31 a pool of ten 1cm long root segments - were collected from both uncolonized and colonized wild type 32 ROC lines; and three replicates were collected from the corresponding *dmi3-1* lines. Uncolonized 33 root segments were selected from lateral roots with a similar development and position in the root 34 system architecture as the colonized segments from inoculated ROCs.

Total RNA was isolated from colonized and uncolonized samples and prepared using the RNeasy<sup>TM</sup> Plant Mini kit (Qiagen, Hilden, Germany). Plant material was processed with a mechanical disruption procedure using a TissueLyser system (Qiagen Retsch GmbH, Hannover, Germany) and further treated following the manufacturer's instructions. RNA quality and quantity were evaluated by photometric measurements and on agarose gels.

6 To remove traces of genomic DNA before cDNA synthesis, samples were treated with TURBO<sup>TM</sup>

7 DNase (Ambion) according to the manufacturer's instructions. The RNA samples were routinely 8 checked for DNA contamination by RT-PCR analysis, using primers for *MtTEF* (Supporting 9 Information Table S1). Conventional PCR assays on fungal genomic DNA excluded any cross-10 hybridization of *M. truncatula* specific primers.

11 For single-strand cDNA synthesis, about 700 ng of total RNA was denatured at 65 °C for 5 min and

12 then reverse-transcribed at 25 °C for 10 min, 42 °C for 50 min and 70 °C for 15 min. The final volume

13 was 20 $\mu$ l and contained 10 $\mu$ M of random primers, 0.5mM dNTPs, 4 $\mu$ l 5Xbuffer, 2  $\mu$ l 0.1 M DTT and

- 14 1µl Super-Script II (Invitrogen, Carlsbad,CA, USA).
- 15 Specific primers for *MtTPLATE*, *MtAP2A1*, *MtCHC1* and *MtCHC2* were designed on putative *M*.
- 16 truncatula CDS sequences from NCBI database (http://www.ncbi.nlm.nih.gov/) and designed using
- 17 SerialCloner software specifically for qRT-PCR analyses (Supporting Information Table S1). Primer
- 18 validation was obtained with conventional PCR asssay on *M. truncatula* cDNA to avoid overlapping

19 with unspecified sequences, and testied on fungal genomic DNA.

- 20 qRT-PCR experiments were carried out in a final volume of 15µl containing 7.5µl of iTaq<sup>™</sup> 21 Universal SYBR® Green Supermix (Bio-Rad), 1µl of 3µM specific primers, and 10 ng of cDNA. 22 Samples were run in the Rotor Gene apparatus (Qiagen) using the following program: 10min pre-23 incubation at 95°C, followed by 40 cycles of 15s at 95°C, and 30s at 60°C. Each amplification was 24 followed by melting curve analysis (60–94°C) with a heating rate of 0.5°C every 15s. All reactions 25 were performed with three technical replicates and only Ct values with a standard deviation that did 26 not exceed 0.3 were considered. The comparative threshold cycle method (Rasmussen, 2001) was 27 used to calculate relative expression levels using the plant *MtTEF* as reference genes.
- 28

# 29 Accession numbers

Sequence data from this article can be found in the NCBI data libraries under the following accession
numbers: LjT06B17 (*LjTPLATE* promoter from 47759 to 49830); MTR\_5g012010 (*MtKNOLLE*);
MTR\_7g089730 (*MtCYC1*); MTR\_5g082900 (*MtCHCL1*); MTR\_3g070940 (*MtCHCL2*);
MTR\_7g031450 (*MtTPLATE*); MTR\_2g084610 (*MtAP2A1*); MTR\_1g105120 (*MtBCP*);
MTR\_6g021805 (*MtTEF*).

# 2 **Results**

3

#### 4 TPLATE expression is activated in AM colonized areas of the root

5 In order to highlight the involvement of the endocytosis and cell division marker TPLATE in the 6 process of AM fungal accommodation, we studied the pattern of TPLATE promoter activity by 7 expressing a *pLjTPLATE:GUS* transcriptional reporter (2kb promoter) in *Agrobacterium rhizogenes*-8 transformed roots from wild type Lotus japonicus and applying the GUS assay to excised roots. More 9 than 50 roots were screened in brightfield microscopy and intense GUS-positive staining was 10 observed in most root tips (Supporting Information Fig. S2a,b), consistent with the role of TPLATE 11 in cell division. In roughly 1 out of 5 root tips, the expression pattern was less uniform and limited to 12 small groups of meristematic cells (Supporting Information Fig. S2c,d) or isolated cell pairs 13 (Supporting Information Fig. S2e). Taken together, the diverse meristematic patterns of 14 *pLjTPLATE:GUS* staining appear to highlight a transient upregulation of the TPLATE promoter above a low basal level as previously described in Arabidopsis (Van Damme et al., 2006; Brady et 15 16 al., 2007), even if literature data and public databases indicate that TPLATE mRNA is expressed in 17 virtually all cells due to its constitutive role in endocytic processes, as confirmed by both protein 18 localization specifically in root and hypocotyl epidermal cells as well as functional studies on A. thaliana seedlings (Van Damme et al., 2006, Van Damme et al., 2011; Gadeyne et al., 2014, Sanchez-19 20 Rodriguez et al., 2018). Moreover, GUS staining was often observed in the root vascular cylinder in 21 both young (Supporting Information Fig. S2a) and differentiated areas of the root (Supporting 22 Information Fig. S2f, h), possibly related to endocytic processes linked to phloem activity (Liesche 23 et al., 2008).

24 AM colonization induced an evident change in this TPLATE expression pattern, with the appearance 25 of intense GUS-positive areas associated with intraradical fungal structures (Supporting Information 26 Fig. S2g), independent of the sampling time (14, 21 or 42 days post inoculation). The observation of 27 over 60 vibratome sections from 10 independent root samples consistently indicated that GUS activity 28 was particularly intense in inner cortical cells containing arbuscules (Supporting Information Fig. 29 S2i). This was further confirmed by the analysis of acid fuchsin counter-stained samples (20 root 30 segments overall), where the precise location of intraradical fungal structures could be detected more 31 easily in longitudinal sections (Fig. 1a-c).

In conclusion, the GUS assay provided strong indications that AM colonization boosts *TPLATE* promoter activity in arbusculated root cortical cells.

- 34
- 35

#### 1 Split cells are observed in the cortex of colonized roots

In addition to the correlation between arbuscule presence and *TPLATE* promoter activation, a consistent feature was observed in dozens of longitudinal root sections: the presence of one or more couples of arbuscule-containing cells, half the size of the surrounding (either colonized or uncolonized) cortical cells (Fig. 1d-f). In fact, the roughly square shape of such cells made them stand out very clearly among the more elongated cortical cells.

7 To better investigate this intriguing observation and exclude that ectopic cell divisions could be 8 caused by our use of A. rhizogenes-transformed composite plants, we performed an extensive 9 screening of longitudinal root sections from both L. japonicus and M. truncatula wild type plants, 10 grown in pots for 8 weeks with or without AM inoculation. Fluorescent staining of fungal and plant 11 walls made split cells easy to recognize by confocal imaging (see materials and methods for details). 12 Representative images of L. japonicus samples (from a total of 6 colonized and 6 uninoculated root 13 segments) are shown in Fig. 2. Couples of short cortical cells were only occasionally observed in 14 uninoculated plants (Fig. 2a, b). By contrast, split cells were regularly found in AM colonized roots 15 (Fig. 2c-e), alongside arbusculated and non-arbusculated cortical cells of regular size. In more detail, 16 both unpenetrated (Fig. 2c, d) and arbusculated split cells (Fig. 2c-e) were observed in the colonized 17 area, including couples where one cell contained an arbuscule and the second one did not (Fig. 2c), 18 and couples where the two arbuscules had different (Fig. 2c,d) or comparable stages of development 19 (Fig. 2e).

The table in Fig. 2, resumes our quantitative analysis of split cell occurrence in control and mycorrhizal plants of both *L. japonicus* and *M. truncatula*. We quantified the average number of split cell couples per image and the number of images containing at least one couple of split cells. In both plants, AM colonization significantly increased (t test <0.01) the average number of split cells: over 6 times in *L. japonicus* (from 0.71 to 4.80) and over 4 times in *M. truncatula* (from 0.58 to 2.53). The percent of images containing split cells was also increased in both AM hosts, from 36% to 96% in *L. japonicus* and from 36% to 66% in *M. truncatula*.

In short, the induction of anticlinal cell divisions was confirmed in the inner cortex of colonized areas in untransformed *L. japonicus and M. truncatula* plants, excluding a causal connection with *A. rhizogenes* infection. Secondly, our detailed quantification of split cell occurrence demonstrated a statistically supported relation between AM colonization and inner cortical cell divisions. Lastly, our observations of split cells where each contained differently developed arbuscules, alongside uncolonized split cells next to the arbusculated area, were suggestive of cell divisions occurrence in advance of arbuscule development.

#### 1 Live imaging of ectopic cortical cell divisions in mycorrhizal roots using TPLATE-GFP

2 To clarify whether the AM-dependent induction of cortical cell division was indeed triggered before 3 fungal entry, we expressed a 35S::AtTPLATE-GFP fusion (originally developed in Arabidopsis 4 thaliana and tobacco BY-2 cells by Van Damme et al., 2004; 2006), in A. rhizogenes-transformed root organ cultures (ROCs). Since L. japonicus ROCs can only very rarely be colonized in our 5 6 experimental conditions, we used M. truncatula (legume) and D. carota (non-legume) ROCs instead, 7 two more amenable models for live cell imaging of symbiotic interactions (Chabaud et al., 2002; 8 Genre et al., 2008; Fournier et al., 2015). 9 In vivo observations of at least 10 uninoculated M. truncatula ROCs confirmed the predicted 10 localization of TPLATE-GFP (Van Damme et al., 2004; 2006) with a major accumulation of the 11 construct along newly formed cell walls in the apical meristem (Supporting Information Fig. S3a,b). 12 By increasing microscope sensitivity, a weaker background fluorescence could also be recorded in 13 the cytoplasm (likely related to 35S-driven overexpression) and at the plasma membrane of both

14 meristematic and differentiated cells (Supporting Information Fig. S3c), in agreement with the 15 reported plasma membrane targeting of TPLATE (Van Damme *et al.*, 2011, Gadeyne *et al.*, 2014) in 16 non-dividing cells.

Our observation of over 50 *G. gigantea* infection units, intended as root areas underlying the sites of hyphopodium contact, revealed an intense TPLATE-GFP labeling along a few anticlinal cell walls in the inner root cortex of both *M. truncatula* (Fig. 3a,b) and *D. carota* ROCs (Fig. 3c; Supporting Information Fig. S4a,b).

21 In particular, GFP-marked anticlinal cell walls were observed in the inner cortex of both plant species 22 as soon as hyphopodia developed on the root surface (Fig. 3a,c; Supporting Information Fig. S4a) 23 and until the first hyphae reached the inner cortex, about 2 days later (Fig. 3b; Supporting Information 24 Fig. S4b). Such GFP-labeled walls appeared to separate two square-shaped cells that corresponded in 25 shape to the previously described split cells (Fig. 1; Fig. 2). In fact, couples of split cells were 26 constantly observed in the colonized areas of the inner cortex of TPLATE-GFP-expressing ROCs 27 (Fig. 3d; Supporting Information Fig. S4b,c), although their walls were not as strongly labeled by 28 GFP in more advanced stages of root colonization (Supporting Information Fig. S4c) nor after 29 arbuscule development (Fig. 3d).

Comparable couples of square-shaped cells were only occasionally observed in the inner cortex of corresponding control segments from uninoculated roots of the same age and size as the colonized segments, (Supporting Information Fig. S4d), and their transverse wall was anyway never highlighted by intense TPLATE-GFP accumulation. These observations were strongly supported by consecutive imaging of a single site in an inoculated carrot root with a gap of 24 hours between each observation (Fig. 3e-h). At the first time point (Fig 3e-g), hyphae developing from a single hyphopodium had colonized the epidermis (Fig. 3e) and outer cortex (Fig. 3f). A deeper focal plane reaching the inner cortex (Fig. 3g) revealed an intensely labeled cell plate being formed. When the same focal plane was imaged 24 hours later (Fig 3h), the TPLATE-GFP signal had significantly faded from the mature cross wall and - in the meantime - four additional inner cortical cells had divided in the same area. On the same line, quantitative morphometric analyses performed on both *D. carota* and *M. truncatula* 

confirmed a significant increase in the number of split cells in mycorrhizal compared to uninoculated
ROCs (Supporting Information Table S2).

9 Based on cell morphology and the analogous TPLATE-GFP labeling of recent cell walls in the root
10 meristem and in the colonized area of the mature cortex, we interpreted our observation as evidence
11 of an early, AM-dependent induction of cell division in the root inner cortex.

12

# 13 Ectopic cortical cell division is not induced in *Mtdmi3-1* mutants

14 Because cortical cell divisions were observed since the early steps of root colonization and often 15 occurred in cells that were not in direct contact with the fungus, we decided to investigate whether 16 this process was induced in the CSSP mutant *Mtdmi3-1*, where fungal colonization is arrested at the 17 epidermis level, with the development of extensive and branched hyphopodia (Catoira et al., 2000; 18 Morandi et al., 2005). By expressing the same TPLATE-GFP construct in this genotype, we first 19 confirmed construct functionality observing the expected GFP localization at newly formed cell walls 20 in the root tip meristem (Fig. 4a). Our careful observation of 8 independent hyphopodium-associated 21 root segments (Fig. 4b-e) did not reveal any fungal penetration of the root inner tissues and no 22 statistically significant change was observed in the number of split cells between uninoculated and 23 inoculated *dmi3-1* roots (Supporting Information Table S2). On this basis we concluded that the 24 activation of cortical cell divisions in M. truncatula is dependent on fungal presence inside the root -25 which in turn requires DMI3-mediated signaling within the CSSP.

26

## 27 Cell division and TPC-associated endocytic markers are upregulated in mycorrhizal roots

Based on our observations of cell division induction and TPLATE involvement in AM fungal accommodation processes, we used qRT-PCR to analyse the expression of cell division markers and TPC-interacting proteins that act in endocytic processes associated with cell plate assembly.

31 Gene expression was analyzed in *M. truncatula* ROCs (Chabaud et al., 2011; Genre et al., 2013)

32 obtained from both wild type and *dmi3-1* plants (Genre *et al.*, 2005).

33 Two acknowledged cell division markers were chosen. MtKNOLLE encodes a syntaxin (homolog to

34 *AtSYP111*) exclusively localizing to cell-plate directed vescicles and essential for cytodieresis (*knolle* 

1 mutants develop incomplete cell walls and multinucleated cells; Lukowitz et al., 1996). Our second 2 cell division marker was MtCYC1, encoding a putative Cyclin-like F-box protein, previously reported 3 to be upregulated by 3,01 log2 FC in arbusculated vs non-colonized cells (data from the M. truncatula 4 genome array performed on a population of root cortical cells colonized with *Rhizophagus irregularis* 5 and isolated through laser capture microdissection by Gaude et al., 2012). Both transcripts were 6 significantly upregulated in mycorrhizal root samples compared to uninoculated roots in WT, while 7 no upregulation was recorded in *dmi3-1* mutants (Fig. 5). This result was consistent with our 8 hypothesis that cell division-related mechanisms are involved in AM colonization.

9 We also analysed the expression of four TPC-related endocytic markers. MtTPLATE and the alpha 10 subunit of the Adaptor Protein complex 2 (MtAP2A1) - both involved in clathrin-mediated 11 endocytosis (Gadeyne et al., 2014; Di Rubbo et al., 2013) - were constantly and significantly more 12 abundant in mycorrhizal root segments compared to controls in wild type M. truncatula, while their 13 expression did not change significantly in *dmi3-1* (Fig. 5). A weak but statistically significant 14 upregulation was observed for *Clathrin Heavy Chain 2 (MtCHC2)*, in mycorrhizal WT samples, while 15 this was not the case for *Clathrin Heavy Chain* 1 (Fig. 5). 16 Lastly, as a marker of active AM colonization, we analyzed the expression of the AM-specific Blue

*Copper-binding Protein MtBCBP* (Parádi *et al.*, 2010) in the same samples. This confirmed that the upregulation of our selected markers was strictly associated with *MtBCBP* expression (Supporting Information Fig. S5). On this basis, we propose that cell division markers and actors of clathrinmediated endocytosis act during AM fungal accommodation, in line with our microscopic observations.

22

- 1 Discussion
- 2

#### 3 The recruitment of cell division mechanisms in AM

4 During symbiotic interface biogenesis, a novel membrane-bound compartment is assembled in the 5 middle of the cell, an area that is normally occupied by the vacuole in fully differentiated root cells. 6 Displacing the tonoplast is indeed among the first consequences of PPA-associated nuclear 7 movements in both epidermal and cortical cells (Genre et al., 2005; 2008). As the PPA assembly 8 proceeds, the vacuole is further deformed (and reduced in volume), until a mass of cytoplasm (the 9 PPA aggregate) occupies the portion of the cell volume where the symbiotic interface and 10 intracellular hyphae are about to develop (Genre et al., 2012). An analogous mechanism is 11 orchestrated when differentiated plant cells undergo mitosis: the vacuole has to be reduced - and 12 eventually split - so that the correct positioning of nucleus, mitotic spindle, phragmoplast and cell plate can generate the cell division plane (Van Damme, 2009; Smertenko et al., 2017). Also in this 13 14 case, initial vacuole remodeling is obtained by the coordinated movement of the nucleus and the 15 assembly of phragmosome cytoplasmic bridges that cross the vacuole along the equatorial plane and 16 eventually fuse into one central mass of cytoplasm encompassing the nucleus (Lloyd & Buschmann, 17 2007).

In addition to such similarities in cell dynamics, our results suggest a more direct relationship between the two processes. The upregulation of several cell division markers in colonized root areas and *TPLATE* expression pattern, that indicate an activation of gene expression in colonized cortical cells, are both consistent with a reactivation of cell cycle-related processes. This is reinforced by the recurring presence of split cells in the colonized region of roots from phylogenetically distant hosts and the direct observation of dividing cortical cells in the vicinity of intraradical hyphae and hyphopodia.

25 Altogether, we interpret our results as indicative of the reactivation of cell cycle-related processes in 26 the inner cortex since early root colonization. The biological meaning of this mechanism remains to 27 be fully clarified; nevertheless, literature on AM fungal accommodation provides intriguing clues: 28 the symbiotic interface materials have been described as very similar to the primordial wall found in 29 cell plates (Balestrini & Bonfante, 2014); a temporary, global re-routing of exocytic processes has 30 been proposed to take place in both cell plate (Lam et al., 2008) and periarbuscular interface 31 deposition (Pumplin et al., 2012); lastly, evidence of ploidy increases in mycorrhizal root systems 32 (Berta et al., 2000; Lingua et al., 2002; Fusconi et al., 2005; Bainard et al., 2011) suggests the onset 33 of endoreduplication (deriving from the arrest of the mitotic cycle) during AM colonization, even if 34 the precise location and timing of such endored uplication events remains to be defined (Genre et al.,

2008). In the light of our current results, it is tempting to speculate that a mechanistic analogy links
the exocytic processes that lead to cell plate deposition and symbiotic interface biogenesis, even if
further investigation is needed to test this hypothesis.

4

#### 5 Membrane dynamics during fungal accommodation: a role for endocytosis?

Our results on the upregulation of endocytic effectors such as TPLATE, CHC2 and AP2A1 during
AM fungal accommodation are suggestive of a role for clathrin-mediated endocytosis (CME)-related
processes in this symbiotic interaction.

9 Even if further investigation is needed to clarify this hypothesis, the involvement of CME 10 complements previous descriptions of the exocytic origin of the symbiotic interface in both AM and 11 nitrogen fixing symbiosis (Ivanov *et al.*, 2012; Genre *et al.*, 2012; Fournier *et al.*, 2015; Zhang et al., 12 2015; Barker *et al.*, 2017): all major exocytic events in the plant cell are associated with endocytic 13 membrane recycling (Samaj *et al.*, 2004; Ketelaar *et al.*, 2008). Significantly, this also applies to cell 14 plate formation (Dhonukshe *et al.*, 2006; McMichael & Bednarek, 2013). 15 Indeed, CME has recently been shown to play a fundamental role in legume-rhizobium interactions: 16 a 60% reduction in the abundance of *L. immunicus Clerknin Hermy Chrin L (LiCHCL)* transmitte (but

16 a 60% reduction in the abundance of *L. japonicus Clathrin Heavy Chain 1 (LjCHC1)* transcripts (but 17 not *LjCHC2*) has been shown to strongly reduce rhizobial infection and nodule development (Wang 18 *et al.*, 2015). In the light of this finding, the consistent 30% upregulation of *M. truncatula CHC2* (but 19 not *MtCHC1*) during AM colonization opens the intriguing possibility that the two genes act in 20 alternative pathways controlling legume mycorrhization or nodulation, although more detailed 21 analyses going beyond the scope of the present research will be required to clarify this point.

22

#### 23 **Evolutionary context**

24 The reactivation of the cell cycle is not unique to AM interactions, and has indeed been described 25 much earlier in legume-rhizobium interactions. The study by Xiao and colleagues (2014) provides 26 one of the most thorough descriptions of cell proliferation during nodule development, using 27 Medicago truncatula as a model system. In this legume, rhizobium inoculation triggers anticlinal 28 divisions that start in the pericycle and then extend to the endodermis, inner and middle cortical cell 29 layers, with the latter eventually generating the nodule meristem. The authors describe this process 30 as reminiscent of lateral root development, in line with the hypothesis of nodule evolutionary origins 31 from modified lateral roots. In the case of AM, however, cell cycle-related processes (including 32 marker gene regulation and the appearance of split cells) were restricted to a few, often sparse cells 33 in the inner cortical layer. This important difference indicates that the AM-related response is more 34 limited and also lacks the coordination observed in neo-organogenesis of both nodules (Xiao et al.,

1 2014) and lateral roots (Downie, 2014). On this basis, we cannot define a direct relationship between 2 the AM-dependent induction of cortical cell divisions and nodule initiation. Nevertheless, isolated 3 split or endoreduplicated cells are often observed in the vicinity of the infection thread, as it develops 4 across the root cortex. Such cell divisions are clearly unrelated to those that originate the nodule primordium (Xiao et al., 2014), and their observation led to the conclusion that pre-infection thread 5 6 assembly (a transvacuolar cytoplasmic aggregation that is remarkably similar to the PPA) could 7 involve reactivation of cell cycle (Yang et al., 1994) and cell division-related mechanisms (Downie, 8 2014). Based on our current results we interpret the triggering of cortical cell divisions as a conserved 9 developmental trait in both symbioses.

The induction of ectopic cell divisions in roots colonized by the evolutionarily more ancient AM symbiosis is particularly intriguing - also in the light of the frequent observation of couples of short arbusculated cells in *Horneophyton ligneri* fossils from the Rhynie Chert (Strullu-Derrien *et al.*, 2018) - and suggests that this response is not related to developmental innovations introduced by legumes during their co-evolution with nitrogen-fixing bacteria, but rather a conserved strategy derived from the 400 million year-old AM symbiosis.

16 17

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28 29

## 30 Author contributions

G.R. designed the experiments, developed the transgenic lines, performed microscopy, morphometric and gene expression analyses and wrote the text. G.C. performed gene expression and morphometric analyses, transgenic line production and contributed to the writing. V.F. performed gene expression analyses and developed the *LjTPLATE:GUS* vector. M.C. developed the *LjTPLATE:GUS* vector.

1	V.V. performed gene expression analyses, transgenic line production and contributed to the writing.
2	D.V.D. provided the AtTPLATE-GFP vector and contributed to the writing. A.G. designed the
3	research and experiments and wrote the text.
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21	Fig. 1. <i>TPLATE</i> expression pattern in mycorrhizal roots of <i>L. japonicus</i> . Following the GUS assay
22	reaction (blue), root samples were counterstained with acid fuchsin (purple) to outline <i>F. mosseae</i>
23	hyphae (hy) and arbuscules (ar). A young colonization unit is shown in (a) with limited intraradical
24	development from a single hyphopodium (hp) at 14 days post inoculation (dpi); (b) intense TPLATE
25	promoter activity is associated with arbuscules in the inner cortex. (c) Details of GUS-positive
26	staining in arbusculated cells. The largest hyphal branches are indicated by arrowheads. (d-f) GUS
27	staining of arbuscule-containing cells in the inner cortex at 14 dpi (c,d) and 21 dpi (e,f): a few
28	arbuscules have developed inside split cells (white dashed line). Bars = $100 \mu m$ in a; $20 \mu m$ in (b,d;
29	40 µm in (c,e,f).
30	
31	Fig. 2. Qualitative and quantitative analysis of cortical cell divisions in colonized roots of pot-
32	grown L. japonicus and M. truncatula. (a-d) Representative images of untransformed L. japonicus
33	samples where the double staining of <i>F. mosseae</i> walls with WGA-FITC (green) and plant walls with

34 propidium iodide (red) indicated an increase in the number of split cells in AM colonized roots. (a,b)

1 Control uninoculated plants only occasionally displayed couples of cortical cells (white dashed line) 2 that were roughly half as long as the neighboring cells (asterisk). By contrast, split cells were frequent 3 in AM colonized roots (c-e), alongside cortical cells of regular size (asterisks). Four main conditions 4 were found: uncolonized split cells in the vicinity of arbuscules (c, left; d, right); split cells where one 5 cell contained an arbuscule and the second one did not (c, center); split cells containing arbuscules at 6 different stages of development (c, right; d, left); split cells containing arbuscules with comparable 7 morphology (e). Bars =  $40\mu m$  (a, c);  $30\mu m$  (b, d, e).

- 8 The table reports the results of our quantitative analysis of split cell occurrence in 375x375 µm optical 9 sections (as those shown in a and c) from control and mycorrhizal plants. Two parameters were 10 considered: the average number of split cells per image (reported +/- the standard error) and the 11 number of images containing at least one couple of split cells. For L. japonicus, AM colonization 12 significantly increased the average number of split cells over 6 times (t test <0.01) as well as the 13 percent of images containing split cells (from 36% to 96%). An analogous situation was observed in 14 *M. truncatula*, where the average number of split cells per images increased over 4 times (T-test; 15 pvalue <0.01) and the percent of split cell-containing images passed from 36% in controls to 66% in 16 mycorrhizal roots. Overall these analyses correlate the presence of AM colonization with a consistent 17 increase in the number of split cortical cells.
- 18

19 Fig. 3. AtTPLATE-GFP highlights AM-induced cell divisions in the inner cortex. All panels 20 show longitudinally oriented optical sections cutting through the inner cortex from living M. 21 truncatula (a,b) or D. carota (c-h) ROCs colonized by G. gigantea. In the presence of a hyphopodium 22 (hp) and penetrating hyphae (hy) in the epidermis (a,b), several cell divisions are visible in the M. 23 truncatula inner cortex, marked by the intense TPLATE-GFP fluorescence at anticlinal walls 24 (arrowheads); the faint and diffuse GFP fluorescence in the cytoplasm and on the plasma membrane 25 was obtained by increasing the microscope sensitivity to provide a general outline of root cell shapes; 26 asterisks mark neighboring undivided cortical cells. (c) shows an analogous situation in carrot, where 27 bright TPLATE-GFP labeling (arrowheads) highlights recent cell divisions in the inner cortex 28 underlying a hyphopodium; also in this case, the asterisk marks an undivided cortical cell. (d) 29 Following arbuscule development, the compact shape of several divided cells is recognizable 30 (arrowheads), but the lack of intense anticlinal wall labeling by TPLATE-GFP suggests that cell 31 divisions have not occurred in the preceding hours; one cell (arrow) appears to have divided twice, 32 as suggested by the extremely reduced size of two neighboring cells. Serial optical sections recorded 33 at increasing depth from the same carrot root are displayed in (e,f,g). A large hyphopodium is visible 34 on the root epidermis in (e). Several intracellular hyphae extend from the hyphopodium to the outer 1 cortex (f), where particularly intense accumulation of GFP signal is associated to a cortical PPA 2 (arrowheads). (g) Fluorescence of comparable intensity marks a forming cell wall (arrowhead) in the 3 inner cortex; white dashed lines mark long, undivided cortical cells. (h) displays the same focal plane 4 as (g) 24 hours later; the previously partial cell wall is now completed (arrowhead) and all the cells 5 marked by the dashed line have divided. Bars =  $50\mu m$ .

6

**Fig. 4.** *At***TPLATE-GFP localization in** *M. truncatula dmi3-1* **mutant ROCs.** Recent cell divisions in the root tip meristem were marked by intense TPLATE-GFP accumulation (arrowheads), as shown in (a). In contrast to our observations on WT ROCs, the presence of a *G. gigantea* hyphopodium (hp) on the surface of a *dmi3-1* root (b) did not induce ectopic cell divisions in the underlying inner cortex ((c), asterisk). An analogous example is shown in (d,e) where two optical sections from the same root area are presented, one focused on the root epidermis (d) and one on the inner root cortex (e). Bars =  $20 \mu m$  in (a);  $40\mu m$  in (b-e).

14

15 Fig. 5. Regulation of cell division and endocytic markers in wild type and *dmi3-1 M. truncatula* 16 upon AM colonization. The expression of the cell divisions markers KNOLLE and CYC1, alongside 17 TPLATE and TPLATE interactors in clathrin mediated endocytosis AP2A1, CHC1 and CHC2, were 18 analyzed by quantitative RT-PCR 48 hours post hyphopodium development on ROC segments. 19 TPLATE and AP2A1 transcripts were significantly upregulated in wild type inoculated root segments 20 (striped histograms) compared to controls (white histograms); no significant upregulation was 21 observed in *dmi3-1* mutants, where symbiotic signaling and intraradical colonization are blocked. A 22 differential regulation was observed for the two Clathrin heavy chain genes: while no change in gene 23 expression was observed for CHC1, a weak but statistically significant upregulation of CHC2 was 24 recorded in inoculated wild type roots. Relative gene expression levels (fold change) are normalized 25 to the corresponding control; bars represent standard errors. Different letters indicate statistically 26 significant differences; T-test p value < 0.04; at least three biological replicates were used for each 27 condition.

28

# 29 Table S1. Primers used in this work

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Table S2. AM colonization causes the appearance of split cells in the inner root cortex. Two parameters were considered for quantitative analysis of split cell occurrence in 375x375 µm optical sections: the average number of split cells per image (reported +/- the standard error) and the number of images containing at least one couple of split cells. A marked increase in the number of split cells was observed upon fungal colonization in both *D. carota* and wild type *M. truncatula* (T-test; p value <0.01). By contrast, when the same analysis was done on *M. truncatula dmi3-1* mutants, no
 significant change was observed.

- 3
- 4 Figure S1. *Lotus japonicus TPLATE* promoter sequence used for GUS analysis.
- 5

6 Figure S2. TPLATE expression pattern in roots of L. japonicus expressing the pLjTPLATE:GUS 7 **construct.** (a) Intense blue GUS staining in a root tip meristem (asterisk). While GUS activity often 8 extended to the whole meristem, as shown in (b), alternative expression patterns were occasionally observed (c, d), likely related to differences in meristematic activity: as shown here, TPLATE 9 10 promoter activity was limited to small groups of meristematic cells (arrowheads). (e) Higher magnification of a root tip meristem, showing two pairs of GUS-positive meristematic cells; these 11 12 observations suggest that *pTPLATE* is upregulated in actively dividing cells. (f) GUS staining in the 13 central cylinder (cc) was much more prominent than in the other tissues of the differentiated zone, 14 where hardly any GUS staining could be detected during the reaction period, indicating much lower 15 expression levels than in the central cylinder. (g) 42 days post-inoculation with the AM fungus 16 Funnelliformis mosseae, GUS activity was evident in the colonized areas of the root cortex, where 17 arbuscules (ar) develop. (h, i) Transverse sections from control (h) and mycorrhizal (i) roots showing 18 GUS-staining in the central cylinder and arbuscule-containing cells. Bars =  $200 \mu m$  in (a, f, g); 100 19 µm in (b, c, d); 30 µm in (e); 50 µm in (h, i).

20

21 Figure S3. AtTPLATE-GFP localization in uninfected M. truncatula ROCs. New-laid cell walls 22 are highlighted in the root apical meristem (a,b) by an intense GFP signal (arrowheads). The nuclei 23 (n) of four recently-divided meristematic cells in (a) are recognizable as dark areas in the faintly 24 fluorescent cytoplasm and allow an easy identification of relative cell positions. A deep longitudinal 25 section from an uninfected root is shown in (c); root cortex layers are recognizable between the 26 epidermis (ep) and central cylinder (cc): the lack of intense labeling in any anticlinal wall and the 27 regularity of cell sizes indicate that no recent cell division has occurred. Bars =  $20 \,\mu m$  in (a) and (b); 28 50µm in (c).

29

Figure S4. AtTPLATE-GFP-labeled cell divisions in colonized carrot ROCs. Longitudinal optical sections cutting through the inner root cortex. (a) Intense GFP fluorescence (arrowheads) labels a few anticlinal walls in the inner cortical layers underlying a hyphopodium (hp); such cell walls divide regular cells in two smaller daughter cells. (b) (c) Couples of split cells (dashed line) and AtTPLATE-

- GFP marked walls (arrowheads) are visible in the vicinity of an intraradical hypha (h) reaching the
  inner cortex. (d) No divided cells are recognizable in uninoculated roots. Bars = 50µm.
- 3

Figure S5. Validation of sample mycorrhizal status by *BCBP* expression analysis. The
statistically significant upregulation of *Blue copper-binding protein* (*BCBP*) gene - an acknowledged
plant marker for AM colonization - confirmed AM establishment in the samples used for gene
expression analyses. Relative gene expression levels (fold change) are normalized to the control.
Different letters indicate statistically significant differences; T-test p value < 0.04.</li>