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To cite this version:
Simone Belmondo, Valentina Fiorilli, Jacob Pérez-Tienda, Nuria Ferrol, Roland Marmeisse, et al.. A dipeptide transporter from the arbuscular mycorrhizal fungus Rhizophagus irregularis is upregulated in the intraradical phase. Frontiers in Plant Science, Frontiers, 2014, 5 (436), 10.3389/fpls.2014.00436 . hal-02487466

HAL Id: hal-02487466
https://hal-univ-lyon1.archives-ouvertes.fr/hal-02487466
Submitted on 27 May 2020

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A dipeptide transporter from the arbuscular mycorrhizal fungus *Rhizophagus irregularis* is upregulated in the intraradical phase

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Arbuscular mycorrhizal fungi (AMF), which form an ancient and widespread mutualistic symbiosis with plants, are a crucial but still enigmatic component of the plant micro biome. Nutrient exchange has probably been at the heart of the success of this plant-fungus interaction since the earliest days of plants on land. To characterize genes from the fungal partner involved in nutrient exchange, and presumably important for the functioning of the AM symbiosis, genome-wide transcriptomic data obtained from the AMF *Rhizophagus irregularis* were exploited. A gene sequence, showing amino acid sequence and transmembrane domains profile similar to members of the PTR2 family of fungal oligopeptide transporters, was identified and called RiPTR2. The functional properties of RiPTR2 were investigated by means of heterologous expression in *Saccharomyces cerevisiae* mutants defective in either one or both of its di/tripeptide transporter genes PTRA2 and DAL5. These assays showed that RiPTR2 can transport dipeptides such as Ala-Leu, Ala-Tyr or Tyr-Ala. From the gene expression analyses it seems that RiPTR2 responds to different environmental clues when the fungus grows inside the root and in the extraradical phase.

**Keywords:** arbuscular mycorrhizal fungi, organic nitrogen, dipeptide transporter, *Rhizophagus irregularis*, symbiosis

## INTRODUCTION

Nitrogen (N) is quantitatively the most important soil nutrient for plant growth and productivity. Plants acquire N not only as nitrate and ammonium that are converted to amino acids in the root or shoot tissues, but also as organic N forms (i.e., amino acids, peptides, and proteins) from the soil (Näsholm et al., 2009). Free amino acids generally only account for a small fraction of organic N pool of soil solution whereas peptide- and protein-bound amino acids may contribute to most of the soil organic N pool (Farrell et al., 2011; Hill et al., 2012; Warren, 2014).

Although the ecological significance of organic N uptake for plant N nutrition is still a matter of discussion, several lines of evidence suggest that plants inhabiting ecosystems characterized by slow N mineralization rates may, to a significant degree, rely on organic N forms for growth (Lipson and Näsholm, 2001).

To make the picture more complex, many plant species form intimate symbioses with mycorrhizal fungi, which constitute a large proportion of the microbial biomass in many soils and may play a crucial role in the N nutrition of their host plants (Girlanda et al., 2007; Fitter et al., 2011).

Mycorrhizal fungi are also known to use different N sources, depending on specific biochemical, physiological, and ecological features of the fungus involved (Girlanda et al., 2007).

Acquisition of organic N has traditionally been associated with ectomycorrhizal fungi, which are localized in the upper, organic matter-enriched soil horizons. And, indeed, some ectomycorrhizal fungi have been shown to possess proteolytic capabilities and organic N uptake systems (Wipf et al., 2002; Guidot et al., 2005; Benjdia et al., 2006; Shah et al., 2013). Arbuscular mycorrhizal (AM) fungi, which form relationships with the majority of plant families in most ecosystems, improve the mineral nutrition of their hosts via an efficient uptake of mineral nutrients from the soil. AM fungi could also be involved in the acquisition of organic and inorganic N (Cappellazzo et al., 2008; Lanfranco et al., 2011; Pérez-Tienda et al., 2012). Arbuscule-containing cells are thought to be the main site for such a nutrient exchange (Bonfante and Genre, 2010). Arbuscules are highly branched fungal structures that develop inside a living cortex cell: each fungal branch is surrounded by a proliferating plant plasma membrane called the periarbuscular membrane.

It has been shown that organic N uptake is greatly enhanced by AM colonization (Cliquet et al., 1997; Hawkins et al., 2000) and that AM symbiosis could both enhance the decomposition of N and increase N capture from organic patches (Hodge et al., 2001). The uptake of exogenously supplied Arg has also been observed in the extraradical mycelium (ERM) grown in *in vitro* conditions.
cultures (Govindarajulu et al., 2005; Fellbaum et al., 2012). Apart from the capability of taking up amino acids, there is increasing evidence that AM fungi could increase N capture from more complex organic material (Hodge et al., 2001; Leigh et al., 2009; Whiteside et al., 2013).

N uptake requires the activity of transporters that transfer the N compounds across cellular membranes. Regarding organic N, the only transporter so far described in AM fungi is GmosAAP1, an amino acid permease from Funneliformis mosseae (Cappelazzello et al., 2008) which may play a role in the first steps of amino acid uptake from the soil. No data are so far available for peptide transporters.

At least four distinct plasma membrane, proton-coupled peptide transport systems have been described in fungi, and in other eukaryotic organisms. Transporters belonging to PTR, DAL5 and FOT families transport di- and tripeptides, while those belonging to OPT family transport longer tetra- and pentapeptides (Hauser et al., 2001; Homann et al., 2005; Reuß and Morschhäuser, 2006; Perry et al., 1994). More recent studies showed the allantotriaedulosuccinate permease DAL5, contributes to the uptake of the di- and tripeptides that a ptr2Δ mutant continued to assimilate (Hommann et al., 2005; Cai et al., 2007).

Several PTR members have also been identified in plants and exhibit various functions (Dietrich et al., 2004; Karim et al., 2005; Komarova et al., 2008). For example, in Arabidospis thaliana AtPTR5 facilitates peptide transport into germinating and possibly maturing pollen, ovules, and seeds while AtPTR1 has a role in uptake of peptides by roots (Komarova et al., 2008).

In this study we performed the first functional characterization of a putative dipeptide transporter Ript2 from the AM fungus R. irregularis which was previously shown to be differentially expressed between mycorrhizal roots and ERM (Tisserant et al., 2012). We further addressed its regulation pattern in intra- and extraradical fungal structures challenged with different N sources in order to understand its potential role in the fungal nutrition and symbiotic interaction.

**MATERIALS AND METHODS**

**BIOLOGICAL MATERIALS, GROWTH CONDITIONS, AND TREATMENTS**

Rhizophagus irregularis (Syn. Glomus intraradices, DAOM 197198; Krüger et al., 2012) inoculum for seedlings and root organ cultures (ROCs) mycorrhization, was produced through in vitro monocentric cultures. These were established in bi-compartmental Petri dishes with a watertight plastic wall separating the root compartment (RC) from the hyphal compartment (HC) (Fortin et al., 2002). The RC was filled with 25 ml of solid M minimal medium and the HC with 25 ml of solid M medium lacking sugar (M-C). Cultures were started by placing an explant of Glomus intraradices transformed-chicory (Cichorium intybus) roots colonized with the AM fungus in the RC. Once the mycelium of R. irregularis had grown on the plastic wall and completely filled the HC compartment, the medium was dissolved with sterile 10 mM citrate buffer, pH 6.0. Spores were then collected and used for plant inoculation.

To obtain the ERM, when the fungus profusely colonized the HC, its content was removed, and the HC was filled with 15 ml liquid M-C medium containing either 3.2 mM (100% N) or 0.8 mM (25% N) KNO₃. The mycelium was allowed to colonize this medium over the subsequent 2 weeks. Petri dishes were examined regularly and roots were trimmed as required to prevent crossing into the HC.

For the dipeptide treatments, ERM was grown for 2 weeks in liquid M medium (100% N as KNO₃). At this point, the medium of the HC was replaced and replaced by fresh liquid M-C medium containing as nitrogen sources 3.2 mM nitrate, 3.2 mM nitrate and 10 mM Ala-Leu, 10 mM Ala-Leu or no N. ERM was harvested after 24 h.

In both experiments, ERM was collected with tweezers, rinsed with sterilized water, dried with sterilized filter paper, immediately frozen in liquid N and stored at −80°C until used.

To obtain seedlings colonized by R. irregularis, the Millipore sandwich method (Giovannetti et al., 1993) was used. Seeds of Medicago truncatula Gaertn cv Jemalong were first scarified using sandpaper P180–200, sterilized with 5% commercial bleach for 3 min and rinsed three times for 10 min with sterile distilled water. Germination was induced under sterile conditions in 0.6% agar/water, incubated for 5 days in the dark (25°C) and then exposed at the light for 4 days. Plants were watered with a modified Long-Ashton (LA) solution containing 3.2 µM Na₂HPO₄·12H₂O and 0.5 mM NaN₃ nitrate as P and N sources, respectively (Hewitt, 1966) and were grown in a growth chamber under 14 h light (24°C)/10 h dark (20°C) regime. Plants were harvested 60 days post-inoculation (dpi).

For the dipeptide treatment, in a first experiment, M. truncatula mycorrhizal roots were obtained in pot cultures watered with a LA solution containing 1 mM nitrate. Two months after inoculation, mycorrhizal roots were treated for 24 h in hydroponic conditions with a LA solution containing 10 mM Ala-Leu or 1 mM nitrate or no N. In a second experiment M. truncatula mycorrhizal roots were grown in the sandwich system as described above for 60 days with a modified LA solution containing 2.5 mM Ala-Leu and 0.25 mM nitrate as N sources.

For mycorrhizal plants, only portions of the root system showing extraradical fungal structures were collected under a stereo-microscope. The colonization level was assessed according to Trouvelot et al. (1986). For the molecular analyses, roots were immediately frozen in liquid nitrogen and stored at −80°C.

**SEQUENCE ANALYSES**

Fungal protein sequences homologous to PTR2 transporters were identified by BLASTp searches within the Mycosocsm database (Grigoriev et al., 2014). Prediction of trans-membrane domains was performed using TMHMM (http://www.cbs.dtu.dk/services/TMHMM-2.0/), SOSUI (http://harrier.nagahama-i-bio.ac.jp/sosu1/), HMMPred (http://www.enzim.hu/hmmpred/) and TMPred (http://www.ch.embnet.org/software/TMPRED_form.html) programs.

Amino acid sequences were aligned using MUSCLE (Edgar, 2004) and their phylogenetic links inferred by using the
Maximum Likelihood method based on the JTT matrix-based model (Jones et al., 1992) as implemented in MEGA5 (Tamura et al., 2011).

YEAST COMPLEMEMENTATION ASSAYS

The coding region of RpPTR2 was amplified from R. irregularis cDNAs by PCR using Phusion DNA-Polymerase (Finnzymes, Espoo, Finland) and the two following oligonucleotides containing a NotI restriction site: forward primer, 5′-TGACATTTGCGCGCATATAAGGAGACATTTCA-3′; reverse primer, 5′-ACTTCCAGGGCCCGCTGACTATTTC TTCGAGATTAA-3′. The PCR product was cloned into NotI sites downstream of the S. cerevisiae constitutive PGK1 promoter in the pFL61 E. coli–yeast shuttle vector (Minet et al., 1992). Recombinant (pRiPTR2) and empty vector (EV) were used to transform W303 (MATa ura3 can1–100) yeast mutants deleted of only one of its two known dipeptide transporter genes, PTR2 and DAL5 (Homann et al., 2005).

Yeast transformation was performed using standard protocols (Rose et al., 1990) and yeast transformants propagating either the EV or the pRiPTR2 recombinant vector were initially selected on a yeast nitrogen base minimal medium lacking uracil. Single colonies of transformants were grown over night in 5 ml of NH4Cl-containing yeast base carbon (YCB) liquid medium. Once the OD600nm reached 0.9, serial dilutions (1:10–1:100–1:1000–1:10,000) in sterile water were prepared and 5 ml plated on YCB solid medium containing H2O2 (negative control), NH4Cl (positive control), Ala-Tyr, Tyr-Ala or Ala-Leu (0.25 mM each). Plates were incubated at 30°C for 4 days and photographed.

NUCLEIC ACID EXTRACTION AND RT-qPCR ASSAYS

Total genomic DNA was extracted from R. irregularis extraradical structures and M. truncatula and C. intybus roots using the DNeasy Plant Mini Kit (Qiagen), according to the manufacturer’s instructions.

Total RNA was isolated from about 100 mg of seedling roots and 20 mg of ERM using the RNeasy Plant Mini Kit (Qiagen). Samples were treated with TURBO™ DNase (Ambion) according to the manufacturer’s instructions. The RNA samples were used for reverse transcription (RT)-PCR amplification.

The 5′- and 3′-UTR sequences of the R. irregularis dipeptide transporter genes M. truncatula and C. intybus were determined by 5′-RACE and 3′-RACE respectively (Pérez-Tienda et al., 2011). The coding sequences were identified by comparing the 5′- and 3′-RACE sequences with the R. irregularis genome sequence (Minet et al., 2011). The coding region of RpPTR2 was amplified from R. irregularis cDNAs by PCR using Phusion DNA-Polymerase (Finnzymes, Espoo, Finland) and the two following oligonucleotides containing a NotI restriction site: forward primer, 5′-TGACATTTGCGCGCATATAAGGAGACATTTCA-3′; reverse primer, 5′-ACTTCCAGGGCCCGCTGACTATTTC TTCGAGATTAA-3′. The PCR product was cloned into NotI sites downstream of the S. cerevisiae constitutive PGK1 promoter in the pFL61 E. coli–yeast shuttle vector (Minet et al., 1992). Recombinant (pRiPTR2) and empty vector (EV) were used to transform W303 (MATa ura3 can1–100) yeast mutants deleted of only one of its two known dipeptide transporter genes, PTR2 and DAL5 (Homann et al., 2005).

Yeast transformation was performed using standard protocols (Rose et al., 1990) and yeast transformants propagating either the EV or the pRiPTR2 recombinant vector were initially selected on a yeast nitrogen base minimal medium lacking uracil. Single colonies of transformants were grown overnight in 3 ml of NH4Cl-containing yeast base carbon (YCB) liquid medium. Once the OD600nm reached 0.9, serial dilutions (1:10–1:100–1:1000–1:10,000) in sterile water were prepared and 5 ml plated on YCB solid medium containing H2O2 (negative control), NH4Cl (positive control), Ala-Tyr, Tyr-Ala or Ala-Leu (0.25 mM each). Plates were incubated at 30°C for 4 days and photographed.

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qRT-PCR experiments were carried out in a final volume of 20 μl containing 10 μl of iQ™ Universal SYBR® Green Supermix (Bio-Rad), 1 μl of 3 μM specific primers, and about 20 ng of cDNA. Samples were run in the iCycler iQ apparatus (Bio-Rad) using the following program: 10 min pre-incubation at 95°C, followed by 40 cycles of 15 s at 95°C, and 1 min at 60°C. Each amplification was followed by melting curve analysis (60–94°C) with a heating rate of 0.5°C every 15 s. All reactions were performed with three technical replicates and only Ct values with a standard deviation that did not exceed 0.3 were considered. The comparative threshold cycle method (Rasmussen, 2001) was used to calculate relative expression levels using the plant MiTEF or the fungal RfTef as reference genes for plant and fungal genes, respectively. The analyses were performed on three independent biological replicates. Statistical tests were carried out through one-way analysis of variance (One-Way ANOVA) and Tukey’s post-hoc test, using a probability level of p < 0.05. All statistical analyses were performed using the PAST statistical package (version 2.16; Hammer et al., 2001).

SEMI-QUANTITATIVE RT-PCR ON LASER MICRODISSECTED CELLS

M. truncatula roots colonized by R. irregularis obtained with the millipore sandwich system, as described above, were cut into 5–10 mm-long pieces, treated with ethanol and glacial acetic acid (3:1) under vacuum for 30 min and then placed at 4°C overnight. Roots were subsequently dehydrated in a graded series of ethanol (50%–70%–90% in sterilized water and 100% twice) followed by Neoclear (twice) with each step on ice for 30 min. Neoclear was gradually replaced with paraffin (Paraplast Plus; Sigma-Aldrich, St. Louis) according to Pérez-Tienda et al. (2011). A Leica AS LMD system (Leica Microsystems, Inc.) was used to collect colonized cortical cells from paraffin root sections as described by Balestrini et al. (2007).

RNA was extracted from dissected cells using the PicoPure kit protocol (Arcturus Engineering). A DNAse treatment was performed using an RNA-free DNase Set (Qiagen) in a Pico Pure column, according to the manufacturer’s instructions and RNAs were eluted in 21 μl of sterile water.

All RT-PCR assays were carried out using the One Step RT-PCR kit (Qiagen). DNA contaminations were assessed using the MiTEF primers described above. Reactions with RpPTR2 or MtPTR4 specific primers were carried out in a final volume of 10 μl containing 2 μl of 5 × buffer, 0.4 μl of 10 mM dNTPs, 1 μl of each primer 10 μM, 0.2 μl of One Step RT-PCR enzyme mix, and 1 μl of a total RNA diluted 1:1. The samples were incubated for 30 min at 50°C, followed by 15 min incubation at 95°C. Amplification reactions were run for 40 cycles of 94°C for 30 s, 60°C for 30 s, and 72°C for 40 s. RT-PCR experiments were conducted on two different biological replicates of 1500–2000 microdissected cells each.

RESULTS

With the aim to characterize fungal genes involved in fungal/plant nutrient exchange and, possibly, in the functioning of arbuscules, the key structures of the AM symbiosis, we
exploited transcriptomics data (Tisserant et al., 2012) generated for the AM fungus *R. irregularis* (Syn. *G. intraradices*; Krüger et al., 2012). We focused our attention on an EST (contig step3_c3279 of MIRA v2 assembly) that, from the microarray data, showed a 33-fold up-regulation in the intraradical mycelium (IRM) compared to the ERM (Tisserant et al., 2012). The sequence, containing a full length ORF of 547 amino acids, had been annotated as an oligopeptide transporter. BLAST searches indicated a similarity to fungal Major Facilitator Superfamily (MFS) di/tripeptide transporters. In particular, the sequence possesses the PTR2[ pfam00854] conserved domain that characterizes the proton-dependent oligopeptide transporters (PTR, T.C.2.A.17) family. For this reason, it was called RiPTR2.

Recent releases of the complete genome sequence of *R. irregularis* (Tisserant et al., 2013; Lin et al., 2014) allowed us to assess that RiPTR2 is a single copy gene containing four introns (Figure S1). Twelve transmembrane domains were predicted for RiPTR2 amino acid sequence (Figures S1,S2) using HMMTOP, TMHMM, SOSUI, and TMPRED programs. This structure is similar to that of the well-characterized PTR2 sequence from *S. cerevisiae* which shows 40.8% identity at the amino acid level to RiPTR2.

Two putative homologs were found within an extensive transcripts dataset of another AM fungus, *Gigaspora margarita* (Salvioli et al., in preparation), which is phylogenetically distant from *R. irregularis*. The two deduced amino acid sequences, named GmarPTR2A and GmarPTR2B show 56.2 and 51.7% identity to RiPTR2, respectively.

A phylogenetic analysis was carried out by Maximum Likelihood method using PTR2 sequences from a number of fungi representative of distinct taxonomic groups retrieved from the MycoCosm (Grigoriev et al., 2014) genomic database (Figure 1; Table S1). RiPTR2 grouped with the two *G. margarita* sequences to form a well-supported sequence cluster clearly separated from sequences from Ascomycota, Basidiomycota and Zygomycota species.

**RiPTR2 ENCODES A FUNCTIONAL DIPEPTIDE TRANSPORTER**

To verify whether RiPTR2 encodes a functional peptide transporter, it was expressed in *S. cerevisiae* W303 mutant strains.

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**FIGURE 1 | Phylogenetic position of RiPTR2 protein sequence in comparison with homologous sequences from selected species representative of the major fungal phyla (Table S1).** The evolutionary history was inferred by using the Maximum Likelihood method based on the JTT matrix-based model as implemented in MEGA5. The percentage of trees higher than 75% in which the associated taxa clustered together is shown next to the branches. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site.
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deleted of either one or both of its two endogenous di/tripeptide transporter genes, PTR2 and DAL5. Both single (ptr2Δ or dal5Δ) or double (ptr2Δ dal5Δ) mutant strains are unable to use many dipeptides as N sources (Homann et al., 2005).

RiPTR2-expressing yeast transformants were plated onto selective media, containing one of three different dipeptides as N sources; Ala-Tyr, Tyr-Ala or Ala-Leu. Tyr-Ala is known to be a substrate for the yeast Ptr2p but not for Dal5p, while Ala-Leu is used by Dal5p but not by Ptr2p, and Ala-Tyr is neither used by Ptr2p nor by Dal5p in the W303 background (Homann et al., 2005). In addition, H2O and NH4+ were used as negative and positive control, respectively.

All transformants (empty or recombinant vector) were unable to grow in N-free medium (H2O) although a background growth could be observed probably due to residual traces of N compounds present in the agar. As expected, all transformants were able to grow in the NH4+-containing medium. Interestingly, ptr2Δ dal5Δ cells expressing RiPTR2 grew on Ala-Leu, Ala-Tyr and also Tyr-Ala, whereas no growth was observed for ptr2Δ dal5Δ cells transformed with the EV, indicating that RiPTR2 is able to transport these three dipeptides (Figure 2).

Ala-Leu and Ala-Tyr transport in the dal5Δ single mutant expressing RiPTR2 was also evident (Figure 2). Results obtained with ptr2Δ single mutant were difficult to interpret since this mutant, transformed with the EV, showed a high background growth on all three dipeptides. However, RiPTR2 transformants clearly displayed an increased growth on all three dipeptides.

Taken as a whole these results demonstrated that RiPTR2 codes for a functional dipeptide transporter.

RiPTR2 EXPRESSION PROFILES

We first compared the expression profile of RiPTR2 in intraradical (IRM) and extraradical (ERM) mycelia by quantitative RT-PCR from M. truncatula plants colonized by R. irregularis grown in 0.5 mM nitrate in the sandwich system. RNA extractions were performed on ERM and on roots fragments from which the ERM was carefully removed to produce the IRM sample. Samples were normalized with the RiTEF housekeeping gene. A strong RiPTR2 expression was observed in the IRM while RiPTR2 transcripts were less abundant, and almost barely detected, in the ERM (Figure 3A). A similar result was obtained considering mycorrhizal roots of C. intybus devoided of ERM, grown in the root compartment and ERM developed in the hyphal compartment of the same ROC (Figure 3B). These results thus confirmed the strong expression in the IRM as observed in the microarray data (Tisserant et al., 2012).

We also performed a time course experiment in the sandwich system to determine RiPTR2 transcript abundance at different times (7, 14, 28, and 60 days) post-inoculation (dpi) of

**FIGURE 3** Relative expression of RiPTR2 assessed by qRT-PCR in intraradical mycelium (IRM) and extraradical mycelium (ERM) from mycorrhizal roots of M. truncatula grown in the sandwich system (A) and mycorrhizal roots of C. intybus grown in monoaxenic culture (B). Data for each condition are presented as mean ± standard deviation. Different letters indicate statistically significant difference (p < 0.05, ANOVA).

**FIGURE 2** RiPTR2 complements growth defects of ptr2Δ, dal5Δ or ptr2Δ dal5Δ yeast transporter mutants on dipeptides. Serial dilutions (1:10–1:100–1:1,000–1:10,000 from left to right) of cultured strains transformed with RiPTR2 or the empty vector (EV) were plated on YCB solid medium containing H2O (negative control), NH4+ (positive control), 0.25 mM Ala-Tyr, Tyr-Ala or Ala-Leu. Plates were incubated at 30°C for 4 days.
M. truncatula plants. Morphological analyses of roots showed almost no fungal structures at 7 or 14 dpi, while mycorrhization frequency increased from 28 to 60 dpi. Arbuscules were visible starting from 28 dpi and were slightly less abundant at 60 dpi than at 28 dpi (Figure S3). Since in the ERM RipTR2 is expressed at negligible levels, gene expression was evaluated in whole mycorrhizal roots, without making a distinction between IRM and ERM. RipTR2 mRNA abundance increased in parallel to the development of the intraradical phase as demonstrated by morphological data and the parallel mRNA accumulation of MtPT4, the M. truncatula phosphate transporter-encoding gene which is considered a molecular marker of arbuscule-containing cells (Harrison et al., 2002; Figures 4A,B).

The laser microdissection technique was used to specifically obtain RNA from M. truncatula arbusculated cells. The authenticity of the samples was verified using MtPT4 specific primers. RipTR2 mRNA was detected in the two independent samples analyzed, indicating that, under these conditions, in planta RipTR2 expression occurred in arbuscules (Figure 4C).

We then investigated whether the presence of a dipeptide (Ala-Leu), a candidate substrate of RipTR2 as indicated by the yeast heterologous expression, could modulate RipTR2 expression levels. This was tested in both a short- and in a long-term exposure experiments where we evaluated the RipTR2 expression in whole mycorrhizal roots, without making a distinction between IRM and ERM.

In the first experiment (short term exposure), M. truncatula mycorrhizal plants were obtained in pot cultures watered with a Long Ashton (LA) nutrient solution containing 1 mM nitrate. After 2 months mycorrhizal roots were treated for 24 h in hydroponic conditions with a LA solution containing 10 mM Ala-Leu or 1 mM nitrate or no N. No significant difference was observed in RipTR2 expression levels among the different treatments (Figure 5A). Plants showed rather similar mycorrhization degrees as revealed by the MtPT4 expression levels; although no N samples had higher values, they were not statistically different from those of the other two treatments (Figure 5B).

We also investigated whether the RipTR2 expression in the ERM developed in the ROC system was responsive to 24 h exposure to 10 mM Ala-Leu. As previously observed RipTR2
expression in the ERM was extremely low (Figure 3) and this treatment led to an apparent decrease of RiPTR2 mRNA abundance compared to the ERM kept for 24 h in a medium containing 3.2 mM nitrate or no nitrogen (Figure 6).

In the second experiment (long term exposure), *M. truncatula* mycorrhizal and non-inoculated plants were grown in the sandwich system for 60 days with a nutrient solution containing 2.5 mM Ala-Leu and 0.25 mM nitrate as N sources. Mycorrhizal plants grown in standard nutrient solution, that is 0.5 mM nitrate, were used as controls. Independently from the AM colonization, plants grown in the presence of Ala-Leu + nitrate have total (shoot + root) biomasses lower than plants grown in only nitrate (Figure 7A). Interestingly, when shoot to root ratio is considered, it appears that the mycorrhization has a positive effect on the plant in the presence of Ala-Leu + nitrate, in particular in resource allocation into shoots (Figure 7B). This result suggests that the AM fungal colonization contributes to the more efficient use of N, including dipeptides. This result may also be due to the higher colonization level as revealed by the MtPT4 expression (Figure 8B). No difference in RiPTR2 transcript abundance was observed in the two conditions (Figure 8A).

**DISCUSSION**

Organic N dominates the N pool of the majority of soils (Schimel and Bennet, 2004). A key concept under revision is that soil microbes have a greater ability to use organic N than plants: indeed there is evidence that plants not only take up organic N (Komarova et al., 2008) but may also compete with microbes for it, although it remains debated how competitive plants are (Kuzyakov and Xu, 2013).

Recent works have shown that, in soils, short peptides represent a greater proportion of N than free amino acids, although there is no detailed information on the availability of individual peptides due to their multiplicity and the lack of a suitable detection system (Hill et al., 2012). Among the peptide uptake systems, the PTR family comprises proton driven symporters which are present in bacteria, fungi, plants and animals. Human PTR transporters have been extensively studied as they are the main route through which the body absorbs and retain dietary protein (Newstead, 2014). Despite their potential importance in the N cycle (Schmidt et al., 2014), still very little is known on PTR from plants or soil fungi.

In this work we have characterized RiPTR2, a PTR transporter from the fungus *R. irregularis*, a model species for the ecologically and agriculturally important group of the Glomeromycota. Searches within the two complete genome sequences of *R. irregularis* so far available (Tisserant et al., 2013; Lin et al., 2014) showed that RiPTR2 is a single copy gene. Interestingly, two putative homologs have been identified within a transcriptome of another AM fungus, *G. margarita*, which belongs to a distinct family of Glomeromycota. PTR2 gene redundancy has also been found in the genomes of several filamentous fungi (Vizcaino et al., 2006), including the ectomycorrhizal fungus *Hebeloma cylindrosporum* which is, to our knowledge, the only mycorrhizal fungus with characterized PTR2 genes (Benjdia et al., 2006).

The RiPTR2 sequence was able to complement the growth defects of yeast mutants defective of the two well studied dipeptide transporters. At least in the heterologous system, RiPTR2 was able to transport Ala-Leu, Ala-Tyr and Tyr-Ala but it is likely that other dipeptides can be transported. Substrate promiscuity is in fact a common feature shared by all peptide uptake systems although the molecular basis of this phenomenon is still an open question (Newstead, 2014). On the other hand, this low substrate specificity has the advantage to avoid restricting uptake to only a subset of the available peptides.

**FIGURE 6** | Relative expression of RiPTR2 assessed by qRT-PCR in the extraradical mycelium developed in ROC exposed for 24 h to 10 mM Ala-Leu or 3.2 mM nitrate or no N. Data for each condition are presented as mean ± standard deviation. Different letters indicate statistically significant difference (p < 0.05, ANOVA).

**FIGURE 7** | Total fresh weight (A) and shoot to root ratio (B) of *M. truncatula* plants grown in the sandwich system inoculated (Myc) or not (C) and watered with either 2.5 mM Ala-Leu and 0.25 mM nitrate (Ala-Leu + nitrate) or 0.5 mM nitrate (LA). Data for each condition are presented as mean ± standard deviation. Different letters indicate statistically significant difference (p < 0.05, ANOVA).
Gene expression data clearly indicated that RiPTR2 is expressed in the ERM, though at a low level, suggesting a potential role in the uptake of dipeptides from the soil solution. However, at least in the analyzed conditions, mRNA abundance was higher in the intraradical phase. The time course experiment clearly showed an up-regulation of RiPTR2 during the colonization process and the establishment of mature mycorrhizas. The analysis of laser-microdissected arbusculated cells confirmed that RiPTR2 is consistently expressed in arbuscules. Since RiPTR2 mRNA abundance does not perfectly match MtPT4 mRNA accumulation (at 28 or 60 dpi) we suggest that RiPTR2 may be expressed not only in arbuscules but also in intercellular hyphae. Evidence that nutrient exchange goes beyond the interface of arbusculated cells and could involve intercellular hyphae as well has been obtained from the expression profile of a fungal monosaccharide transporter (Helber et al., 2011).

A high expression level in the intraradical phase, including arbuscules, where the flux is commonly expected to be directed toward the host plant cells, is puzzling. However, it is worth noticing that additional transporters, a phosphate one (Balestrini et al., 2007; Tisserant et al., 2012; Fiorilli et al., 2013) and an ammonium one (Pérez-Tienda et al., 2011) from the fungal symbiont were found to be consistently expressed in arbuscules. Dealing with organic N, R. irregularis transcriptomic data also showed the up-regulation of a putative amino acid transporter (contig Glomus_c20826) in the IRM vs. the ERM (Tisserant et al., 2012). These findings suggest that the fungus may reabsorb nutrients released in the periarbuscular space and thus exerts a control over the amount of nutrients delivered to the host. The interface compartment that surrounds the arbuscules is considered acidic (Guttenberger, 2000) and provides a gradient in the electrochemical potential used for the energization of uptake processes. It could be an ideal environment for PTR2 proteins, since they work as proton-coupled transporters and usually show the highest transport activity at acidic pH (Benjdia et al., 2006; Komarova et al., 2008). Interestingly, two M. truncatula PTR genes were also described as mycorrhiza-responsive and their transcripts detected in arbusculated cells (Gomez et al., 2009). We confirmed the up-regulation of these two M. truncatula genes in mycorrhizal roots compared to control roots in the long term experiment (data not shown).

RiPTR2 gene expression is responsive to the Ala-Leu dipeptide, a possible substrate. We observed a down-regulation of RiPTR2 in the ERM grown in ROC after 24 h exposure to 10 mM Ala-Leu. A down-regulation was also found for a dipeptide transporter of the ectomycorrhizal fungus H. cylindrosporum after exposure to millimolar dipeptide concentrations (Benjdia et al., 2006). By contrast, RiPTR2 expression in whole mycorrhizal roots did not change after 24 h exposure to 10 mM Ala-Leu. Since these expression values are likely to be mainly representative of the intraradical phase, we can argue that 24 h are not sufficient to perceive the dipeptide, and that the intraradical RiPTR2 expression is probably also regulated by the plant endogeneous N status which may not be affected by the short term exposure.

Interestingly, our data from the long term experiment suggest that the AM fungal colonization contributes to the more efficient use of N when nitrate is present in limiting condition and together with a dipeptide.

A part from yeasts (S. cerevisiae, Schizosaccharomyces pombe) where PTR2 expression is induced by either the presence of amino acids or dipeptides (Kitamura et al., 2012; Ljungdahl and Daignan-Fornier, 2012), very little is known on the regulation of PTR genes in filamentous fungi, with the exception of H. cylindrosporum (Benjdia et al., 2006). A sequence showing high similarity to PTR genes was recently found to be expressed during growth on protein-containing substrates in the ectomycorrhizal fungus P. involutus (Shah et al., 2013). A PTR2 gene from the biocontrol agent Trichoderma harzianum was up-regulated when the fungus interacted with the plant pathogen Botrytis cinerea (Vizcaíno et al., 2006) and during growth on cell wall of Fusarium solani (Vieira et al., 2013) indicating that PTR genes are involved in the mycoparasitic process. Furthermore, since Glomeromycota are very distantly related to commonly studied fungal species, we may anticipate that regulation of their PTR genes can respond to different environmental clues.
In their whole our results show that RiPTR2 is expressed in the ERM, the fungus–soil interface, suggesting a role in the uptake of organic N from soil; however, a stronger expression is consistently observed in the in planta phase, including arbuscules, pointing to a function in the mobilization of organic N in mycorrhizal roots. Further investigations, not only on the fungal but also on the plant side, will help to obtain a more comprehensive view of the dipeptide metabolism in the AM symbiosis.

AUTHOR CONTRIBUTIONS
Valentina Fiorilli and Simone Belmondo carried out the majority of the experiments and drafted the manuscript. Nuria Ferrol and Jacob Pérez-Tienda contributed to the experiments on the ERM grown in monoaxenic cultures. Roland Marmeiss participated in the design of the work and helped with the yeast complementation. Luisa Lanfranco coordinated the project and wrote the manuscript. All authors read and approved the final manuscript.

ACKNOWLEDGMENTS
The Authors express their thanks to Cristina Calcaro for her help in the microdissection experiments and Ascensión Valderas for technical assistance. Research was funded by the University grant (60%) to Luisa Lanfranco and the Project AGL2012-35611 from the Spanish Ministry of Economy and Competitiveness to Nuria Ferrol. We thank P. Bonfante for access to an unpublished genome dataset of Gigaspora margarita (BEG 34). We are grateful to F. Martin and the Mycorrhizal Genomics Initiative consortium for access to unpublished genome data. We also thank S. Torres and collaborators for the use of unpublished genome sequence data from Mucor circinelloides, M. Berbee and collaborators for those of Gonapodya prolifera and Comidiobolus coronatus, and J. Stajich, A. Grygansky, R. Vilgalys and collaborators for those of Umbelopsis ramanniana and Lichtheimia hyalospora. The genome sequence data were produced by the US Department of Energy Joint Genome Institute in collaboration with the user community.

SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at: http://www.frontiersin.org/10.3389/fpls.2014.00436/abstract

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Received: 27 June 2014; accepted: 15 August 2014; published online: 03 September 2014.

Citation: Belmondo S, Fiorilli V, Pérez-Tienda J, Ferrol N, Marmeisse R and Lanfranco L (2014) A dipeptide transporter from the arbuscular mycorrhizal fungus *Rhizophagus irregularis* is upregulated in the intraradical phase. *Front. Plant Sci.* 5:436. doi: 10.3389/fpls.2014.00436

This article was submitted to Plant Traffic and Transport, a section of the journal *Frontiers in Plant Science*.

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