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The SlZRT1 Gene Encodes a Plasma Membrane-Located ZIP (Zrt-, Irt-Like Protein) Transporter in the Ectomycorrhizal Fungus Suillus luteus

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Zinc (Zn) is an essential micronutrient but may become toxic when present in excess. In Zn-contaminated environments, trees can be protected from Zn toxicity by their root-associated micro-organisms, in particular ectomycorrhizal fungi. The mechanisms of cellular Zn homeostasis in ectomycorrhizal fungi and their contribution to the host tree’s Zn status are however not yet fully understood. The aim of this study was to identify and characterize transporters involved in Zn uptake in the ectomycorrhizal fungus Suillus luteus, a cosmopolitan pine mycobiont. Zn uptake in fungi is known to be predominantly governed by members of the ZIP (Zrt/IrtT-like protein) family of Zn transporters. Four ZIP transporter encoding genes were identified in the S. luteus genome. By in silico and phylogenetic analysis, one of these proteins, SlZRT1, was predicted to be a plasma membrane located Zn importer. Heterologous expression in yeast confirmed the predicted function and localization of the protein. A gene expression analysis via RT-qPCR was performed in S. luteus to establish whether SlZRT1 expression is affected by external Zn concentrations. SlZRT1 transcripts accumulated almost immediately, though transiently upon growth in the absence of Zn. Exposure to elevated concentrations of Zn resulted in a significant reduction of SlZRT1 transcripts within the first hour after initiation of the exposure. Altogether, the data support a role as cellular Zn importer for SlZRT1 and indicate a key role in cellular Zn uptake of S. luteus. Further research is needed to understand the eventual contribution of SlZRT1 to the Zn status of the host plant.

Keywords: Suillus luteus, mycorrhiza, zinc transporter, zinc homeostasis, zinc deficiency, metal uptake

INTRODUCTION

Zinc (Zn) is an essential micronutrient required by all living organisms (Eide, 2006). Due to its unique set of chemical properties, Zn plays both a functional and a structural role in many proteins. The metal is characterized by a small radius and has thus a highly concentrated charge (Clemens, 2006a). Furthermore, Zn is a Lewis acid with a high affinity for ligands with sulfur- (S),
Zn concentrations are included in the ZIP I subfamily (Gaither and Eide, 2001). However, the same properties that make Zn an indispensable nutrient can also induce toxicity (Clemens, 2006b). When present in excess, Zn can cause protein damage and inactivation by uncontrolled high-affinity binding to functional groups within proteins (Clemens, 2006b). For this reason, it is necessary for living cells to tightly regulate Zn concentrations in the cytoplasm. The physiological concentration range of Zn between deficiency and toxicity is extremely narrow and organisms are consequently equipped with a number of homeostatic mechanisms to tightly regulate cytoplasmic Zn concentrations (Eide, 2006). Especially transporter proteins play a crucial role in maintaining Zn homeostasis (Eide, 2006).

In eukaryotes most of the Zn transport is achieved by two protein families: the ZIP (Zrt/Irt-like protein) and CDF (Cation Diffusion Facilitator) transporter families (Gaither and Eide, 2001). Proteins belonging to the ZIP transporter family increase cytoplasmic Zn levels by transporting Zn across the plasma membrane or by mobilizing stored Zn from intracellular compartments. Whereas members of the CDF family transport Zn in the direction opposite to that of the ZIP proteins. Efflux or compartmentalization of Zn is promoted by transporting Zn from the cytoplasm into the lumen of an organelle or out of the cell (Gaither and Eide, 2001).

Transporters belonging to the ZIP family typically possess 5 to 8 transmembrane domains (TMDs). The protein sequence is most conserved in TMD IV and the region adjacent to TMD IV (Eng et al., 1998). The ZIP family can be divided into four subfamilies based on a higher degree of sequence similarity: the ZIP I, ZIP II, GufA and LIV-I subfamily (Guerinot, 2000; Gaither and Eide, 2001). Members of the ZIP family are well-studied in Saccharomyces cerevisiae, which is an excellent fungal model cell (Gaither and Eide, 2001). Currently, four ZIP Zn transporter genes have been identified in baker’s yeast: ZRT1, ZRT2, ZRT3, and YKE4. The yeast ZRT1 gene was the first influx Zn transporter gene from any organism to be characterized at the molecular level (Zhao and Eide, 1996). The ZRT1 gene encodes a high-affinity Zn uptake system induced by Zn limitation, whereas the ZRT2 transporter corresponds to a low-affinity uptake system that is active in Zn repleted cells (Eide, 1996; Zhao and Eide, 1996). Zn uptake in yeast is predominantly governed by these two plasma membrane-located transporters (Eide, 1996; Zhao and Eide, 1996). Both transporters are included in the ZIP I subfamily (Gaither and Eide, 2001). The third characterized yeast ZIP protein, ZRT3, belongs to the GufA ZIP subfamily (Gaither and Eide, 2001). This transporter localizes to the vacuolar membrane and mobilizes Zn under Zn deficiency (MacDiarmid et al., 2000). Lastly, Kumárovsics et al. (2006) characterized YKE4, a bidirectional Zn transporter in the endoplasmic reticulum (ER) of S. cerevisiae, which regulates Zn concentrations in the ER and cytoplasm. YKE4 is a LIV-I subfamily transporter (Gaither and Eide, 2001).

Additional to the research in yeast, other ZIP transporters and mechanisms of Zn homeostasis in fungi are primarily characterized and studied in human fungal pathogens. Membrane Zn importers of the ZIP I subfamily have been shown to be crucial for the acquisition of Zn and the virulence of several human pathogenic fungi (Crawford and Wilson, 2015). This was observed in Candida albicans for the Zn transporter CaZRT1 (Cititu et al., 2012), in Cryptococcus neoformans for CnZIP1 (Do et al., 2016) and in Aspergillus fumigatus for AfZrtI (Amich et al., 2014). These transporters enable pathogenic fungi to overcome Zn deficiency within the Zn-limited host environment (Jung, 2015). Zn and fungal ZIP transporters are therefore considered to be key players in this kind of pathogenic host-microbe interactions.

In the current study we aim to identify plasma membrane localized Zn importers and their role in cellular Zn homeostasis in the ectomycorrhizal fungus Suillus luteus. Ectomycorrhizae are mutualistic host-microbe interactions between tree roots and ectomycorrhizal fungi. The mycobiont offers the tree a balanced nutrient supply in exchange for photosynthetic sugar (Martin et al., 2016). Although Zn is not expected to be a key regulator of ectomycorrhizal development nor to be extremely scarce at the symbiotic interface, availability of this element may have an impact on the fitness of both individual symbiotic partners and the mutualism in particular environments. Micronutrient deficiencies are rarely observed in natural forests but severe Zn deficiency in tree plantations has been reported previously (Thorn and Robertson, 1987; Boardman and McGuire, 1990). Moreover, trees are sensitive to high soil Zn concentrations. We previously demonstrated that well-adapted ectomycorrhizal fungi can protect host trees from Zn toxicity when Zn is present in excess (Adriaensen et al., 2004, 2006). An improved knowledge on the mechanisms of cellular Zn homeostasis in ectomycorrhizal fungi, going beyond the general focus on detoxification by vacuolar sequestration and including Zn uptake and deficiency, will be the first step toward a better understanding of the contribution of ectomycorrhizal fungi to host tree Zn homeostasis.

MATERIALS AND METHODS

Fungal Strains and Culture Conditions
The monokaryotic S. luteus isolate UH-Slu-Lm8-n1 (Kohler et al., 2015) and the dikaryotic isolate UH-Slu-P4 (Colpaert et al., 2004) were used in this study. Cultures were maintained on solid Fries medium according to Colpaert et al. (2004). Preceding Zn exposure assays, 1-week-old exponentially growing mycelia were harvested and liquid cultures were initiated and maintained according to Nguyen et al. (2017). Three gram of spherical mycelia grown for 1 week in liquid culture were transferred to petri dishes containing 25 ml modified liquid Fries medium supplemented with 0, 20, 500, or 1000 μM ZnSO4·7H2O. These Zn concentrations were chosen to induce Zn deficiency, Zn...
sufficiency and mild Zn toxicity (Ruytinx et al., 2017). The petri dishes were incubated on a shaking incubator at 23°C. Metal exposure was performed in triplicate. Mycelia (400 mg) were sampled at 0, 1, 2, 4, 8, and 24 h after initiation of exposure, flash frozen in liquid nitrogen and stored at −70°C.

ZIP Identification and Phylogenetic Tree Construction

The S. luteus reference genome was searched for ZIP transporter encoding genes. A BLASTp search using characterized fungal ZIP transporters (Supplementary Table 1) and a Pfam domain search were performed at the S. luteus genome portal at MycoCosm of the Joint Genome Institute (JGI)1 (Grigoriev et al., 2012; Kohler et al., 2015). Full-length amino acid sequences of previously characterized ZIP transporters were obtained from the transporter classification database2, the Swissprot database3 and the National Center for Biotechnology Information (NCBI) server4. All sequences, including the newly identified S. luteus ZIP sequences, were aligned with the Multiple Alignment using Fast Fourier Transform (MAFFT) alignment logarithm version 7 (Katoh and Standley, 2013) and imported into the Molecular Evolutionary Genetics Analysis (MEGA) package version 6.06 (Tamura et al., 2013). A phylogenetic tree was constructed using the neighbor-joining (NJ) method (Poisson correction model for distance computation) to infer evolution of the identified S. luteus ZIP transporters and to predict their function more precisely.

Cloning of SIZRT1

Total RNA was extracted from S. luteus mycelium ground in liquid nitrogen using the RNeasy Plant Mini kit (Qiagen, Germany) and a cDNA library was constructed using the SMARTer cDNA synthesis kit (Clontech, United States) according to the manufacturer’s instructions. Specific primers were designed for amplification of the full-length coding sequence of SIZRT1 (left: 5’ CCT CAAACTATGTCAAGTTAAATT 3’; right: 5’ TGCCCA ACGCCCCAGGAGC 3’). The PCR reaction contained: 10x High Fidelity PCR buffer, 0.2 mM dNTP-mixture, 2 mM MgSO₄, 0.2 μM SIZRT1 forward and reverse primer, 5 ng cDNA and 0.5 U Platinum Taq High Fidelity DNA polymerase (Invitrogen, United States). RNase-free water was added to obtain a final reaction volume of 30 μl. The following PCR cycling conditions were used: 2 min at 95°C; 35 cycles of 30 s at 95°C + 30 s at 56°C + 1 min at 68°C, and 1 cycle of 3 min at 68°C. 5 μl of the PCR product was visualized on an agarose gel to verify the reaction specificity and the length of the amplicon. The remaining 25 μl PCR product was purified using the GeneJET PCR Purification Kit (ThermoScientific, United States). The purified PCR-product was cloned into the gateway entry vector pCR8/GW/TOPO (Invitrogen) and subsequently transferred by the Gateway LR-clonase II Enzyme Mix (Invitrogen) to destination vectors pYES-DEST52 (Invitrogen, United States) and pAG426GAL-ccdB-EGFP (Alberti et al., 2007) for functional analysis in yeast. The insert was sequenced in both directions to verify correct orientation and fusion.

Yeast Mutant Complementation and Subcellular Localization

SIZRT1 was heterologously expressed in S. cerevisiae. Yeast strains used are CM30 (MATα, ade6, can1-100, his3-11, 15 leu2-3, trp1-1, ura3-52) and CM34 or Δzrt1Δzrt2 (CM30, zrt1::LEU2, zrt2::HIS3) (MacDiarmid et al., 2000). Yeast cells were transformed according to the LiAc/PEG method as described by Gietz and Woods (2002). Transformed yeast cells were selected on synthetic defined medium without uracil [SD-URA; 0.7% w/v yeast nitrogen base (Difco), 2% w/v D-glucose, and 0.2% w/v Yeast Synthetic Drop-out Mix without uracil (Sigma)]. Plates were incubated at 30°C.

For metal tolerance assays, transformed yeasts were grown to mid log phase (OD₆₀₀ ± 1.5) in liquid SD-URA medium with 2% w/v D-galactose instead of D-glucose (induction medium). Yeast cells were pelleted, washed with sterile distilled water, and adjusted to OD₆₀₀ = 1. A 1/10 dilution series was prepared (10⁻⁰, 10⁻¹, 10⁻², and 10⁻³). Drop assays were performed for three independent yeast clones on SD-URA control induction medium (1 mM Zn) and induction medium supplemented with 50, 100, or 200 μM ethylenediaminetetraacetic acid (EDTA) (MacDiarmid et al., 2000). For subcellular localization of SIZRT1::EGFP fusion proteins, yeast transformants were grown to mid-log phase OD₆₀₀ = 1 on induction medium. Plasma membrane of the cells was stained at 0°C by FM4-64 (Molecular Probes, Invitrogen) according to Vida and Emr (1995). Afterward, a 3 μl droplet of yeast cells was analyzed at 0°C with a Zeiss LSM 510 META laser scanning confocal microscope (Carl Zeiss, Jena, Germany), using a Zeiss 40x NA1.1 water immersion objective (C-Apochromat 40x/1.1 W Corr., Carl Zeiss). Enhanced green fluorescent protein (EGFP) fluorescence analysis was performed with the 488 nm excitation line of an argon-ion laser and a band-pass 500–550 nm emission filter. FM4-64 (ThermoFisher) fluorescence analyses were performed with a 543 nm HeNe laser and a long-pass 560 nm emission filter. Image processing was carried out with ImageJ (NIH, Bethesda, MD, United States) software.

Zn Content Analysis of Transformed Yeast

Transformed yeast cells were cultured at 30°C in liquid induction medium without Zn until culture saturation. Three rounds of Zn deprivation were completed by re-inoculating 0.5 ml of saturated yeast suspension to new Zn-less induction medium. Zn-starved cells were grown to mid log phase (OD₆₀₀ ± 1.5) and diluted to OD₆₀₀ = 1. One ml of yeast suspension was added to Erlenmeyer flasks containing 20 ml liquid induction medium without Zn and medium supplemented with 500 μM Zn (repletion). Zn treatments were performed for five independent yeast clones. Cultures were allowed to grow for 24 h at 30°C. Yeast cells were collected by centrifugation, washed three times with 20 mM PbNO₃ and milli-Q water. Afterward cells were resuspended

1http://genome.jgi.doe.gov/Suilu2/Suilu2.home.html
2http://www.tcdb.org/
3http://www.uniprot.org/
4http://www.ncbi.nlm.nih.gov/
in 0.5 ml of milli-Q water, frozen (−20°C) and lyophilized. Lyophilized cells were acid digested (HNO3/HCl) and Zn content was determined by inductively coupled plasma optical emission spectrometry (ICP-OES).

**RNA Extraction, cDNA Synthesis, and qPCR**

Total RNA was extracted from mycelium pulverized in liquid nitrogen using the RNeasy Plant Mini Kit (Qiagen). The TURBO DNA-free kit (Invitrogen (Life Technologies), United States) was used to perform a desoxyribonuclease (DNase) treatment to remove residual genomic DNA. RNA integrity and absence of DNA contamination was verified via agarose gel analysis. RNA concentration and purity were evaluated on a NanoDrop ND-1000 Spectrophotometer (Isogen Life Science, the Netherlands). One µg of each RNA sample was converted to cDNA with the Primerscript RT Reagent Kit (Perfect Real Time) (TáKaRa Clontech, United States). A 10-fold dilution of the cDNA was prepared in 1/10 diluted Tris-EDTA (TE) buffer and stored at −20°C.

Real-time reverse transcription polymerase chain reaction (qRT-PCR) was performed in a 96-well optical plate with an ABI PRISM 7500 Fast Real-Time PCR System (Life Technologies) according to Ruytinx et al. (2016). A SIZRT1 specific primer pair was designed using Primer3 (Rozen and Skaletsky, 2000) (5′ GCCAAACGGACAAACTTG 3′; 5′ GACAGGCGCAGGATGAAAG 3′; efficiency = 92.1%). Data were expressed relative to the sample with the highest expression level via the formula 2−(Ct−Ctmin) and normalized using five reference genes (TUB1, ACT1, GR975621, AM085296, and AM085296). Reference genes were selected previously by Ruytinx et al. (2016) and their stability was confirmed within the current experimental conditions using GeNorm (Vandesompele et al., 2002). A normalization factor was calculated as the geometric mean of the relative expression levels of the reference genes. Mean values of four biological replicates were calculated, rescaled to the control condition (20 µM Zn) within each time point and log2 transformed. A two-way analysis of variance (ANOVA) followed by a Tukey’s HSD test was run in “R” version 3.2.2 (R Core Team, 2015) to assess differences in SIZRT1 expression level.

**RESULTS**

**Identification of a ZIP Transporter in**

**S. luteus**

The BLASTp search of the *S. luteus* genome (UH-Slu-Lm8n1 v2.0) with characterized fungal ZIP transporters identified four *S. luteus* genes (protein IDs 720881, 22926, 229544, and 811220) predicted to encode ZIP proteins. A NJ phylogenetic tree including previously characterized ZIP transporters was constructed to predict the function of the newly identified *S. luteus* genes. The different ZIP subfamilies are well-supported in the tree as indicated by high bootstrap values (>90; Figure 1). Three of the identified *S. luteus* proteins (IDs 720881, 22926, and 229544) cluster within the ZIP I subfamily. Within this subfamily, proteins with ID 720881 and 22926 cluster together with the *S. cerevisiae* Zn importers ScZRT1 and ScZRT2; the protein with ID 229544 clusters with *S. cerevisiae* ATX2, a Golgi transporter involved in manganese (Mn) homeostasis. The fourth identified *S. luteus* gene encodes a protein (ID 811220) clustering close to the *S. cerevisiae* YKE4 (ER localized Zn transporter) within the LIV-1 subfamily of ZIP transporters. We were not able to detect a member of the Gu H and ZIP II subfamily of ZIP transporters within the *S. luteus* reference genome.

One identified *S. luteus* gene, encoding the protein with ID 22926 was selected for further analysis because of its high sequence similarity with the yeast Zn importers ScZRT1 and SpZRT1. Reciprocal BLASTp suggest the *S. luteus* protein to be orthologous to the high-affinity Zn importers ScZRT1 and SpZRT1 and was therefore named SIZRT1. SIZRT1 is predicted to have a 1398 bp open reading frame with eight exons encoding a 338 amino acid polypeptide. The encoded peptide shows several characteristics that are typical for proteins belonging to the ZIP family (Figure 2). Eight TMDs were predicted by the topology program TMHMM and a long variable cytoplasmic loop is present between TMD3 and TMD4. A histidine rich motif HXX(HX)3, suggested to function as Zn binding site, is present in the variable cytoplasmic loop of SIZRT1 and two other histidines that are typically conserved in ZIP transporters were identified (Figure 2). One of these conserved histidines is located in the conserved TMD4, which contains the ZIP signature sequence described by Eng et al. (1998). SIZRT1 matches 13 of the 15 amino acids of this ZIP signature sequence.

**Functional Analysis of SIZRT1 in Yeast**

SIZRT1 was heterologous expressed in yeast to confirm that it encodes a plasma membrane-located ZIP Zn importer, which was predicted by the phylogenetic analysis. Figure 3 and Supplementary Figure 1 illustrate that transformation with SIZRT1 partly restored the growth of the zinc-uptake-deficient yeast strain Δzrt1Δzrt2 on medium supplemented with different concentrations of EDTA. Transformation with the empty vector did not result in complementation of the Zn deficient phenotype (Figure 3 and Supplementary Figure 1). Expression of the SIZRT1::EGFP fusion protein indicates a localization of SIZRT1 on the plasma membrane. Yeast cells transformed with SIZRT1::EGFP showed a bright green fluorescent ring surrounding the cells, which co-localized with FM4-64 plasma membrane staining (Figure 4).

**Zn and Fe Content Analysis of**

**Transformed Yeast**

Zn and Fe content were measured in Zn starved (Figure 5A) and Zn replete (Figure 5B) yeast cells in order to obtain more insight into the function of SIZRT1. Figure 5A illustrates that Δzrt1Δzrt2 yeast mutants transformed with SIZRT1 contained the same amount of Zn as the wild type (WT) yeast after starvation (0 µM Zn) while Δzrt1Δzrt2 mutants transformed with the empty vector had a significantly lower
Zn content. Similarly, 24 h after Zn repletion (500 µM Zn) SlZRT1 transformed yeast mutants and WT yeast accumulated significantly more Zn than empty vector transformed yeast mutants (Figure 5B). A small difference in Zn accumulation was observed between WT yeast cells and SlZRT1 transformed Δzrt1Δzrt2 yeast cells. Additionally, the Fe content in the yeast transformants was analyzed, since some ZIP transporters can also use Fe as a substrate. Yet, no significant differences in Fe content were observed among the yeast transformants exposed to 0 or 500 µM Zn (Supplementary Figures S2A,B).

**SlZRT1 Gene Expression Analysis in S. luteus**

In *S. luteus*, SlZRT1 gene expression was determined at early time points (0, 1, 2, 4, 8, and 24 h) after exposure to different concentrations of Zn [0, 20 (control), 500, and 1000 µM]
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FIGURE 2 | Alignment of SlZRT1 and ScZRT1 protein sequences. The eight transmembrane domains (TMDs) predicted by topology prediction program TMHMM2.0 are indicated in green, the putative metal binding domains in blue, the ZIP signature sequence as described by Eng et al. (1998) in red, and two conserved histidines with an asterisk (located in TMD4 and TMD5).

FIGURE 3 | Functional complementation of the zinc-uptake-deficient yeast strain Δzrt1Δzrt2 by SlZRT1. Wild type (WT) and mutant yeast cultures with an OD$_{600}$ = 1 were 10-fold serial diluted (10$^0$, 10$^{-1}$, 10$^{-2}$, and 10$^{-3}$) and spotted on control (1 mM Zn) or ethylenediaminetetraacetic acid (EDTA) supplemented synthetic drop-out (SD) medium. The WT strain was transformed with the empty vector (EV, pYES-DEST52; Invitrogen), the mutant strain Δzrt1Δzrt2 with either the EV or the vector containing SlZRT1. The experiment was carried out for three independent clones and pictures were taken after 4 days of growth.
to assess the role of the \textit{SIZRT1} in Zn homeostasis. Results clearly illustrate that mRNA levels of \textit{SIZRT1} are dependent of external Zn concentration (Figure 6). Exposure to mildly toxic Zn concentrations (500 and 1000 \textmu M) results in an almost immediate significant downregulation of \textit{SIZRT1} gene expression. The expression patterns upon exposure to 500 and 1000 \textmu M Zn are similar regardless differences in external Zn concentrations. In contrast, in the absence of external Zn, \textit{SIZRT1} expression is quickly induced to reach a maximum level after 2 h, declines to control levels after 4 h and tends to be higher again in the long term (24 h).

**DISCUSSION**

Transition metals, such as Zn, Fe, Mg, are essential to all living organisms. However, when present in excess these metals may become toxic. To overcome metal toxicity, it is crucial for cells to tightly control cytoplasmic metal concentrations (Eide et al., 2005). Metal transporter proteins play a crucial role in the regulation of cytoplasmic metal concentrations and cellular metal homeostasis (Migeon et al., 2010). Among fungi, mechanisms involved in Zn homeostasis are mostly studied in \textit{S. cerevisiae}. Transporters of the ZIP family were shown to be vital to prevent Zn deficiency in this species (Eide, 2006) and several other fungi (Kiranmayi et al., 2009; Jung, 2015). \textit{S. cerevisiae} possess two plasma membrane localized Zn importers of the ZIP family (ScZRT1 and ScZRT2) and one tonoplast localized ZIP transporter (ScZRT3) for re-mobilization of vacuolar stored Zn. In the current study, we identified four ZIP transporter encoding genes in the genome of the ectomycorrhizal fungus \textit{S. luteus}. Three of the newly identified proteins are members of the ZIP I subfamily of ZIP transporters, one belongs to the LIV-1 subfamily (Figure 1). With the exception of ScZRT3, a tonoplast transporter
involved in Zn mobilization from the vacuole, homologs for all characterized S. cerevisiae ZIP transporters were identified within the S. luteus genome. So far, no homologs for the ScZRT3 protein have been identified in members of the Basidiomycota. Nevertheless, several basidiomycetes including S. luteus store excess Zn into their vacuoles (Sackey et al., 2016; Ruytinx et al., 2017). Transporters belonging to other protein families likely evolved in these species to re-mobilize stored Zn in absence of external environmental Zn. In accordance with what has been found in other fungi, there was no member of the ZIP II subfamily of ZIP transporters detected in S. luteus. This subfamily consists mainly of metazoan representatives (Guerinot, 2000).

Reciprocal BLASTp suggested the S. luteus protein with ID 22926, named SIZRT1 to be orthologous to the S. cerevisiae ScZRT1 transporter. ScZRT1 functions as a high-affinity Zn uptake transporter and receptor (Schothorst et al., 2017). Together with its homolog, the plasma membrane transporter ScZRT2, ScZRT1 is responsible for Zn uptake in Zn deficient yeast cells (Gaither and Eide, 2001). SIZRT1 and ScZRT1 show 39% sequence identity. An important difference in the sequence of SIZRT1 and ScZRT1 is found within the putative Zn binding domain (histidine rich domain, HRD) localized within the cytoplasmic loop between TMD3 and TMD4 (Figure 2). SIZRT1’s binding domain (HDVHGHGHG) shows a HXX additional to the classical (HX)3 domain of ScZRT1 (HDHTHD). This difference might correspond to an altered affinity toward Zn and/or a modified function of the protein. Mutation of the histidines in the HRD of ScZRT1 results in a 70% reduction in the maximum uptake rate of ScZRT1 (V\text{max}), whereas the substrate concentration at which the reaction rate is half of V\text{max} (K\text{m}) remains unaffected (Gitan et al., 2003). Also for other ZIP1 subfamily transporters a reduction in Zn uptake due to mutation of histidines in the HRD was observed (Mao et al., 2007) and some of these histidines are even necessary for the protein to be functional, i.e., able to transport Zn across the plasma membrane (Milon et al., 2006).

Heterologous expression and subcellular localization in yeast are common experimental procedures to study eukaryotic gene function and protein localization (Zhao and Eide, 1996; Mokdad-Gargouri et al., 2012). Heterologous expression of SIZRT1 in the Δzrt1Δzrt2 yeast double mutant, which is defective in Zn uptake, resulted in an almost complete restoration of the phenotype (Figure 3) and SIZRT1::EGFP fusion proteins localize at the plasma membrane of yeast cells (Figure 4). These results support a role as plasma membrane localized Zn transporter for the SIZRT1 protein. However, kinetics of the transporter might be different from the ScZRT1 protein. SIZRT1 did not fully complement ScZRT1 as observed in the drop assays (Figure 3) and Zn starved SIZRT1 transformed Δzrt1Δzrt2 yeast cells accumulate less Zn within 24 h after Zn replenishment than WT yeast cells do (Figure 5). No significant differences in Fe content were observed (Supplementary Figure 2), indicating a high Zn specificity of the transporter.

In yeast, ScZRT1 expression is regulated both at the transcriptional and the post-transcriptional level by Zn (Gitan and Eide, 2000). Post-translationally, Zn induces the removal of ScZRT1 from the plasma membrane via ubiquitination (Gitan et al., 2003). After endocytosis the protein is degraded in the vacuole. This regulatory system ensures a rapid shutdown of Zn uptake in yeast cells exposed to high concentrations of Zn (Gitan and Eide, 2000). In S. luteus SIZRT1 expression is regulated by excess Zn. SIZRT1 expression level is significantly lower after exposure to potentially toxic concentrations of Zn (500 and 1000 µM) as compared to the control (20 µM) and this already 1 h after initiation of the exposure (Figure 6). In contrast, absence of external Zn results in a rapid accumulation of SIZRT1 mRNA. Two hours after initiation of Zn starvation in S. luteus mycelium, SIZRT1 gene expression peaks and declines again to reach control levels after 4 h of growth in absence of Zn. After 24 h of growth in the absence of Zn, the SIZRT1 expression level in S. luteus mycelium is slightly higher again compared to the level in mycelium grown in control conditions. These fluctuations in expression level could possibly reflect the cell’s Zn status. A similar expression pattern, though delayed in time was detected by Schothorst et al. (2017) in S. cerevisiae for ScZRT1.
in conditions of Zn deprivation. *ScZRT1* transcripts peak at 2 days under Zn deprivation and decline again afterward. A fast transcriptional response on limited environmental Zn concentrations is common for plasma membrane localized Zn transporters of the ZIP I subfamily. Induction of transcription in the absence of external Zn was reported previously for fungal ZIP I subfamily Zn importers which were identified in Ascomycota (*ScZRT1* of *Schizosaccharomyces pombe*, ZrfA, ZrfB and ZrfC of *Aspergillus fumigatus*, Tzn1 and Tzn2 of *Neurospora crassa*, CaZRT1 and CaZRT2 of *Candida albicans*) and in Basidiomycota (*CgZIP1* and *GgZIP2* of *Cryptococcus gattii*) (Dainty et al., 2008; Kiranmayi et al., 2009; Jung, 2015).

Altogether, our data support a function as plasma membrane localized Zn importer with an important role in Zn homeostasis of *S. luteus* for SIZRT1. Likely, SIZRT1 is responsible for an adequate supply of Zn to the cell when environmental Zn is limited. With our current data, we cannot conclude on a role for SIZRT1 as Zn receptor for signaling in order to adjust primary metabolism to external Zn availability. Such a role was reported recently for *ScZRT1* (Schothorst et al., 2017) and is certainly worth investigation in *S. luteus* and mycorrhizal fungi in general. Ectomycorrhizal fungi are well-known to offer their host plant a balanced nutrient supply by efficiently collecting limited nutrients and reducing the transfer of excess, potentially toxic elements. In relation to Zn, ectomycorrhizal fungi in general, and *S. luteus* in particular, are reported to protect their host plant from Zn toxicity (Colpaert et al., 2011). As trees in general do not tolerate high Zn soil concentrations, this protective feature of *S. luteus* is interesting for phytoremediation purposes. Further research is needed to better understand the regulation and function of SIZRT1 within the *S. luteus* - host ectomycorrhizal association and to assess the contribution of SIZRT1 to the Zn status of the host plant.

**AUTHOR CONTRIBUTIONS**

LC, JR, and JC designed the study. LC, AT, ES, and NA performed the experiments. LC, EM, AK, and JR analyzed the data. LC and JR wrote the manuscript. LC, ES, NA, MODB, JR, and JC contributed in manuscript editing. All authors read and approved the final version of the manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2017.02320/full#supplementary-material

**FIGURE S1** Functional complementation of the zinc-uptake-deficient yeast strain Δzrt1Δzrt2 by SIZRT1:EGFP. Wild type (WT) and mutant yeast cultures with an OD<sub>600nm</sub> = 1 were 10-fold serial diluted (10<sup>2</sup>, 10<sup>−1</sup>, 10<sup>−2</sup>, and 10<sup>−3</sup>) and spotted on control (1 mM Zn) or ethylenediaminetetraacetic acid (EDTA) supplemented synthetic drop-out (SD) medium. The WT strain was transformed with the empty vector (EV, pAG306GAL-ccdB-EGFP; Alberti et al., 2007), the mutant strain Δzrt1Δzrt2 with either the EV or the vector containing SIZRT1:GFP. The experiment was carried out for three independent clones and pictures were taken after 4 days of growth.

**FIGURE S2** Fe concentration in transformed yeast cells (A,B). The WT strain was transformed with the EV (pYES-DEST52, Invitrogen), the mutant strain with either the EV or the vector containing SIZRT1:GFP. Data are the average ± SE of five biological replicates, significant differences (p < 0.05) are indicated by different letters. (A) in control conditions (0 µM Zn), (B) after exposure to Zn (800 µM).
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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.