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Role of cytosolic, tyrosine-insensitive prephenate dehydrogenase in *Medicago truncatula*

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Abstract

L-Tyrosine (Tyr) is an aromatic amino acid synthesized de novo in plants and microbes downstream of the shikimate pathway. In plants, Tyr and a Tyr pathway intermediate, 4-hydroxyphenylpyruvate (HPP), are precursors to numerous specialized metabolites, which are crucial for plant and human health. Tyr is synthesized in the plastids by a TyrA family enzyme, arogenate dehydrogenase (ADH/TyrA_p), which is feedback inhibited by Tyr. Additionally, many legumes possess prephenate dehydrogenases (PDH/TyrA_c), which are insensitive to Tyr and localized to the cytosol. Yet the role of PDH enzymes in legumes is currently unknown. This study isolated and characterized *Tnt1*-transposon mutants of *MtPDH1* (*pdh1*) in *Medicago truncatula* to investigate PDH function. The *pdh1* mutants lacked *PDH* transcript and PDH activity, and displayed little aberrant morphological phenotypes under standard growth conditions, providing genetic evidence that *MtPDH1* is responsible for the PDH activity detected in *M. truncatula*. Though plant PDH enzymes and activity have been specifically found in legumes, nodule number and nitrogenase activity of *pdh1* mutants were not significantly reduced compared with wild-type (Wt) during symbiosis with nitrogen-fixing bacteria. Although Tyr levels were not significantly different between Wt and mutants under standard conditions, when carbon flux was increased by shikimate precursor feeding, mutants accumulated significantly less Tyr than Wt. These data suggest that *MtPDH1* is involved in Tyr biosynthesis when the shikimate pathway is stimulated and possibly linked to unidentified legume-specific specialized metabolism.

KEYWORDS

legumes, prephenate dehydrogenase, the shikimate pathway, TyrA dehydrogenase, tyrosine

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1 | INTRODUCTION

l-Tyrosine (Tyr) is an aromatic amino acid synthesized de novo in plants and microbes, but not animals; thus, humans must acquire Tyr through their diet or by enzymatic conversion of l-phenylalanine (Phe, Fitzpatrick, 1999). In addition to its involvement in protein synthesis, Tyr and a Tyr-pathway intermediate 4-hydroxyphenylpyruvate (HPP) are the precursors to numerous specialized metabolites crucial for plant and animal health (Schenck & Maeda, 2018). Tyr-derived plant specialized metabolites have roles as photosynthetic electron carriers (plastoquinone, Metz, Nixon, Rogner, Brudvig, & Diner, 1989), pollinator attractants (betalain pigments, Strack, Vogt, & Schliemann, 2003) and defense compounds (dhurrin and rosmarinic acid Møller, 2010; Petersen, 2013). In grasses, Tyr also serves as a precursor to lignin, the main phenolic polymer in plants (Barros et al., 2016; Higuchi, Ito, & Kawamura, 1967; Rosler, Krekel, Amrhein,

& Schmid, 1997). Humans have co-opted some of these natural products to serve nutritional and medicinal roles such as some benzylisoquinoline alkaloids, which have antitussive, analgesic, and antimicrobial activities (Barken, Geller, & Rogosnitzky, 2008; Beaudoin & Facchini, 2014; Kries & O'Connor, 2016), and the antioxidant properties of tocopherols (collectively referred to as vitamin E, Bramley et al., 2000).

The aromatic amino acids (AAAs; Tyr, Phe, and tryptophan [Trp]) are synthesized in the plastids from chorismate, the final product of the shikimate pathway (Maeda & Dudareva, 2012; Tzin & Galili, 2010). Chorismate mutase (CM, EC number 5.4.99.5) catalyzes the committed step of Tyr and Phe synthesis—the pericyclic Claisen rearrangement of chorismate into prephenate (Eberhard et al., 1996; Goers & Jensen, 1984; Kuroki & Conn, 1989; Mobley, Kunkel, & Keith, 1999). Prephenate is converted into Tyr via two reactions, oxidative decarboxylation catalyzed by a TyrA

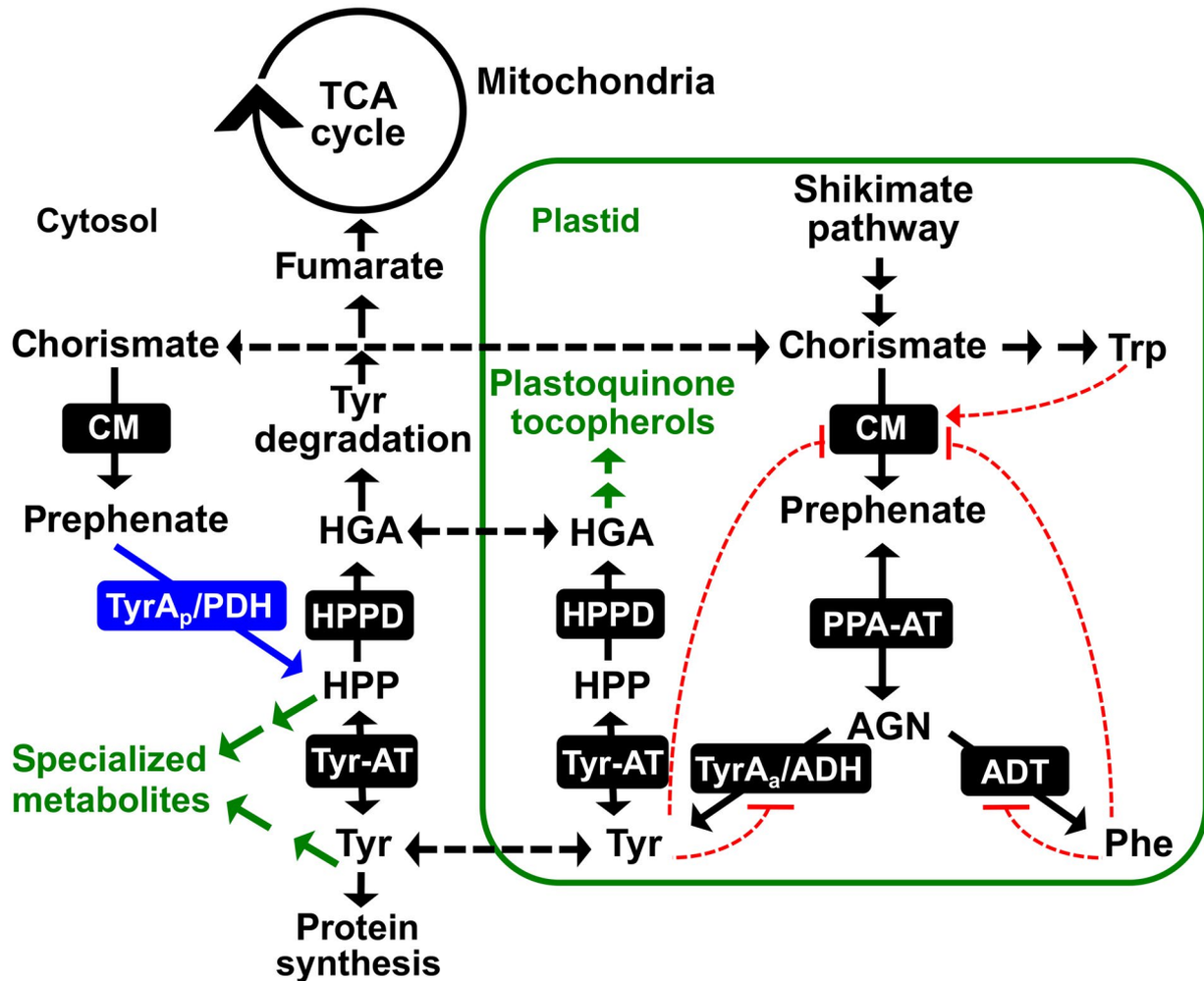


FIGURE 1 Tyr biosynthesis and associated pathways in legumes. In addition to a highly regulated plastid localized ADH pathway for Tyr biosynthesis, legumes possess a Tyr-insensitive, cytosolic PDH enzyme (blue). Tyr synthesized in the plastids is exported into the cytosol where it can be incorporated into proteins, enter the Tyr degradation pathway to the TCA cycle or serve as a precursor to specialized metabolism. Black dotted lines denote known or potential transport steps, and red dotted lines represent feedback regulation with arrows meaning induction and hashes inhibition. AGN, arogenate; ADT, arogenate dehydratase; CM, chorismate mutase; HGA, homogentisate; HPP, 4-hydroxyphenylpyruvate; HPPD, HPP dioxygenase; PPA-AT, prephenate aminotransferase; TyrA_a/ADH, arogenate dehydrogenase; TyrA_p/PDH, prephenate dehydrogenase; Tyr-AT, tyrosine aminotransferase.

dehydrogenase enzyme and transamination. These reactions can occur in either order, leading to alternative Tyr pathways (Figure 1, Schenck & Maeda, 2018). In most microbes, a prephenate-specific TyrA dehydrogenase (PDH/TyrA_p, EC 1.3.1.13, Figure 1) first converts prephenate into HPP followed by transamination to Tyr via Tyr-aminotransferase (Tyr-AT or TyrB, EC2.6.1.5). In plants, these reactions occur in the reverse order with transamination of prephenate to form aroenate by a plastidic prephenate aminotransferase (PPA-AT, EC 2.6.1.78), followed by oxidative decarboxylation catalyzed by a plastidic aroenate-specific TyrA dehydrogenase (ADH/TyrA_a, EC 1.3.1.78) producing Tyr (Figure 1). The ADH-mediated Tyr pathway appears to be essential for normal growth and development in plants as indicated by the severe phenotype of *Arabidopsis thaliana adh2/tyra2* knockout mutant, which was further exacerbated by transient suppression of the other *ADH1/TyrA1* gene (de Oliveira et al., 2019). PDH and ADH are the key enzymes in their respective pathways, as they compete for substrates that are shared with Phe biosynthesis, and are generally feedback inhibited by Tyr (Connelly & Conn, 1986; Gaines, Byng, Whitaker, & Jensen, 1982; Rippert & Matringe, 2002; Rubin & Jensen, 1979).

PDH activity, which is commonly found in microbes, has been detected in tissue extracts of some plants, all restricted to the legume family (Gamborg & Keeley, 1966; Rubin & Jensen, 1979; Siehl, 1999). In soybean, leaf tissue had the highest PDH activity of all analyzed tissues (Schenck, Chen, Siehl, & Maeda, 2015). Phylogenetic analyses of plant and microbial TyrAs showed that *PDH* genes are uniquely present in legumes, suggesting that legume PDH enzymes are the result of a recent legume-specific gene duplication event of a plant *ADH* rather than horizontal gene transfer from a *PDH*-possessing rhizobia (Schenck et al., 2015). However, not all legumes possess *PDH*, which suggests gene loss in some legumes (Schenck et al., 2015; Schenck, Holland, et al., 2017). *PDH* recombinant enzymes from *Glycine max* (soybean; GmPDH1 and GmPDH2) and *Medicago truncatula* (Medicago; MtPDH1) preferred prephenate over aroenate as their substrates (Schenck et al., 2015; Schenck, Holland, et al., 2017) and, unlike plant *ADH* enzymes, were insensitive to Tyr inhibition and localized to the cytosol (Schenck et al., 2015; Schenck, Holland, et al., 2017). These recently diverged plant *ADH* and *PDH* enzymes were used to identify a single amino acid residue (Asn222 of GmPDH1) of TyrA dehydrogenases that switches TyrA substrate specificity and underlies the evolution of legume *PDH* enzymes (Schenck, Holland, et al., 2017; Schenck, Men, & Maeda, 2017). While these biochemical and evolutionary studies established that some legumes have an alternative Tyr-insensitive cytosolic *PDH* enzyme(s), its *in vivo* function is unknown.

To address this issue, here we hypothesized and tested four non-mutually exclusive *in planta* functions of the *PDH* enzyme in legumes. Hypothesis I: *PDH* functions as a part of a redundant Tyr biosynthetic pathway in the cytosol (Figure 1). Although AAA biosynthesis is localized to the plastids (Bickel, Palme, & Schultz, 1978; Jung, Zamir, & Jensen, 1986), some plants including

legumes possess cytosolic isoforms of CM and Tyr-AT, which catalyze immediately up- and down-stream steps of *PDH* and complete the cytosolic Tyr biosynthetic pathway from chorismate (Figure 1) (D'Amato, Ganson, Gaines, & Jensen, 1984; Ding et al., 2007; Eberhard et al., 1996; Schenck et al., 2015; Wang, Toda, & Maeda, 2016). Also, a cytosolic Phe pathway was recently identified in plants (Qian et al., 2019; Yoo et al., 2013). Hypothesis II: *PDH* is involved in the production of Tyr or HPP-derived metabolite(s) (Figure 1). Tyr is the precursor of many plant specialized metabolites (Schenck & Maeda, 2018), and duplicated primary metabolic enzymes can be co-opted to efficiently provide Tyr and HPP precursors to support their downstream specialized metabolism (Maeda, 2019; Moghe & Last, 2015; Weng, Philippe, & Noel, 2012). Hypothesis III: *PDH* is involved in the Tyr catabolism pathway to the tricarboxylic acid (TCA) cycle. Tyr catabolism proceeds through HPP, the product of *PDH*, and feeds intermediates (e.g., fumarate) of the TCA cycle (Figure 1). Thus, HPP produced from *PDH* may be directly incorporated into the Tyr degradation pathway. Hypothesis IV: *PDH* is involved in the legume-rhizobia symbiosis. Many legumes form a symbiotic relationship with rhizobia, and *PDH* activity is uniquely present in legumes. Furthermore, *MtPDH1* expression is upregulated in the nodules after treatment with nitrate (NO₃⁻, Benedito et al., 2008) or phosphinothricin (PPT, Seabra, Pereira, Becker, & Carvalho, 2012), both of which stimulate nodule senescence (Matamoros et al., 1999; Pérez Guerra et al., 2010; Streeter & Wong, 1988). To systematically test the above four hypotheses, this study isolated and characterized *pdh* mutants using genetics, biochemistry, metabolomics, histochemical staining, microscopy, and gene expression analyses in the model plant *M. truncatula*, which conveniently has a single *PDH* gene compared with some other legumes that have multiple copies (Schenck et al., 2015; Schenck, Holland, et al., 2017).

2 | MATERIALS AND METHODS

2.1 | Plant materials and growth conditions

Medicago truncatula accession R108 wild type and *pdh1* mutant seeds were scarified in concentrated sulfuric acid for eight minutes and repeatedly washed with water. Seeds were then surface sterilized with bleach for 1.5 min followed by repeated water washes. Seeds were placed in sterile water for 16 hr at 4°C and then transferred to germination medium (0.5× MS medium, 0.8% agar, 1 μM GA₃, pH 7.6), wrapped in aluminum foil and placed at 4°C. After 48 hr, plates with sterilized seeds were moved to 22°C. After 24 hr, the aluminum foil was removed and the germinated seedlings were transferred to standard potting soil and placed in a growth chamber (Convion). Pots were watered with 1× Hoaglands solution when dry, and grown under 12-hr light:dark cycles with 200 μE light intensity and ~60% humidity.

For extended dark treatments, plants were grown under the above conditions for six weeks. Leaves were then excised and floated in

water and returned to the above growth conditions with dark-treated leaves being covered with a cardboard box. After the treatment, leaves were dried using a kimwipe and used for metabolite extraction.

2.2 | Genotyping

A single young leaf from 6-week-old plants was placed in 1.7-ml microcentrifuge tube with 600 μ l of DNA extraction buffer (10 mM Tris-HCl pH 8.0, 25 mM ethylenediaminetetraacetic acid (EDTA), and 0.5% SDS), pulverized with a mini plastic blue pestle, and incubated at 55°C for 15 min. The solution was cooled to room temperature and 200 μ l of 5 M ammonium acetate was added, vortexed for 20 s, and centrifuged at 14,000 g for 3 min. The supernatant was transferred to a fresh tube and 600 μ l of isopropanol was added, followed by centrifugation at 14,000 g for 1 min. The supernatant was decanted and the pellet washed with 400 μ l of 70% ethanol followed by centrifugation at 14,000 g for 1 min. The supernatant was decanted and the pellet dried for 1 hr in a sterile hood. The resulting DNA was dissolved in 50 μ l of H₂O. Genotyping PCRs (20 μ l total) contained 1 μ M gene and insertion-specific primers (Table S2), 1x EconoTaq PLUS master mix (Lucigen), and 1 μ l of genomic DNA. DNA was amplified in a thermocycler with the following conditions: an initial denaturation at 95°C for 5 min, 35 cycles of amplification at 95°C for 20 s, 60–65°C for 20 s, 72°C for 60 s, with a final extension at 72°C for 5 min.

2.3 | Quantitative reverse transcription PCR (qRT-PCR)

RNA was extracted from leaves of 6-week-old plants. About 50 mg of tissue was pulverized in liquid N₂ using a mini pestle. RNA extraction buffer (68 mM sodium citrate, 132 mM citric acid, 1mM EDTA and 2% SDS) was added and vortexed immediately for 10 s, and the tubes were then placed on their sides for 5 min at 22°C. The solution was centrifuged at 12,000 g for 2 min and 400 μ l of the supernatant was transferred to a fresh tube. 100 μ l of 1 M NaCl was added and mixed by pipetting, followed by addition of 300 μ l of chloroform, inverted multiple times and centrifuged at 4°C for 10 min at 12 000 g. The upper phase was transferred to a new tube and an equal volume of isopropanol was added, mixed by inverting and placed at 4°C for 10 min. The solution was then centrifuged at 4°C for 10 min at 12,000 g and the supernatant decanted. The pellet was washed with 70% ethanol, followed by centrifugation for 3 min at 12,000 g and the supernatant decanted and the pellet dried in a sterile hood until all residual ethanol was evaporated. The resulting pellet was redissolved in 25 μ l of nuclease-free H₂O (Promega). To remove DNA, the RNA solution was treated with DNase (Turbo DNase, Fisher) following the manufacturer's protocol. The remaining RNA was quantified using a nano-drop spectrophotometer (Thermo Scientific) and diluted to a 20 ng/ μ l concentration. RNA was converted into cDNA using reverse transcriptase (Applied biosystems) with an oligo d(T) primer.

For qPCR, cDNA was diluted to 5 ng/ μ l and additional fivefold dilutions were made to calculate primer efficiency. All primer pair efficiencies were between 90%–100%. cDNA was mixed with GoTaq qPCR master mix (Promega) containing SYBR green and 300 nM of each primer. Reactions were placed in an Stratagene Mx3000P (Agilent) thermocycler using the following PCR cycle an initial denaturation at 95°C for 10 min, 45 cycles of amplification at 95°C for 15 s, 60°C for 30 s, 72°C for 30 s.

For relative quantification, Ct values were extracted for each reaction and used to quantify initial cDNA concentration using 2^{- $\Delta\Delta$ Ct} method normalized to a housekeeping gene (*MtPI4K*, Kryvoruchko et al., 2016, Table S2) using the LinRegPCR program (Ruijter et al., 2013). For absolute quantification, pET28a vectors carrying *MtPDH1*, *MtncADH* and *MtADH* were used as the template to amplify a fragment of the corresponding genes. The resulting fragments were purified from 0.8% agarose gels using a QIAquick gel extraction kit (Qiagen) following manufacturer's protocol. The DNA concentration was quantified and repeated fivefold dilutions were made and used to obtain a standard curve in qRT-PCR with gene-specific primers (Table S2). Ct values were extracted for each qRT-PCR reaction and used to quantify initial cDNA concentration by using the linear range created as above for the respective gene.

2.4 | Enzyme extraction and ADH and PDH assays

Leaf tissue from 6-week-old plants were ground to a fine powder in a prechilled mortar and pestle under liquid N₂. Extraction buffer (25 mM HEPES, pH 7, 50 mM KCl, 10% ethylene glycol, 1% polyvinylpyrrolidone (PVP) and 1 mM dithiothreitol (DTT)) was added in a 1:3 ratio of tissue to buffer (w/v). The slurry was centrifuged for 20 min at 4°C, at 20,000 g and the resulting supernatant desalted using a gel filtration column (Sephadex G50-80 resin, Sigma-Aldrich) equilibrated with extraction buffer without DTT and PVP. Protein concentrations were determined by a Bradford assay (Bio-Rad Protein Assay, Bio-Rad).

The desalted crude enzyme extracts were used in ADH and PDH reactions that contained 25 mM HEPES (pH 7.5), 50 mM KCl, 10% ethylene glycol, 1 mM NADP⁺, 1 mM substrate (L-arogenate or prephenate, respectively). Arogenate was prepared by enzymatic conversion from prephenate (Sigma-Aldrich), as previously reported (Maeda, Yoo, & Dudareva, 2011). Reactions were initiated by addition of enzyme from various sources and incubated at 37°C for 45 min. The resulting assays were injected into HPLC equipped with a ZORBAX SB-C18 column (Agilent) to directly detect the final product of the assay as described previously (Schenck et al., 2015).

For PDH and ADH activities from various legumes (Figure S9), leaf material was obtained from the University of Wisconsin-Madison Botany Department greenhouse or from identified trees on campus. Plants were grown under varying light and temperature conditions and leaf material was collected at different developmental stages. Enzyme extractions and ADH and PDH assays were performed as described above.

2.5 | Metabolite extraction and detection

Tissue from 6-week-old plants was added into 1.7-ml microcentrifuge tubes with 400 μ L of extraction buffer (2:1 (v/v) methanol:chloroform, 0.01% butylated hydroxytoluene (BHT), 100 μ M norvaline and 1.25 μ g/ml tocol as previously described (Collakova & DellaPenna, 2003), with 3 glass beads (3 mm). Samples were vigorously shaken for 3 min at 1,000 rpm using a genogrinder (MiniG 1600, SPEX SamplePrep). Additional chloroform (125 μ l) and water (300 μ l) were added and vortexed for 30 s. Samples were centrifuged for 10 min at 20,000 rpm and the polar and non-polar phases were transferred to new tubes and dried using a vacuum concentrator (Labconco).

For tocopherol analysis, the dried nonpolar phase was resuspended in methanol with 0.01% BHT. Samples were injected into a HPLC (Agilent 1260) equipped with a ZORBAX SB-C18 column (Agilent) using a 30-min isocratic elution of 95% methanol, 5% water. Tocopherols in the extractions were detected using fluorescence excitation at 290 nm and emission at 330 nm and compared with authentic standards (Sigma-Aldrich) and normalized to an internal control (tocol).

For anthocyanin detection, 1 M HCl of methanol was added to the polar phase in a 1:1 ratio. A spectrophotometer was used to measure absorbance at 520 nm. Absolute levels were estimated using an extinction coefficient of anthocyanin absorbance (Lee et al., 2005) of $33,000 \text{ L} \times \text{M}^{-1} \times \text{cm}^{-1}$.

The polar and non-polar phases were analyzed with GC-MS. The dried polar phase was redissolved in pyridine with 15.0 mg/ml methoxyamine-HCl. Samples were vortexed for 30 s followed by sonication for 10 min and incubated for 60 min at 60°C. This was repeated once more, then the samples were derivatized with an equal volume of *N*-methyl-*N*-(tert-butyldimethylsilyl)trifluoroacetamide + 0.1% *tert*-butyldimethylchlorosilane (MTBSTFA + 0.01% t-BDMCS, Sigma-Aldrich) and incubated for 60 min at 60°C. Samples were then injected into GC-MS (Trace 1310, ISQ LT, Thermo Scientific). The dried non-polar phase was redissolved in 800 μ l of chloroform with 50 ppm BHT and 500 μ l of 1.25 M HCl in methanol. Following incubation at 50°C for 4 hr, samples were completely dried under nitrogen gas. The dried non-polar phase was resuspended in 70 μ l of pyridine and derivatized with 30 μ l of MTBSTFA + 0.01% t-BDMCS (Sigma-Aldrich). Following transfer to a glass vial, 1 μ l of the polar and non-polar phases were injected onto a 30 m column (TG-5MS, Thermo Scientific) using a 10:1 split ratio, and an oven ramp method of 5°C per minute for 46 min and held at 300°C for 10 min. Detected compounds were compared with NIST library matches and peak areas based on ion abundance were normalized to an internal standard (norvaline).

2.6 | Acetylene reduction assay (ARA)

Plants used for ARA were scarified and germinated as described previously. Seedlings were transferred to 12" square plates containing modified solid Fahræus medium supplemented with 0.5 mM NH_4NO_3 (Catoira et al., 2000). Plates were grown under the same conditions as pot grown plants; however, plates were placed at a

$\sim 60^\circ$ angle so that the roots would not penetrate the agar. After 5 days of growth on plates, each plant was inoculated with 1 ml of water containing *Sinorhizobium meliloti* Rm1021 at an OD_{600} 0.02. Plants were then allowed to grow for 14, 21 and 28 dpi at which point nodules were counted and plants were moved to 10-mL glass jars with 1 ml of sterile water at the bottom. Glass jars were sealed with a rubber stopper, then injected with 1 ml of acetylene gas (10% acetylene final concentration). After 48 hr of incubation at 37°C, 1 ml of the gas from the glass jars was injected into a gas chromatograph (GC-2010, Shimadzu) to measure the production of ethylene and ARA activity was calculated as described in Hardy, Holsten, Jackson, & Burns, 1968.

2.7 | Histochemical staining of lignin composition

For cell wall composition staining, thin cross sections of stem tissue from 6-week-old plants were prepared using a razor blade. For Mäule staining, stem sections were placed in a 1% potassium permanganate solution for 5 min, then rinsed with water followed by addition of a 12% HCl (v/v) for 5 min and rinsed with water. A 1.5% solution of sodium bicarbonate was added to facilitate a color change to dark red and visualized using an epifluorescence microscope (Olympus BX60).

For phloroglucinol staining, similar stem cross sections were placed in a well plate with 1 ml of 10% phloroglucinol (w/v) solution in 95% ethanol with 500 μ l of 10 N HCl and incubated for 5 min. Stem sections were transferred to a glass slide and washed with water and visualized using an epifluorescence microscope (Olympus BX60).

GUS staining was performed on nodules developed after inoculation with *S. meliloti* Rm1021 carrying a *PnifH::Uida* fusion, to localize expression of the bacterial *nifH* gene, which is required for nitrogen fixation. Nodules were embedded in 4% agarose and 50–100 μ M sections were made with a vibratome® 1000 plus (Leica). Sections were immersed in a staining solution (2.5% 5-Bromo-4-chloro-3-indoxyl-beta-D-glucuronic (X-Gluc), 0.2 M sodium phosphate buffer (pH 7), 0.1 M potassium ferricyanide, 0.1 M potassium ferrocyanide, 0.25 M Na_2EDTA , and 10% Triton X-100) and vacuum infiltrated for 10 min, incubated in the dark at 37°C for 30 min, and rinsed with phosphate buffer. Sections were visualized using bright field microscopy.

3 | RESULTS

3.1 | Isolation of *pdh* mutants in *Medicago truncatula*

To investigate the in vivo function of PDH in legumes, two independent homozygous alleles of *Tnt1*-transposon mutants of *M. truncatula* (Tadege et al., 2008) were identified for the *MtPDH1* locus (*Mt3g071980*) (Cheng et al., 2014). The *pdh1-1* and *pdh1-2* mutants carried a transposon insertion in the first and last exons of the *MtPDH1* gene, respectively (Figure 2a). *MtPDH1* was constitutively

expressed across many tissues and under various conditions with the highest and lowest expression being detected in aerial tissues and seeds, respectively (Figure S1). Both alleles did not produce any *PDH* transcript in the leaves (Figure 2b). The *M. truncatula* genome contains two *ADH* genes, *MtADH* (Mt4g115980) and *MtncADH* (Mt5g083530) (Schenck, Holland, et al., 2017), but neither were enhanced in the *pdh1* mutants (Figure 2b). Consistent with their transcript levels, PDH activity was almost completely abolished in both *pdh1* mutants without significant reduction in ADH activity (Figure 2c). After six weeks of growth, *pdh1-1* showed no phenotypic difference from wild-type (Wt; R108). The *pdh1-2* mutant, on the other hand, had a slight bushy phenotype, which could be due to an unknown secondary transposon insertion(s) (Figure 2d); however, multiple backcrossing attempts were unsuccessful. These data provide genetic evidence that *MtPDH1* is responsible for the PDH activity detected in *M. truncatula*. The minor impacts of eliminating PDH activity on overall plant growth in two independent *pdh1* mutants suggest that the PDH enzyme is not essential during standard growth conditions in *M. truncatula*.

3.2 | Tyr and Tyr-derived compounds are unaltered in *pdh* mutants during standard growth condition

To test the hypothesis that the PDH pathway serves as a redundant Tyr biosynthetic route (hypothesis I), Tyr and Tyr/HPP-derived compounds were analyzed in Wt and mutants. Metabolites were extracted from root and leaf tissue of 6-week-old plants grown under standard conditions. Surprisingly, the levels of Tyr were not

significantly different between Wt and mutants in either leaf or root tissue (Figure 3a). Additionally, the levels of Phe and Trp in the leaves and roots were not significantly reduced in mutants as compared to Wt, though a slightly higher Trp level was observed in *pdh1-2* (Figure 3a). Potential effects on tocopherols, HPP-derived metabolites, were also tested (hypothesis II), but their levels were not significantly different between Wt and mutants in both leaf and root tissue (Figure 3b). We also performed non-targeted analysis using GC-MS for both polar and non-polar metabolites; however, no consistent differences were observed between Wt and mutants (Table S1). Together, these data suggest that under standard growth conditions, the lack of the PDH enzyme had no substantial effects on the overall accumulation of AAAs or Tyr/HPP-derived metabolites analyzed.

In grasses, upwards of 50% of the total lignin is derived from Tyr, via Tyr ammonia-lyase (TAL) activity (Barros et al., 2016). Since TAL activity has also been detected in some non-grass species, including legumes (Beaudoin-Eagan & Thorpe, 1985; Giebel, 1973; Khan, Prithviraj, & Smith, 2003), the legume PDH enzyme may synthesize Tyr that is directly incorporated into the phenylpropanoid pathway for downstream products such as lignin (hypothesis II). To test this possibility, stem cross sections from Wt and mutants were stained using two different reagents, Mäule and phloroglucinol, which can detect potential differences in composition or linkages of lignin (Mitra & Loqué, 2014; Pomar, Merino, & Barceló, 2002). Neither staining method showed any obvious differences in stem lignification between Wt and mutants (Figure S2a,b). When phenylpropanoid intermediates, *p*-coumarate and ferulate, involved in lignin biosynthesis were analyzed by

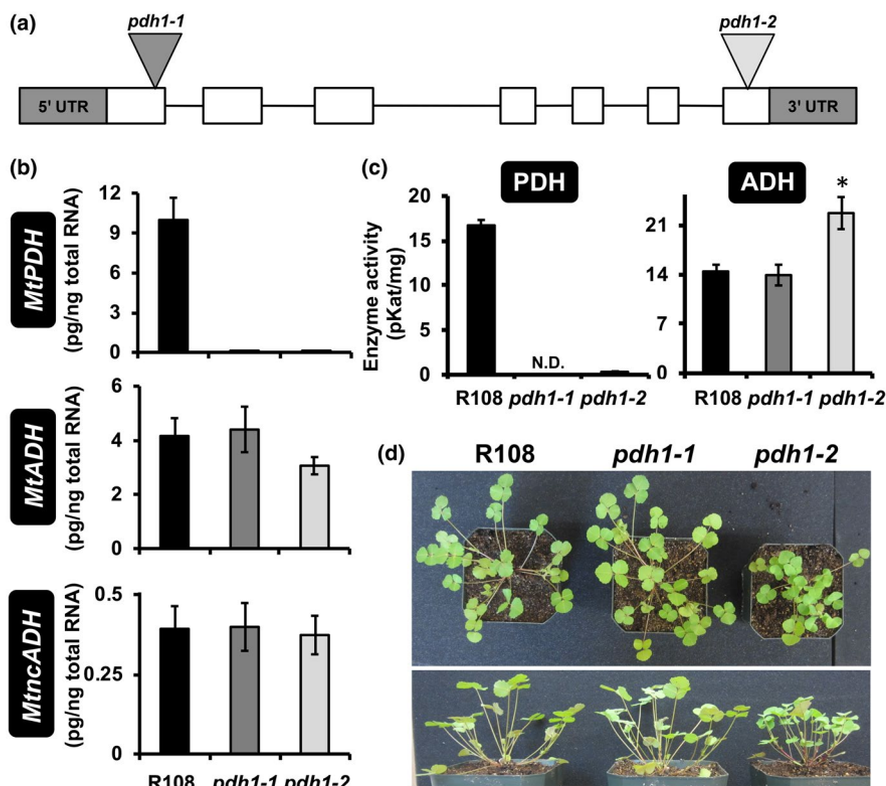
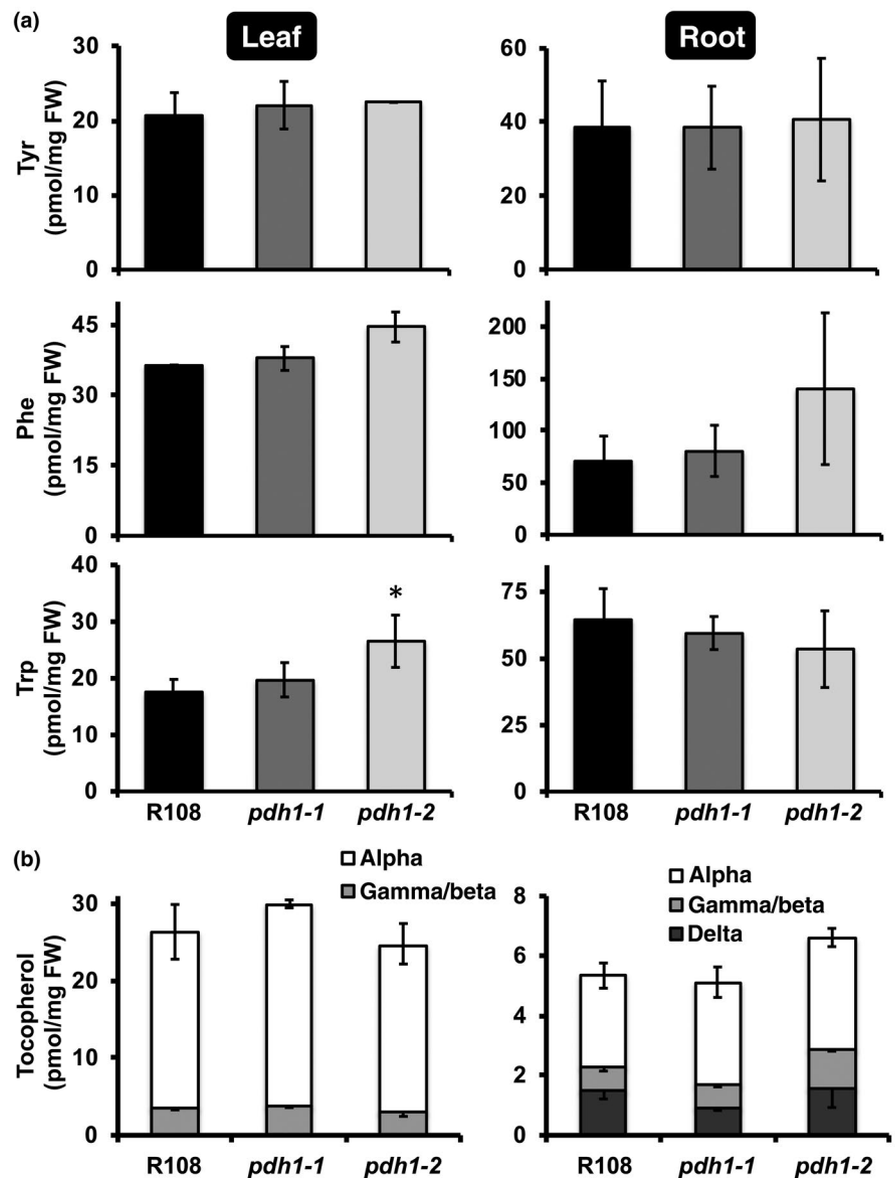


FIGURE 2 Isolation of *Medicago pdh1* mutants. (a) *MtPDH1* (Mt3g071980) genomic structure, exons (white boxes) and introns (lines), 5' and 3' untranslated regions (UTR, gray boxes). Two *Tnt1*-transposon mutants were isolated with insertions in exon one (*pdh1-1*) and seven (*pdh1-2*). (b) *MtPDH1* transcripts were nearly abolished in *pdh1-1* and *pdh1-2*, without affecting expression of either *ADH* homolog. Bars represent average absolute mRNA levels (pg/ng total RNA) \pm SEM of three biological replicates. (c) PDH and ADH activity from mutants and wild-type (Wt, R108). Bars represent average enzymatic activity (pKat/mg) \pm SEM of three biological replicates. Significant differences to R108 control are indicated; * $p \leq .05$. (d) Phenotype of R108 and mutants after 6-week growth under standard conditions

FIGURE 3 Targeted metabolite analysis of *pdh1* mutants and Wt. Leaf and root tissues from 6-week-old plants, grown under standard conditions, were used for metabolite extraction. Bars represent average absolute metabolite levels (pmol/mg fresh weight (FW)) \pm SEM of three biological replicates. (a) Aromatic amino acid levels in leaf and root tissues. Significant differences to Wt (R108) control are indicated; * $p \leq .05$. (b) Tocopherol composition and content from leaf and root tissue



GC-MS, they were somewhat reduced in *pdh1-2* but not consistently in *pdh1-1* (Figure S2c). Thus, the lack of PDH does not have substantial impacts on lignin biosynthesis in *M. truncatula*.

3.3 | Less tocopherols are accumulated in *pdh1* than Wt under high light treatment

Under various biotic and abiotic stresses, the shikimate pathway is induced, which often leads to accumulation of AAAs (Betz et al., 2009; Dyer, Henstrand, Handa, & Herrmann, 1989; Gilbert, Gadush, Wilson, & Madore, 1998; Zhao, Williams, & Last, 1998). To test the potential role of PDH in Tyr biosynthesis under stress (hypothesis I), Wt and *pdh1-1* were subjected to 48-hr high light treatment (Gonzali, Mazzucato, & Perata, 2009), which is known to induce production of AAA-derived antioxidants (Collakova & DellaPenna, 2003). The levels of Tyr were not altered, except at 24 hr when Tyr increased slightly in *pdh1-1* compared with Wt

(Figure S3a). As expected, high light treatment enhanced tocopherol accumulation, but to a significantly lesser extent in *pdh1-1* compared with Wt at both 24 and 48 hr (Figure S3b). The levels of anthocyanins, which are also induced under various stresses including high light stress (Collakova & DellaPenna, 2003), were increased after 24 and 48 hr of high light treatment but were not significantly different between Wt and *pdh1-1* (Figure S3c). These results show that the lack of PDH negatively impacts the accumulation of HPP-derived tocopherols when their production is induced under high light conditions.

3.4 | *MtPDH1* is co-expressed with senescence-related genes but has no major impacts on dark-induced senescence

To identify potential processes and pathways that are coordinately regulated with *PDH*, a gene co-expression analysis with

MtPDH1 was performed using the Medicago gene expression atlas (He et al., 2009). *MtPDH1* was co-expressed with genes mainly involved in senescence-related processes (e.g., nucleases, proteases, and lipases, Figure S1) and the gene encoding HPP dioxygenase (HPPD, Siehl et al., 2014), a senescence-activated enzyme involved in Tyr catabolism (Figure S5a) (Wang, Toda, Block, & Maeda, 2019). To experimentally test the potential involvement of PDH in senescence, *PDH* gene expression and enzymatic activity were monitored at different developmental stages during natural leaf senescence (Figure S4a). To define stages of leaf senescence, expression of a senescence marker gene (*MtVPE*, Pérez Guerra et al., 2010) was monitored, together with loss of chlorophyll in leaves collected at various developmental stages from a single plant (Figure S4b). *MtVPE* was basally expressed in fully green leaves (defined here as the S1 stage) and gradually induced upon senescence (defined here as S2, S3, and S4; Figure S4b), mirroring the loss of chlorophyll. Fully senescent leaves (S5) were collected but did not yield high-quality RNA and proteins, and hence could not be further analyzed. The highest *MtPDH1* expression was detected in green (S1) leaves, and PDH enzymatic activity was not induced upon senescence (Figure S4c,d). Thus, *MtPDH1* is not upregulated during natural senescence in the leaves. Similar to *PDH*, expression of the two *ADH* genes in *M. truncatula* did not follow the developmental pattern of leaf senescence (Figure S4c). Unlike PDH activity, however, ADH enzymatic activity was gradually induced upon senescence (Figure S4d).

Since *MtPDH1* was co-expressed with *MtHPPD* (Figure S1), and PDH together with HPPD provides a direct route for catabolism of Tyr to homogentisate, eventually leading to fumarate (Figure 1, Figure S5a), we investigated potential impacts of *pdh1* deficiency in Tyr catabolism (hypothesis III). The expression of genes encoding all enzymes of the Tyr catabolic pathway was measured in the mutants and Wt under standard growth conditions (Figure S5a, Dixon & Edwards, 2006). The expression of genes encoding the first two steps of the pathway, HPPD and homogentisate dioxygenase (*HGO*), were not significantly altered in the mutants compared with Wt. The subsequent step, maleylacetoacetate isomerase (*MAAI*), was induced by twofold and 2.5-fold in *pdh1-1* and *pdh1-2*, respectively (Figure S5b), though the final step in the pathway, fumarylacetoacetate hydrolase (*FAH*), showed opposite expression patterns in the two mutants (Figure S5b). Thus, no consistent changes in the expression of the Tyr degradation pathway genes, beyond *MAAI*, were observed in *pdh1* mutants.

To further examine Tyr catabolism during leaf senescence under artificial, but controlled, conditions excised leaves from Wt and mutants were exposed to an extended dark treatment (Peng et al., 2015; Wang et al., 2019). Over 7 days, leaves from Wt and mutants responded to dark-induced senescence similarly with no apparent growth phenotypes (Figure S6a). Furthermore, there were no significant differences in α -tocopherol levels between Wt and mutants at any time point analyzed (Figure S6b). These data together suggest that the lack of PDH had no substantial effects on Tyr catabolism or

metabolism under standard growth conditions and at least under the senescence conditions tested here (Figures S5 & S6).

3.5 | Less Tyr is accumulated in *pdh* mutants following shikimate feeding

Although the steady-state levels of Tyr were not altered in the mutants (Figure 3a), some of the above data suggest that the PDH enzyme may contribute to Tyr production (hypothesis I) at least under some stress conditions (Figures S1 and S3b). Thus, we hypothesized that the potential role of PDH in Tyr biosynthesis might be manifested when the carbon flux toward the Tyr pathway is elevated. To test this possibility, an intermediate of the upstream shikimate pathway, shikimate, was exogenously fed to Wt and mutants, and the levels of AAAs were analyzed. Excised leaves were floated in a solution containing shikimate or H₂O for up to 8 hr, rinsed to remove metabolites in the feeding solution, and leaf metabolites were extracted and analyzed using HPLC and GC-MS. As more prolonged feeding and higher concentrations of shikimate led to abnormal leaf phenotypes, 8 hr and 25 mM shikimate were selected as an optimal time and concentration for increased carbon flux without pleiotropic effects. After the feeding, shikimate and Phe levels were increased drastically in all genotypes, suggesting that shikimate was taken up and metabolized by the leaves (Figure 4). Tocopherol levels were generally unaffected upon shikimate feeding, likely because the feeding time was not long enough to convert shikimate into tocopherols (Figure 4). Trp levels were reduced in the mutants compared with Wt after feeding with water, and were induced upon shikimate feeding in *pdh1-1* (Figure 4). The unexpected differences observed with Trp levels under standard growth conditions (Figure 3) and the feeding experiments (Figure 4) may reflect unknown stress responses during the feeding experiments. Nevertheless, after 8 hr of feeding, Tyr levels increased by > 39-fold in Wt, but only by 16- and 13-fold in *pdh1-1* and *pdh1-2*, respectively (Figure 4). Repeated 8 hr shikimate feeding experiments with *pdh1* mutants and Wt yielded similar Tyr accumulation patterns. The difference in the Tyr accumulation suggests that PDH contributes to Tyr biosynthesis when carbon flow through the shikimate pathway is enhanced.

To determine if global metabolite changes occurred after shikimate feeding, additional amino acids and TCA pathway metabolites were analyzed by GC-MS. Interestingly, glutamine levels were higher in H₂O-fed Wt, compared with mutants (Figure S7), and upon shikimate feeding, glutamine levels were further significantly elevated in Wt (Figure S7). The levels of TCA cycle intermediates may indicate the functionality of the Tyr catabolic pathway; however, those TCA cycle intermediates analyzed here, including fumarate, were not consistently altered after shikimate feeding in Wt and mutants (Figure S8). Citrate levels were reduced in *pdh1-1* and *pdh1-2* compared with Wt in H₂O control treatment; however, after shikimate feeding these differences were no longer apparent (Figure S8). These data did not provide evidence to support the involvement of the PDH enzyme in Tyr catabolism to TCA cycle intermediates.

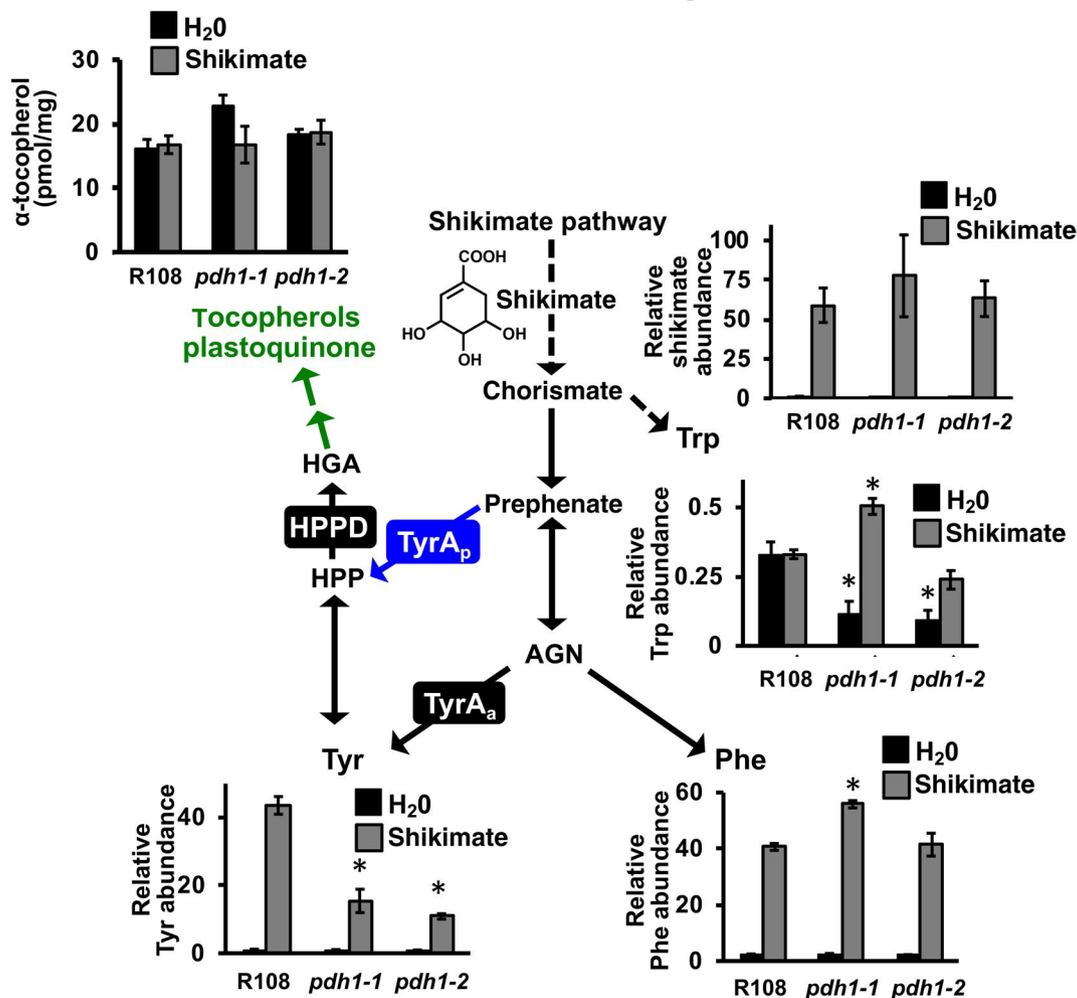


FIGURE 4 Metabolite analysis after shikimate feeding in Wt and *pdh1* mutants. Excised leaves from 6-week-old plants were floated on a solution containing H₂O (black bars) or 25 mM shikimate (gray bars) for 8 hr under constant light. Leaves were then used for metabolite analysis using GC-MS (Tyr, Trp, Phe and shikimate) or HPLC (α-tocopherol). Bars represent average relative metabolite abundance of Tyr, Phe, Trp and shikimate ± SEM of three biological replicates. α-tocopherol is shown as the average absolute metabolite abundance in pmol/mg FW ± SEM of three biological replicates. Additional metabolites measured following shikimate feeding are shown in Figures S7 and S8. Significant differences to Wt (R108) control are indicated; **p* ≤ .05

3.6 | PDH has limited role in the legume-rhizobia symbiosis

To test whether PDH plays a role in legume-rhizobia symbiosis (hypothesis IV), Wt and *pdh1-1* mutants were grown side-by-side on low nitrogen Fahræus medium and inoculated with a well-characterized rhizobium of *M. truncatula*, *Sinorhizobium meliloti* Rm1021. At various time points (e.g., 14, 21, and 28 days post-inoculation (dpi)) *pdh1-1* mutants did not display any phenotypic difference from Wt plants, including the number of nodules produced per root (Figure 5a), suggesting that PDH is not essential for nodule development. *M. truncatula* forms indeterminate nodules that display the four standard zones: I, meristematic; II, infection; III, fixation; and IV, senescence zones in both Wt and *pdh1-1* (Figure 5b, Van de Velde et al., 2006). Expression of the bacterial *nifH* gene, which is required for nitrogen fixation, was monitored by using a *S. meliloti* strain expressing a *PrnifH::UidA* fusion to evaluate nodule maturation on *pdh1-1* and Wt

plants (Starker, Parra-Colmenares, Smith, Mitra, & Long, 2006). After 21 and 28 dpi, β-glucuronidase (GUS) activity and the overall nodule development were not altered between *pdh1-1* and Wt (Figure 5c). To measure nitrogenase activity, acetylene reduction assays (ARA, Wych & Rains, 1978) were also performed at 14, 21, and 28 dpi. Although nitrogenase activity in *pdh1-1* appeared to be slightly reduced compared with Wt at 21 and 28 dpi when nodule senescence might have been initiated, no significant differences were observed (Figure 5d). ARA were repeated at the 21 dpi time point with *pdh1-1* and this time including *pdh1-2*. Again, nitrogenase activity was not significantly different between genotypes, even though *pdh1-1* showed a slight reduction, similar to the first experiment (Figure 5e).

To test whether the presence and absence of PDH activity positively correlated with those of nodulation, we also measured PDH activity from plant tissue extracts of over twenty species sampled across the legume phylogeny (Azani et al., 2017, Figure S9) and compared with their capacity to form nodules (Afkhami et al., 2018). All

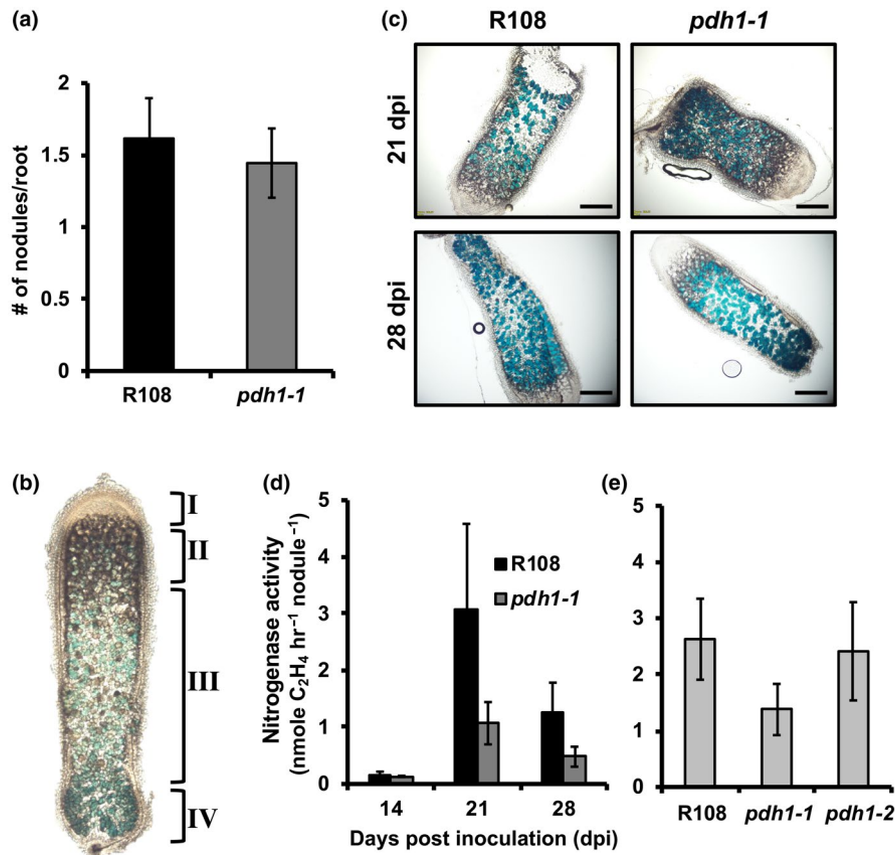


FIGURE 5 The role of PDH in legume-rhizobia symbiosis. (a) Wt and *pdh1-1* were grown on plates with low nitrogen Fahræus medium, inoculated with *S. meliloti*, and the number of nodules was counted at 14 days post-inoculation (dpi). Bars represent average number of nodules per root \pm SEM, $n \geq 11$. (b) Indeterminate nodules from *M. truncatula* consist of four developmental zones: I, meristem; II, infection; III, fixation; IV, senescence located closest to the root. These four developmental zones are shown on a characteristic nodule (enlarged for visualization) from 28 dpi. GUS staining with *PnifH::UidA* shows expression of bacterial *nifH* gene localized to the bacteroids (blue). (c) Thin sections of nodules from 21 and 28 dpi showing expression of *PnifH::UidA*, highlight bacteroid and nodule development in R108 and *pdh1-1*. Approximately equal staining was observed in all nodule developmental zones, suggesting *pdh1-1* is not affected in nodule development or bacteroid number, scale bar = 1 mm. (d) Nitrogen fixation efficiency was measured using an acetylene reduction assay (ARA). Plants were grown under the same conditions as in (a). At 14, 21, and 28 dpi, ethylene production was measured and expressed as the average \pm SEM, $n > 7$. (e) Nitrogen fixation efficiency assay performed in the same way as in (d), but with both mutant alleles and Wt and only at 21 dpi. Activity is expressed as the average \pm SEM, $n > 4$

legumes sister to previously analyzed *G. max* (soybean) and *M. truncatula* (Schenck et al., 2015) showed PDH activity including *Arachis ipaensis* (peanut) (Figure S9a), which possess a TyrA enzyme with similar ADH and PDH activity (Schenck, Holland, et al., 2017). All of these legumes were previously reported to be able to nodulate (Afkhani et al., 2018). When four legume species in the genistoid crown were analyzed (e.g., *Lupinus polyphyllus*), PDH activity was not detectable in any of them, despite the successful detection of ADH activity (Figure S9b). Importantly, these legumes were reported to be able to nodulate, suggesting that legume plants having undetectable levels of PDH activity can still maintain nodulation. Out of four legumes from the mimosoid crown that are known to nodulate, three species showed PDH activity (Figure S9c). In contrast, all four early diverging species tested are incapable of nodulating (Afkhani et al., 2018), but showed PDH activity (Figure S9d). The lack of detectable PDH activity (e.g., the genistoid crown) may be simply due to insufficient sensitivity with HPLC-based assay and/or the use of

tissues that do not express PDH. However, detection of PDH activity in species lacking nodulation (e.g., the early diverging lineages, Figure S9d) strongly support the lack of positive correlation between the presence of PDH activity and nodulation across the legume phylogeny. Phylogenetic sampling (Figure S9) together with *pdh1* mutant analysis (Figure 5) suggest that PDH is not essential for legume-rhizobia symbiosis, though more detailed analyses are needed to fully address its contribution to potentially enhance nitrogen fixation within the nodules in some legumes.

4 | DISCUSSION

In this study, we sought to understand the function of legume-specific PDH enzymes through analysis of *Tnt1*-transposon mutants of PDH in *M. truncatula*. Not only is *M. truncatula* a convenient model system having mutant populations and many genomic/



transcriptomic resources, but *M. truncatula* also has a single *PDH* gene unlike some other legumes, that is, soybean, having two *PDH* genes (Schenck et al., 2015). Analysis of two independent *pdh1* mutant lines suggests that MtPDH1 is responsible for all the *PDH* activity in *M. truncatula* (Figure 1). This observation is consistent with chromatographic separation of *PDH* from *ADH* in soybean, which resulted in the detection of a single *PDH* peak as compared to multiple *ADH* peaks (Schenck et al., 2015). Our genetic data also support that ncADH, and *ADH* enzymes do not contribute to *PDH* activity, consistent with in vitro data of soybean and *M. truncatula* enzymes (Schenck et al., 2015; Schenck, Holland, et al., 2017).

Surprisingly, the null *pdh1* mutants displayed no visible aberrant growth phenotypes (Figure 1). A slight bushy phenotype was observed in the *pdh1-2* allele, but not in *pdh1-1*. Thus, here we focused on consistent responses observed in both alleles to avoid potential pleiotropic effects specific to *pdh1-2*. The lack of substantial phenotypes in *pdh1* is in contrast to the highly compromised growth and leaf development phenotypes of Arabidopsis *adh2/tyra2* mutant (de Oliveira et al., 2019). In *M. truncatula*, we were also unable to recover homozygous mutants of the canonical *ADH*, which is a single copy gene in *M. truncatula* as compared to two *ADH* copies in Arabidopsis (Rippert & Matringe, 2002), further suggesting the essential nature of the *ADH*-mediated Tyr pathway in legumes. Thus, the canonical plastid-localized *ADH* pathway is the predominant Tyr biosynthetic route and cannot be compensated by the cytosolic *PDH* pathway in legumes.

Using the *pdh1* mutants, this study systematically evaluated potential roles of *PDH* in Tyr biosynthesis (hypothesis I), biosynthesis of Tyr- and HPP-derived metabolites (II), Tyr catabolism and senescence (III), and legume-rhizobia symbiosis (IV). The obtained data provide evidence for support of hypothesis I that the *PDH* enzyme is involved in Tyr biosynthesis when carbon flux is increased through the shikimate pathway. Although Tyr levels were not altered between Wt and mutants under standard conditions (Figure 3a), upon feeding with shikimate, Wt accumulated over twofold more Tyr than *pdh1-1* and *pdh1-2* (Figure 4), suggesting that the *PDH* enzyme contributes to synthesis of Tyr under enhanced flux through the shikimate pathway. Although AAA biosynthesis is localized to the plastids (Bickel et al., 1978; Jung et al., 1986; Maeda & Dudareva, 2012), some isoforms of the shikimate and AAA pathway enzymes were detected in the cytosol (Ding et al., 2007; Ganson, D'Amato, & Jensen, 1986; Rubin & Jensen, 1985). Cytosolic CM activities have been detected from various plant species including soybean (Goers & Jensen, 1984; Kuroki & Conn, 1989; Mobley et al., 1999; Schenck et al., 2015), which are not regulated by the AAAs, unlike their plastidic counterparts (Eberhard et al., 1996; Mobley et al., 1999; Westfall, Xu, & Jez, 2014). Tyr-insensitive *PDH* enzymes provide a cytosolic route for conversion of prephenate into HPP, which can be further transaminated to Tyr by cytosolic Tyr-AT enzymes (Figure 1) (Wang et al., 2016). Although the *PDH*-mediated Tyr pathway is not the predominant route for Tyr biosynthesis, it may provide an alternative route for additional Tyr synthesis under some conditions that increase flux through the shikimate pathway.

The current data also partly support hypothesis II that the *PDH* enzyme contributes to the production of specialized metabolites derived from Tyr and/or HPP in legumes. Although Tyr/HPP-derived tocopherols were not altered in the *pdh1* mutants under standard growth conditions (Figure 3b), high light-induced tocopherol accumulation was partially attenuated in the *pdh1* mutant (Figure S3). Therefore, the *PDH* enzyme may also contribute to tocopherol biosynthesis under stress conditions. Furthermore, after shikimate precursor feeding, Wt accumulated more Tyr than mutants (Figure 4), suggesting that the *PDH* enzyme is involved in synthesizing Tyr and possibly downstream compounds when the shikimate pathway flux is increased. Therefore, the alternative *PDH* pathway could be used under certain conditions (e.g., wounding or other stresses) to support production of Tyr/HPP-derived specialized metabolites. Specialized metabolic pathways often emerge through duplication and neofunctionalization of genes from primary metabolic pathways (Moghe & Last, 2015; Schenck & Last, 2019; Weng et al., 2012). The legume *PDH* enzyme emerged as the result of a recent gene duplication of an *ADH* gene followed by a shift in substrate specificity from arogenate to prephenate (Schenck, Holland, et al., 2017). Thus, legumes may be able to divert the shikimate pathway flux to provide additional HPP or Tyr in the cytosol for the synthesis of downstream specialized metabolites including tocopherols. To further test this possibility, additional metabolomics experiments were conducted, but did not identify any putative HPP-derived compounds that were absent or lower in *pdh1* mutants than Wt (Table S1). Although there are many Tyr and HPP-derived specialized metabolites produced in plants (e.g., rosmarinic acid, betalains, dhurrin, and benzylisoquinoline alkaloids), none appear to be specific to the legume lineage or correlate with the distribution of *PDH* activity (Figure S9) (Schenck & Maeda, 2018). Some species of the legume genus *Inga* can accumulate Tyr at 20% of leaf dry weight, which deters insect predation (Lokvam, Brenes-Arguedas, Lee, Coley, & Kursar, 2006). Further analysis identified that *Inga* species with high *PDH* expression have hyper-accumulation of Tyr-derived specialized metabolites, such as Tyr-gallates (Coley et al., 2019). Therefore, more comprehensive analyses of Tyr/HPP-derived metabolites under different conditions, tissue types, and other legume species may identify specialized metabolites downstream of the *PDH* enzyme in legumes.

The current data did not provide sufficient support for hypothesis III that the *PDH* pathway potentially contributes to Tyr catabolism and senescence. As Tyr is the most energetic amino acid (Hildebrandt, Nunes Nesi, Araújo, & Braun, 2015), its catabolism is a crucial process in recovering energy. Mutations in Tyr catabolic genes can have dramatic effects on plant development (Han et al., 2013). Tyr is catabolized into fumarate, a TCA pathway intermediate, first by conversion into HPP via a Tyr-AT enzyme and further through the canonical Tyr catabolism pathway (Figure S5a) (Hildebrandt et al., 2015). Knockout of a soybean *HGO* did not cause lethality but led to an increase in vitamin E content by twofold, suggesting that a significant amount of carbon flux is diverted into vitamin E production when Tyr catabolism is blocked (Stacey et al., 2016). Furthermore, MtPDH1 was co-expressed with HPPD, which together



provide a direct pathway for homogentisate production. This potential coordination of PDH and HPPD, and cytosolic localization, can bypass three enzymatic steps of the ADH-mediated Tyr biosynthetic pathway, catalyzed by PPA-AT, ADH, and Tyr-AT to enter into the canonical Tyr catabolism pathway (Figures 1 & S5a). Additionally, *PDH* was co-expressed with many senescence-related genes (Figure S1, Van de Velde et al., 2006; Kusaba, Tanaka, & Tanaka, 2013; Xi, Chen, Nakashima, Wang, & Chen, 2013) suggesting that *PDH* may function when Tyr catabolism is enhanced. Despite these correlative data, however, experiments to test the potential link between PDH and senescence did not provide evidence to support this hypothesis: upon dark-induced (Figure S6) and natural senescence (Figure S4) conditions that likely stimulate Tyr catabolism, no phenotypic differences were observed between mutants and Wt. Also, genes involved in Tyr catabolism were not consistently altered in mutants as compared to Wt (Figure S5b). Although *PDH* expression was reduced, *PDH* enzymatic activity was unaltered (Figure S4c), suggesting that there may be some unknown post-transcriptional regulation on *PDH* following the induction of senescence. Thus, further experiments under different conditions (e.g., specific stress that induces Tyr catabolism) are needed to address the potential role of *PDH* in Tyr catabolism and senescence.

The data obtained in this study failed to directly support the hypothesis IV that the *PDH* pathway plays a role in legume-rhizobia symbiosis. Legumes initiate a symbiotic relationship with soil-dwelling rhizobia when there is insufficient nitrogen (Oldroyd, Murray, Poole, & Downie, 2011). A chemical communication ensues that ultimately results in compatible rhizobia invading legume roots and formation of a new organ, the root nodule (Oldroyd, 2013). In the nodule, rhizobia fix atmospheric dinitrogen into ammonium, which is assimilated by the plant through the glutamine synthetase-glutamine oxoglutarate aminotransferase (GS-GOGAT) cycle, in which glutamine and glutamate are key amino acid carriers (Krapp, 2015). Mutations in *MtPDH1* did not alter nodule numbers (Figure 5a) or the developmental progression of the nodules (Figure 5c). Furthermore, nitrogenase activity was not significantly affected in mutants at any stage during the symbiotic interaction even when senescence might have been initiated (Figure 5d,e). Alterations in nitrogen assimilation can lead to reduced nodulation in legumes (Matamoros et al., 1999; Streeter & Wong, 1988): for example, plants treated with a GS inhibitor phosphinothricin (PPT) result in loss of nodulation (Seabra et al., 2012) and also stimulate *MtPDH1* expression (Figure S1) as well as many other genes. Interestingly, *pdh1-1* and *pdh1-2* had reduced glutamine levels in the H₂O-treated control leaves after 8 hr (Figure S7), which also persisted after shikimate feeding (Figure S7). Reduced glutamine levels in the *pdh1* mutants may, in turn, affect the symbiotic efficiency with rhizobia, although no statistically-significant reduction in nitrogenase activity was observed (Figure 5d,e). Furthermore, *PDH* activity, which was measured in a phylogenetically diverse group of legumes, does not correlate with ability to nodulate (Afkhami et al., 2018; Azani et al., 2017; Figure S9). Thus, *PDH* may provide an adaptive advantage to some legumes but is likely not directly involved in the legume-rhizobia symbiosis.

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CONFLICT OF INTEREST

The authors declare no conflicts of interest.

AUTHOR CONTRIBUTION

CAS and HAM conceptualized the project. CAS, JMA, LWS, and HAM designed experiments. CAS, JW, DJ, and KG carried out experiments and interpreted results. KSM and JW isolated *Tnt1* mutant lines. CAS and HAM wrote the manuscript. All authors read and edited the manuscript.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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