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## **RUNNING HEAD: Integrated Metabolomics Identifies Saponin P450s**

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**Journal Research Area:** Biochemistry and Metabolism

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4 **TITLE: Integrated Metabolomics Identifies CYP72A67 and CYP72A68**  
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6 **Oxidases in the Biosynthesis of *Medicago truncatula* Oleanate Sapogenins**  
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4 **ABSTRACT (150-250 words)**  
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7 Triterpene saponins are important bioactive plant natural products found in many plant families  
8 including the Leguminosae. Here we characterize two *Medicago truncatula* cytochrome P450  
9 enzymes, MtCYP72A67 and MtCYP72A68, involved in saponin biosynthesis including both *in*  
10 *vitro* and *in planta* evidence. UHPLC-(-)ESI-QToF-MS was used to profile saponin accumulation  
11 across a collection of 106 *M. truncatula* ecotypes. The profiling results identified numerous  
12 ecotypes with high and low saponin accumulation in root and aerial tissues. Four ecotypes with  
13 significant differential saponin content in the root and/or aerial tissues were selected, and  
14 correlated gene expression profiling was performed. Correlation analyses between gene expression  
15 and saponin accumulation revealed high correlations between saponin content with gene  
16 expression of *β-amyrin synthase*, *MtCYP716A12*, and two cytochromes P450 genes, *MtCYP72A67*  
17 and *MtCYP72A68*. *In vivo* and *in vitro* biochemical assays using yeast microsomes containing  
18 MtCYP72A67 revealed hydroxylase activity for carbon 2 of oleanolic acid and hederagenin. This  
19 finding was supported by functional characterization of *MtCYP72A67* using RNAi-mediated gene  
20 silencing in *M. truncatula* hairy roots, which revealed a significant reduction of 2β-hydroxylated  
21 sapogenins. *In vivo* and *in vitro* assays with MtCYP72A68 produced in yeast showed  
22 multifunctional oxidase activity for carbon 23 of oleanolic acid and hederagenin. These findings  
23 were supported by overexpression of *MtCYP72A68* in *M. truncatula* hairy roots, which revealed  
24 significant increases of oleanolic acid, 2β-hydroxyoleanolic acid, hederagenin and total saponin  
25 levels. The cumulative data support that MtCYP72A68 is a multisubstrate, multifunctional oxidase  
26 and MtCYP72A67 is a 2β-hydroxylase, both of which function during the early steps of triterpene-  
27 oleanate sapogenin biosynthesis.  
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48 **Keywords:** Saponin, sapogenin, cytochrome P450, CYP72A67, CYP72A68, *Medicago*  
49 *truncatula*, integrated metabolomics  
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## INTRODUCTION

Saponins are steroidal, steroidal alkaloid or triterpenoid metabolites that are typically conjugated with sugars and present in numerous plant species, including members of the genus *Medicago* (Augustin et al. 2011; Avato et al. 2006; Bialy et al. 1999; Gholami et al. 2014; Huhman et al. 2005; Huhman and Sumner 2002; Pollier et al. 2011; Tava et al. 2011). Many triterpene saponin aglycones (saponins without sugars, also known as sapogenins) are oxidized at various positions on the aglycone (Figure 1). These oxidized positions are often further conjugated with varying numbers of sugars to yield a multitude of saponins. Saponins possess diverse biological activities and plant beneficial properties, which include antifungal, antibacterial, antiviral, antitumor, molluscicidal, insecticidal and antifeedant activities (Augustin et al. 2011; Avato et al. 2006; Dixon and Sumner 2003; Klita et al. 1996; Lu and Jorgensen 1987; Sparg et al. 2004; Yan et al. 2013). In addition, they also affect plant development, including seed germination, vegetative growth and differentiation, fruiting and nodulation (Moses, Papadopoulou, et al. 2014). The pharmacological properties of saponins have been exploited in herbal medicines and, more recently, evaluated for their anticholesterolemic, anticancer, and adjuvant properties (Haridas et al. 2001; Kirk et al. 2004; Kuljanabhagavad et al. 2008; Shibata 2001).

Triterpene saponins constitutively accumulate in plants. However, the saponin biosynthetic pathway and additional accumulation of saponins are further induced during wounding, herbivory and by methyl jasmonate, which is a signaling compound associated with the induction of many defense-responsive plant metabolites (Broeckling et al. 2005; Gholami et al. 2014; Naoumkina et al. 2007; Suzuki et al. 2005). Although saponins are beneficial plant defense compounds, saponins in legume forages, such as alfalfa (*Medicago sativa*), are of particular and substantial economic importance because they result in impaired digestion and reduced weight gain in ruminant animals (Lu and Jorgensen 1987; Sen et al. 1998). Thus, saponins are considered antifeedants in premiere forages such as alfalfa. A detailed molecular and biochemical understanding of saponin biosynthesis would enable future metabolic engineering of crops with increased defense properties resulting in improved fitness and field productivity, and decreased anti-nutrient properties that would result in enhanced livestock weight gain performance. Metabolically engineered legumes with improved performance and

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4 nutritive value would have substantial commercial value that would advance the plant  
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6 biotechnology industry.

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8         Recent studies in the *Medicago* genus have focused on elucidating the relationship  
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10 between the biological activities of saponins and their chemical structures. The aglycone type,  
11 along with the nature and position of the sugar moieties, appear to correlate with different  
12 biological properties (Gholami et al. 2014; Tava et al. 2011). Two distinct classes of sapogenins  
13 can be differentiated in *Medicago* spp. based upon the position and the degree of oxidation: (i)  
14 sapogenins possessing a hydroxyl group at the C-24 position, without any substituent at the C-28  
15 position atom (i.e., soyasapogenols; A, B and E); and (ii) sapogenins possessing a carboxyl  
16 group at the C-28 position that often also contain different oxidized states at the C-23 position  
17 (i.e., oleanate sapogenins with H, OH, CHO, or COOH at the C-23 position) (Carelli et al. 2011;  
18 Fukushima et al. 2011; Gholami et al. 2014). Some saponins possess hemolytic activity that  
19 results from their affinity for membranes, and this activity is related to the nature of the aglycone  
20 moiety (Augustin et al. 2011). No hemolytic activity was observed for soyasapogenols (Yoshiki  
21 et al. 1998), while oleanate derived sapogenins possessed high (hederagenin and medicagenic  
22 acid glycosides) to moderate (zanhic acid glycosides) hemolytic activities (Oleszek 1996).  
23 Recently, ectopic accumulation of bioactive monoglycosylated saponins was suggested to affect  
24 the integrity of *M. truncatula* roots themselves; hence, saponin producing plants need to develop  
25 self-protection mechanisms to allow accumulation of saponins (Pollier et al. 2013).  
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40         The first committed step in triterpene saponin biosynthesis is the cyclization of 2,3-  
41 oxidosqualene. This reaction is catalyzed by a specific oxidosqualene cyclase (e.g.,  $\beta$ -*amyrin*  
42 *synthase*;  $\beta$ AS) which has been functionally characterized in many plant species (Inagaki et al.  
43 2011; Iturbe-Ormaetxe et al. 2003; Sawai and Saito 2011; Suzuki et al. 2002; Thimmappa et al.  
44 2014). Subsequent modifications that impart functional properties and diversify the basic  
45 triterpene backbone include the addition of small functional groups such as hydroxy, keto,  
46 aldehyde and carboxy moieties which are often followed by glycosylation (Augustin et al. 2012;  
47 Miettinen et al. 2018; Moses, Papadopoulou, et al. 2014; Thimmappa et al. 2014). The oxidative  
48 reactions prior to glycosylation are catalyzed by cytochrome P450-dependent monooxygenases  
49 (P450s). To date, several P450s that utilize  $\beta$ -amyrin as a substrate have been identified in  
50 dicotyledonous plants, whereas just one (CYP51H10) has been identified in monocots (Geisler et  
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4 al. 2013; Kunii et al. 2012). In oat (*Avena strigosa*) AsCYP51H10 (*Sad2*) is a multifunctional  
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6 P450 that catalyzes the oxidation of  $\beta$ -amyirin on both the C and D rings to give 12,13 $\beta$ -epoxy-  
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8 16 $\beta$ -hydroxy- $\beta$ -amyirin, an intermediate of root saponin biosynthesis (Geisler et al. 2013).  
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10 Currently, a significant number of other saponin biosynthetic cytochrome P450s have been  
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12 characterized and *in planta* activity inferred via heterologous expression predominantly in yeast  
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14 and to a lesser extent in *Nicotiana benthamiana*. In dicots, GuCYP88D6 in licorice (*Glycyrrhiza*  
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16 *uralensis*, Fabaceae) catalyzes the C-11 oxidation of  $\beta$ -amyirin in glycyrrhizin biosynthesis (Seki  
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18 et al. 2008). Members of the CYP93E subfamily of P450s have been shown to catalyze the C-24  
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20 hydroxylation of  $\beta$ -amyirin in soyasapogenol biosynthesis (Fukushima et al. 2011; Moses,  
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22 Thevelein, et al. 2014; Seki et al. 2008; Shibuya et al. 2006) CYP87D16 catalyzes the C-16 $\alpha$   
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24 hydroxylation of  $\beta$ -amyirin in the biosynthesis of maesasaponins (Moses et al. 2015). Members  
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26 of the CYP716 subfamily of P450s catalyze various oxidations on the  $\beta$ -amyirin backbone,  
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28 including the three-step oxidation of  $\beta$ -amyirin at the C-28 position to yield oleanolic acid by  
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30 CYP716A12 (Carelli et al. 2011; Miettinen et al. 2017), the C-16 $\alpha$  hydroxylation of  $\beta$ -amyirin by  
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32 CYP716Y1 (Moses, Pollier, et al. 2014), and the C-3 oxidation by CYP716A14 (Moses et al.  
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34 2015). In addition, CYP72A154 oxidizes  $\beta$ -amyirin at the C-30 position (Seki et al. 2011) and  
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36 several P450s have been identified that further modify oxidation products of  $\beta$ -amyirin, such as  
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38 MtCYP72A61 and GmCYP72A69 (Sundaramoorthy et al. 2018; Yano et al. 2016) that  
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40 hydroxylate 24-hydroxy- $\beta$ -amyirin at the C-22 and C-21 positions, respectively, in soyasaponin  
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42 biosynthesis, MtCYP72A68 that oxidizes C-23 of oleanolic acid, and MtCYP72A67 that  
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44 catalyzes oxidation at C-2 position (Biazzi et al. 2015; Fukushima et al. 2013). However, several  
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46 P450s and glycosyltransferases involved in saponin biosynthesis still remain uncharacterized and  
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48 *in planta* evidence for tentatively identified genes is minimal.

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50 Correlated gene expression analysis has emerged as a powerful tool for predicting gene  
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52 function as correlation is suggestive of related biological processes (Hirai et al. 2005; Hirai and  
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54 Saito 2004). Such correlations are facilitated by the availability of large quantities of public gene  
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56 expression data which enable the calculation of gene coexpression correlation scores across  
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58 thousands of samples (Usadel et al. 2009). Naoumkina et al. (2010) described a set of  
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60 coexpressed *M. truncatula* genes based on comprehensive clustering of methyl jasmonate-  
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62 induced transcript expression patterns along with chromosomal location analysis (Naoumkina et  
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4 al. 2010). However, the identification of the specific P450 enzymes responsible for the  
5 production of particular metabolites is still a difficult task due to the large numbers and  
6 significant diversity within the P450 multigene family (Augustin et al. 2011).  
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10 We support that an integrated approach that includes genomic, transcript, and metabolite  
11 profiling along with ectopic expression offers a more productive strategy to identify and  
12 functionally characterize new biosynthetic genes (Seki et al. 2011). Such a combined approach  
13 has been successfully used to identify several glycosyltransferase genes involved in the  
14 biosynthesis of triterpene saponins in *M. truncatula* (Achnine et al. 2005; Naoumkina et al.  
15 2010). In the Naoumkina et al., 2010 report, the putative roles of MtCYP72A67 and  
16 MtCYP72A68 in saponin biosynthesis were proposed based upon correlated gene expression  
17 with functionally characterized genes such as those encoding  $\beta$ -amyrin synthase,  
18 MtCYP716A12, MtUGT73F3 and MtCYP93E2 (Carelli et al. 2011; Fukushima et al. 2013;  
19 Naoumkina et al. 2010; Seki et al. 2011). Accordingly, *MtCYP72A67* and *MtCYP72A68*  
20 expression was found to be regulated by the transcription factor TRITERPENE SAPONIN  
21 BIOSYNTHESIS ACTIVATING REGULATOR2 (TSAR2), the regulator of hemolytic  
22 triterpene saponin metabolism in *M. truncatula* (Mertens et al. 2016). Since then, we have also  
23 presented our *in vitro* biochemical assays of recombinant MtCYP72A67 and MtCYP72A68  
24 heterologously expressed in yeast ( Sumner et al. 2012; Tzin et al. 2012a, 2012b). Similar *in vivo*  
25 enzymatic activities have been reported in engineered yeast strains for MtCYP72A61,  
26 MtCYP93E2, MtCYP72A67 MtCYP72A68 and MtCYP716A12 (Biazzi et al. 2015; Fukushima  
27 et al. 2013). However, evidence obtained through studies in heterologous microbial systems does  
28 not always equal *in planta* function. For example, Biazzi *et al*, 2015 associated MtCYP72A68  
29 with medicagenic acid biosynthesis based upon *in vitro* yeast assays. However, we report here  
30 that MtCYP72A68 is a multi-functional oxidase responsible for hederagenin, gypsogenin and  
31 gypsogenic acid biosynthesis based upon *in vivo* and *in planta* evidence. In addition, *in planta*  
32 triterpene engineering has been hampered by a lack of knowledge about the regulatory  
33 mechanisms controlling gene expression (Sawai and Saito 2011). Hence, a challenge for future  
34 triterpenoid research will be to identify the transcription or other regulatory factors that steer  
35 their biosynthesis (Biazzi et al. 2015; Moses et al. 2013)  
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4 **OBJECTIVES**  
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7 In this report, we describe a highly productive approach for the discovery of triterpene  
8 biosynthetic genes and provide both *in vitro* and *in planta* characterization of two CYP72 family  
9 genes. More specifically, *MtCYP72A67* and *MtCYP72A68* were identified based upon large-  
10 scale, correlated metabolite accumulation and gene expression. Highly correlated genes were  
11 then functionally characterized as multisubstrate, mono and multifunctional oxidases in  
12 triterpene saponin biosynthesis using heterologous *in vitro* and *in vivo* yeast expression assays,  
13 heterologous *in vivo* tobacco expression assays, and *in planta* using *M. truncatula* hairy root  
14 cultures and Tnt1 mutants.  
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24 **METHODS**  
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26 **Germplasm plant materials**  
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28 Seeds for the *Medicago truncatula* ecotype collection were obtained from Jean-Marie Prosperi at  
29 L'Institut National De La Recherche Agronomique (INRA;  
30 [http://www.international.inra.fr/the\\_institute](http://www.international.inra.fr/the_institute)). The ecotype collection used in the present study  
31 was described previously (Ronfort et al. 2006). Single seed descent lines for all of the INRA  
32 ecotypes were developed on site at the Noble Foundation.  
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38 **Plant growth conditions**  
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40 Plants were grown in a root cone system (Stewe and Sons, OR) with Turface MVP medium  
41 (Profile Products, Buffalo Grove, IL) in a Conviron TCR180 walk-in growth chamber  
42 maintained at 90% humidity and at an average temperature of 24°C day (16 h) and 20°C night (8  
43 h). Plants were fertilized daily with 15 ppm nitrogen (20-10-20 Peat-Lite Special; The Scotts  
44 Company). Five-week-old plants were harvested, and the Turface was washed quickly from the  
45 roots. Plants were dissected into roots and aerial parts, which were flash frozen in liquid nitrogen  
46 and stored at -80 °C.  
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53 **Metabolomics analyses by UHPLC-(-)ESI-QToF-MS and GC-MS**  
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55 Lyophilized tissues were ground into a fine powder using a mortar and pestle, and 10 mg of  
56 powder was extracted with 1 ml of 80% methanol in a one-dram vial for two hours on an orbital  
57 shaker. An internal standard containing 18 µg/ml umbelliferone was used in all samples.  
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4 Extracted samples were centrifuged for 30 min at 2900 x g at 4°C, and supernatants were  
5 transferred to UHPLC-MS autosampler vials. The HPLC-(-)ESI-QToF-MS analyses were  
6 performed with a Waters Acquity UHPLC system coupled to a Waters Premier hybrid  
7 quadrupole time-of-flight (QTOF) mass spectrometer (<http://www.waters.com>). A reverse-phase,  
8 UPLC BEH 1.7 μm C18, 2.1 mm x 150 mm column (Waters) was used for separations. The  
9 mobile phase consisted of eluent A (0.1% [v/v] acetic acid/water) and eluent B (100%  
10 acetonitrile). Separations were achieved using a linear gradient of 95 % to 30 % A over 30 min,  
11 30 % to 5 % A over 3 min, and 5 % to 95 % A over 3 min. The flow rate was 0.56 mL/min, and  
12 the column temperature was maintained at 60 °C. Mass-to-charge ratios (*m/z*) of the eluted  
13 compounds were determined in the negative ESI mode from *m/z* 50 to 2,000. The Waters QTOF  
14 Premier was operated using the following instrument parameters: desolvation temperature of 385  
15 °C; desolvation nitrogen gas flow of 850 L/h; capillary voltage of 2.9 kV; cone voltage of 48 eV;  
16 and collision energy of 10 eV. The MS system was calibrated using sodium formate, and  
17 raffinose was used as the lockmass compound. *β*-amyrin, erythrodiol, and cycloartenol assays  
18 were extracted twice with 500 μl of ethyl acetate, dried under nitrogen gas, dissolved in 100 μl  
19 pyridine, *N*-Methyl-*N*-(trimethylsilyl) trifluoroacetamide (MSTFA)-derivitized, and analyzed by  
20 GC-MS as described previously (Broeckling et al. 2005). GC-MS of these compounds were  
21 performed because they do not ionize well by negative ESI due to a lower number of hydroxyl  
22 substituents.

### 39 **Data Processing**

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41 Raw UHPLC-(-)ESI-QToF-MS data files were annotated and quantified using MarkerLynx XS  
42 (Waters; [www.waters.com](http://www.waters.com)) or converted to CDF file format, followed by metabolite data  
43 extraction, alignment, and export using MET-IDEA software (Broeckling et al. 2006; Lei et al.  
44 2012). A target ion list containing 143 known and putative triterpene saponin ions of interest was  
45 used for the targeted saponin analyses. This ion list was selected and annotated based on  
46 authentic standards and previous MS and MS/MS analyses of triterpene saponins conducted  
47 internally in the lab (Huhman et al. 2005; Huhman and Sumner 2002). In addition to the targeted  
48 analyses of saponin content, non-targeted analyses of all samples were performed using Waters  
49 MarkerLynx software. The spectral abundance values for all metabolites in a separation were  
50 normalized to the internal standard of 18 μg/ml umbelliferone. Descriptive statistics were  
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4 performed in Microsoft Excel and JMP (SAS Institute Inc; <http://www.jmp.com>). Correlation  
5 coefficients were calculated using a custom MATLAB script (<http://www.mathworks.com/>).  
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### 8 **RNA extraction, quantitative real-time PCR and Medicago Genome Array**

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10 Total RNA was isolated using modified cetyl-trimethyl-ammonium bromide (CTAB) extraction  
11 as described previously (Pang et al. 2007) or RNeasy Plant Mini Kit (Qiagen, Valencia, CA).  
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14 Total RNA was purified and concentrated using the RNeasy MiniElute Cleanup Kit (Qiagen,  
15 Valencia, CA), and then treated with DNase I (Invitrogen, Carlsbad, CA). RNA concentration  
16 and quality were determined with a Nanodrop spectrophotometer (Thermo Fisher Scientific,  
17 Wilmington, DE). First-strand cDNA was synthesized from 2 µg total RNA in a total volume of  
18 20 µL using SuperScript III reverse transcriptase (Invitrogen). Primers for quantitative real-time  
19 PCR were designed using Primer3 software ([http://www.frodo.wi.mit.edu/cgi-](http://www.frodo.wi.mit.edu/cgi-bin/primer3/primer3_www.cgi)  
20 [bin/primer3/primer3\\_www.cgi](http://www.frodo.wi.mit.edu/cgi-bin/primer3/primer3_www.cgi)). Each primer pair was confirmed to give a single PCR product.  
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23 All primers for PCR amplification are listed in Supplemental Table S7. The parameters and  
24 analysis of the qRT-PCR were as described previously (Pang et al. 2007). All reactions were  
25 performed with three technical replicates. Data were analyzed using the SDS 2.2.1 software  
26 (Applied Biosystems). Five hundred nanograms of purified RNA for each of the three biological  
27 replicates were used for probe synthesis using a GeneChip3' IVT express kit, according to  
28 manufacturer's instructions (Affymetrix). Hybridization of probes to Affymetrix GeneChip  
29 *Medicago* genome arrays and scanning of arrays was carried out as described (Benedito et al.  
30 2008). Raw data were normalized by robust multichip averaging (RMA), as previously described  
31 (Irizarry et al. 2003).  
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### 43 **Expression of MtCYP72A67 and MtCYP72A68 genes in WAT11 yeast system**

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45 Coding sequences information for *MtCYP72A67* and *MtCYP72A68* (Li et al. 2007) were  
46 obtained from NCBI genebank: DQ335782 and DQ335780, respectively. Primers were designed  
47 for the coding sequence by using Primer3 software (Rozen and Skaletsky 2000). The upstream  
48 cloning primer for *MtCYP72A67* and *MtCYP72A68* included both a BamHI restriction site and a  
49 kozak yeast translation initiation sequence, where the downstream cloning primer included an  
50 EcoRI cut site. *MtCYP72A67* and *MtCYP72A68* were amplified from cv. Jemalong A17 aerial  
51 tissue cDNA template using Platinum Hi-Fi Taq polymerase (Invitrogen, Carlsbad, CA). The  
52 *CYP72A68* PCR product was cloned into the pGEM-T easy vector (Promega, WI), then  
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4 sequenced using M13 forward and reverse primers. *CYP72A67* and *CYP72A68* were excised  
5 from the p-GEM easy vector via a BamHI and EcoRI restriction digest and sub-cloned into the  
6 pYeDP60 vector (Pompon et al. 1996) and sequenced using the GAL10 promoter. WAT11 yeast  
7 cells were transformed as previously reported (Greenhagen et al. 2003; Urban et al. 1997) and  
8 confirmed by PCR.  
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### 13 **Recombinant expression and microsomal preparations for enzymatic assays**

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15 WAT11 yeast cells were transformed with pYeDP60-MtCYP72A68, pYeDP60-MtCYP72A67  
16 or empty pYeDP60 vector, and microsomes were prepared as previously described (Greenhagen  
17 et al. 2003). For initial *in vitro* studies, 100 µg of total microsomal protein (quantified via  
18 Bradford assay) (Bradford 1976) was incubated for 2 h at 30°C in a 500 µl reaction volume of 50  
19 mM potassium phosphate buffer (pH 7.25) containing 1 mM NADPH and 40 µM purified  
20 authentic substrate of either  $\beta$ -amyrin, cycloartenol, erythrodiol, oleanolic acid or hederagenin.  
21 An NADPH generation system (3.3 mM glucose-6-phosphate, 1.3 mM of NADPH, 3.3 mM  
22 magnesium chloride and 0.4 mM glucose-6-phosphate dehydrogenase) (Mene-Saffrane and  
23 Dellapenna, 2009) was also used. All enzymatic assays were performed in triplicate. All reaction  
24 assays were analyzed using UHPLC-(-)ESI-QToF-MS as described above, or dissolved in 100µl  
25 pyridine, MSTFA-derivitized, and analyzed by GC-MS as previously described (Broeckling et  
26 al. 2005).  
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### 38 **Generation and cultivation of TM3-derived strains**

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40 For expression in TM3-derived yeast strains, *MtCYP72A67* and *MtCYP72A68* were amplified  
41 from *M. truncatula* cDNA with primers P1+P2 and P3+P4, respectively, cloned into  
42 pDONR221, sequence verified and gateway recombined in the yeast expression vector  
43 pAG423GAL-ccdB (Addgene plasmid 14149; (Alberti et al. 2007)). Primer sequences for P1  
44 through P8 are listed in Supplemental Table S8. The construct encoding the self-processing  
45 polyprotein with the *M. truncatula* cytochrome P450 reductase 1 (*MtCPRI*; Medtr3g100160) and  
46 *CYP716A12* was created by amplifying *MtCPRI* without a stop codon and with a 3'-overhang of  
47 the partial T2A sequence with primers P5+P6 and *CYP716A12* with a 5'-overhang of the partial  
48 T2A sequence with primers P7+P8. Subsequently the 2 fragments were fused by PCR with  
49 primers P5+P8, cloned into pDONR221, sequence verified and recombined in the yeast  
50 expression vector pAG425GAL-ccdB (Addgene plasmid 14153; (Alberti et al. 2007)). The yeast  
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4 strains (Supplemental Table S8) were generated from strain TM3 and cultivated as described  
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6 (Moses, Pollier, et al. 2014). Briefly, yeast precultures were grown with agitation in synthetic  
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8 defined (SD) medium containing glucose with appropriate dropout (DO) supplements (Clontech)  
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10 at 30 °C for 18 to 20 h. Gene expression was induced by inoculating washed precultures into  
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12 synthetic defined Gal/Raf medium containing galactose and raffinose with appropriate dropout  
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14 supplements (Clontech) to a starting OD<sub>600</sub> of 0.25 on day 1. The induced cultures were  
15  
16 incubated for 24 h, and on day 2, methionine and methylated β-cyclodextrins (MβCD) were  
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18 added to 1 and 5 mM, respectively. After a further 24 h incubation, MβCD was added once again  
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20 to 5 mM on day 3, and on day 4 all cultures were extracted for metabolite analyses. For organic  
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22 extracts of the spent medium, 1 mL of the yeast culture was extracted twice with 0.5 mL of  
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24 hexane and once with 0.5 mL of ethyl acetate. The organic extracts were pooled, vaporized to  
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26 dryness and trimethylsilylated for GC-MS analysis. GC-MS analysis was carried out as  
27  
28 described (Moses, Pollier, et al. 2014) .

### 29 ***Nicotiana benthamiana* leaf infiltration**

30 For transient expression in *N. benthamiana* leaves, the coding sequence of *GgβAS*,  
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32 *MtCYP716A12*, *MtCYP72A67*, and *MtCYP72A68* were Gateway recombined from their  
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34 pDONR221 entry vectors into the binary vector pK7WG2D (Karimi et al. 2002). The resulting  
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36 constructs were individually introduced into the *A. tumefaciens* strain C58C1, carrying the helper  
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38 plasmid pMP90. *Agrobacterium* strains were grown for 2 d in a shaking incubator (150 rpm) at  
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40 28 °C in 5 mL yeast extract broth medium, supplemented with 100 µg/mL kanamycin, 100  
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42 µg/mL spectinomycin, and 20 µg/mL gentamycin. After incubation, 0.5 mL of bacterial culture  
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44 was used to inoculate 9.5 mL of yeast extract broth medium supplemented with antibiotics and  
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46 containing 10 mM MES (pH 5.7) and 20 mM acetosyringone. After an additional overnight  
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48 incubation (150 rpm, 28 °C), strains for transient coexpression were mixed, collected via  
49  
50 centrifugation, and resuspended in 5 mL of infiltration buffer (100 mM acetosyringone, 10 mM  
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52 MgCl<sub>2</sub>, and 10 mM MES, pH 5.7). The amount of bacteria harvested for each construct was  
53  
54 adjusted to a final OD<sub>600</sub> of 0.3 after resuspension in the infiltration buffer. After 2 to 3 h  
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56 incubation at 150 rpm and 28 °C, the bacteria mixtures were infiltrated to the abaxial side of  
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58 fully expanded leaves of 3- to 4-week-old *N. benthamiana* plants grown at 25 °C in a 14-h/10-h  
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60 light/dark regime. The infiltrated plants were incubated under normal growth conditions for 5 d  
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62 prior to metabolite analysis. *Nicotiana benthamiana* infiltrated leaves were harvested and ground  
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4 to a fine powder in liquid nitrogen for metabolite analyses. Then, 0.4 g of ground leaf material  
5 was extracted with 1 mL of methanol for 10 min and centrifuged for 5 min at 20,800 x g. The  
6 resulting organic extract was evaporated to dryness under vacuum and subsequently resuspended  
7 in 0.5 mL of water and 0.5 mL of ethyl acetate. After centrifuging again for 5 min at 20,800 x g,  
8 the organic phase was removed, vaporized to dryness, and trimethylsilylated for GC-MS analysis  
9 which was carried out as described previously (Moses et al. 2015).  
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### 15 **Ectopic expression of MtCYP72A67 and MtCYP72A68 in *Medicago truncatula* hairy roots**

16 The coding sequence of *MtCYP72A67* and *MtCYP72A68* were amplified from cDNA  
17 synthesized from *M. truncatula* (cv. Jemalong A17) aerial tissue using Platinum Hi-Fi Taq  
18 polymerase (Invitrogen, Carlsbad, CA). The primer sequences used for amplification are listed in  
19 Sup. Table S9. PCR products were cloned into the entry vector pENTR/D/TOPO (Invitrogen)  
20 and sequenced. The entry vectors were recombined into a destination vector, pK7WG2D for  
21 overexpression or pK7GWIWG2D(II) for RNAi (a double-stranded hairpin RNA), by using the  
22 LR clonase reaction (Invitrogen). The vectors were transformed into *Agrobacterium rhizogenes*  
23 (strain ARqua1) by electroporation (Quandt et al. 1993). Transformed colonies were grown on  
24 LB-agar medium at 28°C, with spectinomycin and streptomycin for vector selection. After  
25 confirmation by PCR, transformed agrobacteria were used to transform leaves of *M. truncatula*  
26 (cv. Jemalong A17) and generate hairy roots (Verdier et al. 2012).  
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### 39 **Tnt1 mutant identification of *cyp72a68* lines**

40 The *MtCYP72A68* coding sequence was used for *in silico* blast searches against the Noble  
41 Foundation Tnt1 flanking sequence database [https://medicago-](https://medicago-mutant.noble.org/mutant/database.php)  
42 [mutant.noble.org/mutant/database.php](https://medicago-mutant.noble.org/mutant/database.php), which yielded an insertion event, NF1698 insertion 4 in  
43 R108 ecotype. The Tnt1 insertion in *cyp72a68* was confirmed via cloning and sequencing of the  
44 truncated PCR product using gene-specific and Tnt1 border primers amplified from a  
45 *cyp72a68/cyp72a68* heterozygous NF1698-4 plant, position 503bp. Generation of the *M.*  
46 *truncatula* Tnt1 insertion mutant population and growth of R1 seeds were performed as  
47 described (Tadege et al. 2008). Reverse genetic screening for Tnt1 retrotransposon insertions in  
48 *MtCYP72A68* was performed by using a nested PCR approach (Cheng et al. 2011) and  
49 *cyp72a68*-forward: 5'-GCACGAGGAAAACATTTTCACAC-'3. PCR products from target  
50 mutant NF12169 line were purified with QIAquick PCR purification kit (Qiagen) and sequenced  
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4 by using Tnt1 primers to confirm insertions in *MtCYP72A68* at position. Insertions in  
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6 *Mtcyp72a68* were found at position 503 bp in mutant line NF1698-4 and at 453 bp in line  
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8 NF12169 which were validated by PCR (both calculated from start codon and located on exon).  
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10 All mutants were genotyped however no homozygous Tnt1-*Mtcyp72a67* insertion plants were  
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12 identified.  
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## 15 RESULTS

### 16 **Metabolite profiling of a *M. truncatula* ecotype collection reveals substantial chemical** 17 **diversity and identifies high and low saponin-accumulating ecotypes**

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19 Species-specific germplasm collections (ecotypes; natural genetic variants) are a powerful  
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21 resource for exploring the natural chemical variation in triterpene saponin content (Branca et al.  
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23 2011; Ronfort et al. 2006). In this study, 106 ecotypes were analyzed using high-resolution  
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25 biochemical profiling to characterize the chemical diversity in triterpene saponin content within a  
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27 large *M. truncatula* germplasm collection (collection provided by Dr. Jean-Marie Prospero and  
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29 the French National Institute for Agriculture Research; INRA). The triterpene saponin content in  
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31 each of the above lines was analyzed separately in root and aerial tissues using UHPLC- coupled  
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33 to a hybrid quadrupole time-of-flight mass spectrometer operated in the negative electrospray  
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35 ionization mode (UHPLC-(-)ESI-QToF-MS). Metadata compliant with the Metabolomics  
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37 Standards Initiative (Fiehn et al. 2007; L. W. Sumner et al. 2007) are summarized in  
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39 Supplemental Table S1. Overall, 143 putative and identified triterpene saponins were measured  
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41 based upon unique ion mass-to-charge ratio ( $m/z$ ) and chromatographic retention time pairs.  
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43 Seventeen saponins were rigorously identified based upon co-characterization with authentic  
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45 standards (e.g., 3-Glc-28-Glc-medicagenic acid standard); 53 saponins were tentatively identified  
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47 based upon mass spectral (accurate mass, in-source fragmentation and/or MS/MS) and literature  
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49 data (Huhman et al. 2005; Huhman and Sumner 2002; Pollier et al. 2011); 28 saponins had  
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51 partial annotation based solely on spectral features resulting from probable source fragmentation  
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53 (e.g. possibly Glc-Glc-bayogenin); and the remainder were unknowns. The latter unknown  
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55 saponins were differentiated based upon unique  $m/z$  values and retention times in the same  
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57 manner as the known and putatively identified saponins. Total saponin accumulation values were  
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59 determined for both aerial and root tissues of each ecotype by summing the peak area for each of  
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61 the saponin ion/RT pairs (Supplemental Table S2 and S3). A scatter plot of the total saponin  
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4 content in the aerial tissue versus the root is present in Figure 2. The average value for the total  
5 saponin chromatogram peak areas in 106 ecotypes was 78.32 normalized relative instrument  
6 response to the internal standard (nrir) in the aerial tissue and 285.35 nrir in the roots. Ecotype  
7 ESP105 had the lowest relative content of total saponins in aerial tissue (3.4 nrir), but very high  
8 total saponin accumulation in root tissue (394.1 nrir). In contrast, ecotype GRC43 had the lowest  
9 total saponin accumulation in the root (44.7 nrir) but very high total accumulation in aerial  
10 tissues (131.3 nrir). Thus, these two most diverse ecotypes were selected for further comparative  
11 gene expression analyses.  
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### 21 **Correlated metabolite and gene expression analyses identify *MtCYP72A67* and** 22 ***MtCYP72A68* as cytochrome P450s potentially involved in saponin biosynthesis**

23 Correlations between gene-to-gene expression and gene-to-metabolite accumulation have been  
24 shown to be a powerful tool for the identification of novel natural product biosynthetic genes  
25 (Goossens 2015; Hirai et al. 2005, 2010; Hirai and Saito 2004). Here, correlation analyses  
26 between saponin accumulation and gene expression were performed using selected ecotypes with  
27 high saponin diversity: *M. truncatula* ESP105 and GRC43 (Figure 2, and Supplemental Tables  
28 S2 and S3). These ecotypes were chosen based upon their substantial differential and tissue-  
29 specific accumulation of saponins as described above. We also selected two reference ecotypes:  
30 A17 that was used for genomic sequencing (Young et al. 2011) and R108 that been used in the  
31 generation of a Tnt1 retrotransposon insertion mutant population (Tadege et al. 2008).

32 Affymetrix GeneChip-Medicago Genome Arrays were used for gene expression analyses.  
33 Qualitative and relative quantitative analyses of saponin levels were performed using UHPLC-(-  
34 )ESI-QToF-MS and based upon a unique ion/RT pair list as described above. Saponins were then  
35 grouped according to their triterpene aglycone structures. This grouping of sapogenin-specific  
36 accumulation values was performed for eight different sapogenin aglycones, including oleanolic  
37 acid, hederagenin, bayogenin, medicagenic acid, polygalagenin (putative identification), zanhic  
38 acid, soyasapogenol E and soyasapogenol B (see structures in Figure 1). In addition, two  
39 parameters for total saponin calculations were used: i) total known = sum of the known  
40 sapogenins and ii) total aglycones = sum of total known saponins and unknown aglycones. Gene-  
41 to-gene and gene-to-metabolite Pearson's correlation coefficients (r) were calculated and  
42 clustered using gene expression levels for 23 P450 probe sets implicated in a previous report  
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4 (Naoumkina et al. 2010), *β*-*amyirin synthase* (*βAS*), nine other known genes related to *M.*  
5 *truncatula* terpene biosynthesis, known sapogenin aglycones and total saponin levels (Figure  
6 3). The probe sets of four genes, *MtCYP72A67*, *MtCYP72A68*, *MtCYP716A12* and *MtβAS*, were  
7 positively correlated to each other (Pearson's  $r \geq 0.5$ , upper left triangle) and significant (*P* value  
8  $< 0.01$ , lower right triangle) as highlighted in the Figure 3 heat map. These probe sets were also  
9 highly correlated with the total saponin content and medicagenic acid. The expression levels of  
10 *MtCYP72A67*, *MtCYP72A68*, *MtCYP716A12* and *MtβAS* with the sapogenin aglycones level are  
11 presented in Supplemental Figure S1 and the qRT-PCR verification of the gene expression levels  
12 in Supplemental Figure S2 along with the full list of transcriptome and metabolome data for the  
13 selected *M. truncatula* ecotypes in Supplemental Table S4.  
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23 The correlation data reported here were validated with previous studies which reported  
24 that both *βAS* (the first enzymatic step of triterpene saponin biosynthesis) and *MtCYP716A12*  
25 (enzyme associated with oleanate sapogenin biosynthesis) are key enzymatic steps in triterpene  
26 saponin biosynthesis in *M. truncatula* (Carelli et al. 2011; Fukushima et al. 2011; Suzuki et al.  
27 2002). The data also highlighted two more important genes, *MtCYP72A67*, a P450 suspected to  
28 be involved in *Medicago* saponin biosynthesis (Biazzi et al. 2015; Fukushima et al. 2013;  
29 Naoumkina et al. 2010) and *MtCYP72A68*, which was reported to catalyze the three-step  
30 oxidation of oleanolic acid in a heterologous yeast system (Fukushima et al. 2013).  
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38 Additional correlation analyses and gene expression clustering of the implicated P450  
39 genes were performed using data from the *M. truncatula* Gene Expression Atlas (Benedito et al.  
40 2008). The expression data used for the correlation data are provided in Supplemental Table S5.  
41 These analyses revealed similar clustering of the uncharacterized *MtCYP72A67* and partially  
42 characterized *MtCYP72A68* genes with the previously characterized *MtβAS*, *MtCYP716A12* and  
43 *MtCYP93E2* P450s (Supplemental Figure S3). *MtCYP72A67* and *MtCYP72A68* are also highly  
44 correlated with the accumulation of the oleanate sapogenin medicagenic acid. Therefore, the  
45 cumulative correlation data strongly support that both *MtCYP72A67* and *MtCYP72A68* have a  
46 high potential as putative genes/enzymes involved in triterpene sapogenin biosynthesis.  
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### 56 **Heterologous expression of *MtCYP72A67* in yeast**

57 The potential oleanate sapogenin oxidase activity of *MtCYP72A67* was tested using  
58 heterologous expression in yeast and *in vitro* biochemical assays using yeast microsomes. A  
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4 recent article reports on the identification of CYP72A67 (Fukushima et al. 2013); however, upon  
5 co-expression of the  $\beta AS$ , cytochrome P450 reductase (CPR), CYP716A12, and CYP72A67  
6 genes in yeast, they could not demonstrate CYP72A67 activity (Fukushima et al. 2013). In this  
7 study, we used microsomes from yeast WAT11 cells expressing *MtCYP72A67 in vitro* with a  
8 variety of triterpene sapogenin substrates, and the products were analyzed by UHPLC-(-)ESI-  
9 QToF-MS. NADPH was also added to these assays as a P450 cofactor. When oleanolic acid was  
10 used as a substrate, 2 $\beta$ -hydroxyoleanolic acid was detected as a product in the *MtCYP72A67* (+)  
11 NADPH microsomal samples, but not detected in the *MtCYP72A67* (-) NADPH or empty vector  
12 control samples (Table 1 and Supplemental Figure S4A). When hederagenin was used as a  
13 substrate, 2 $\beta$ -hydroxyhederagenin (*e.g.* bayogenin) was detected as a product in the (+) NADPH  
14 samples but not detected in the *MtCYP72A67* (-) NADPH and empty vector control samples  
15 (Table 1B and Supplemental Figure S4B). Both the empty vector control and assays without  
16 NADPH resulted in no P450 activity (Supplemental Figure S4B). In addition,  $\beta$ -amyrin and  
17 erythrodiol were used as substrates in *MtCYP72A67* assays, but no products were detected using  
18 gas chromatography-mass spectrometry (GC-MS; data not shown; a summary of all substrates  
19 that were used for *MtCYP72A67* yeast assay is listed in Supplemental Table S6). It was  
20 concluded that microsomes containing recombinant CYP72A67 protein possess multi-substrate  
21 C-2 $\beta$ -hydroxylase activity for oleanolic acid and hederagenin, yielding the C-2  $\beta$ -hydroxy  
22 derivatives.  
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40 Encouraged by the *in vitro* assays that clearly indicate the oleanolic acid oxidase activity  
41 of CYP72A67, additional yeast strains KM1 and KM2 were created from a sterol engineered  $\beta$ -  
42 amyrin producing yeast strain TM3 (Moses, Pollier, et al. 2014). All yeast strains generated in  
43 this study are listed in Supplemental Table S8. Both strains express *CYP716A12* and *M.*  
44 *truncatula* cytochrome P450 reductase (*MtCPR1*) from a high-copy number plasmid to produce a  
45 self-processing polyprotein in which the P450 reductase and the P450 are linked via a 2A  
46 oligopeptide (de Felipe et al., 2006). When cultivated in the presence of methylated  $\beta$ -  
47 cyclodextrins (M $\beta$ CD), both strains produce high levels of oleanolic acid (Figure 4A). Yeast  
48 strain KM2 also expresses *MtCYP72A67* from a high-copy number plasmid, whereas KM1 does  
49 not. Comparison of the GC chromatograms of extracts from the spent medium of KM1 and KM2  
50 cultured with M $\beta$ CD showed a single unique peak in strain KM2, but not in strain KM1, that  
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4 corresponds to 2 $\beta$ -hydroxyoleanolic acid (Figure 4A). In summary, MtCYP72A67 was shown to  
5 catalyze the C-2 $\beta$ -oxidation of oleanolic acid, using *in vitro* assays with yeast microsomes and  
6 engineered yeast strains.  
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### 9 10 **Ectopic expression and characterization of *MtCYP72A67* in *planta***

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12 To investigate if MtCYP72A67 functions as a 2 $\beta$ -hydroxylase *in planta*, *M. truncatula* hairy  
13 roots were generated following *Agrobacterium rhizogenes*-mediated transformation with the  
14 *Mtcyp72a67* RNAi, a double-stranded hairpin RNA that triggers post-transcriptional gene  
15 silencing, or with *MtCYP72A67*- overexpressing the *MtCYP72A67* full coding sequence.  
16  
17 Quantitative RT-PCR analysis of the *MtCYP72A67* transcript levels resulted in an average gene  
18 expression reduction of 46% in *Mtcyp72a67*-RNAi hairy roots compared to control. UHPLC(-  
19 )ESI-QToF-MS was used to compare the saponin content in transformed hairy roots relative to  
20 an empty vector control. Figure 5 summarizes the changes in saponin content which were  
21 measured in *Mtcyp72a67* RNAi hairy roots. Deduced enzymatic products of CYP72A67 and  
22 related downstream saponins, including 2 $\beta$ -hydroxyoleanolic acid, bayogenin, polygalagenin  
23 (putative identification), medicagenic acid and zanhic acid were significantly reduced while  
24 substrates oleanolic acid, hederagenin and gypsogenin (putative identification) were significantly  
25 increased in the *Mtcyp72a67* RNAi hairy roots compared to the control. In Figure 5, the oleanate  
26 saponin levels were altered in the *Mtcyp72a67* RNAi hairy roots where the downstream C-2  
27 oxidative derivatives of oleanolic acid were decreased while the non-C-2 oxidative pathway  
28 metabolites were increased.  
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43 *MtCYP72A67* was overexpressed in hairy roots and *MtCYP72A67* transcript levels  
44 showed an average 2.24-fold induction by qRT-PCR compared to the control. Saponin analyses  
45 were performed by UHPLC(-)ESI-QToF-MS for the *MtCYP72A67*-overexpressing hairy roots  
46 and compared to hairy roots transformed with an empty vector. *MtCYP72A67*-overexpressing  
47 hairy roots had significantly increased levels of several aglycones, including 2 $\beta$ -  
48 hydroxyoleanolic acid, polygalagenin (putative identification) and zanhic acid (Supplemental  
49 Figure S5). Unexpectedly, the level of oleanolic acid was also significantly increased which may  
50 indicate a more complex regulatory function of this gene. Ectopic expression of *MtCYP72A67* *in*  
51 *planta* is in agreement with the results from yeast and thus support the conclusion that  
52 MtCYP72A67 possesses multi-substrate 2 $\beta$ -hydroxylase activity for oleanolic acid and  
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4 hederagenin, yielding the C-2 alcohols in the oleanate sapogenin branch of the triterpene  
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6 saponins.

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8 In addition, *G. glabra*  $\beta$ AS and *MtCYP716A12* were transiently expressed with or without  
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10 *MtCYP72A67* in *Nicotiana benthamiana* leaves using *Agrobacterium tumefaciens*-mediated  
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12 infiltration. Similar to yeast, comparison of the GC-MS chromatograms of organic extracts from  
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14 leaves three days after co-infiltration revealed the presence of 2 $\beta$ -hydroxyoleanolic acid in leaves  
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16 that were co-infiltrated with *MtCYP72A67* (Figure 4B). Taken together, *MtCYP72A67* was  
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18 shown to catalyze the 2 $\beta$ -oxidation of oleanolic acid in heterologous tobacco bioassays.  
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### 20 **Heterologous expression of *MtCYP72A68* in yeast**

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22 The potential oleanate sapogenin oxidase activity of *MtCYP72A68* was first tested using  
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24 heterologous expression and *in vitro* enzymatic assays. Microsomes of WAT11 yeast cells  
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26 expressing *MtCYP72A68* were tested for *in vitro* enzymatic activity with various substrates, and  
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28 products analyzed with UHPLC-(-)ESI-QToF-MS or GC-MS as described above. Protein  
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30 activity was measured with oleanolic acid or hederagenin as the substrate and with or without  
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32 NADPH as a P450 cofactor. As shown in Table 2A, when oleanolic acid was used as a substrate,  
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34 hederagenin, gypsogenin, and gypsogenic acid accumulated. None of these products were  
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36 detected in the empty vector control samples (see also Supplemental Figure S6A). The detected  
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38 anion at  $m/z$  469.35 at retention time (RT) of 24.91 min was tentatively identified in this study as  
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40 gypsogenin or 3 $\beta$ -hydroxy-23-oxo-olean-12-en-28-oic acid based upon literature information  
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42 including accurate mass, aglycone anion at  $m/z$  469 and a predicted molecular formula of  
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44  $C_{30}H_{46}O_4$  (Supplemental Figure S6; <http://www.chemspider.com>).

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46 Another detected anion at  $m/z$  485.35 and RT of 21.88 min was previously identified as  
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48 gypsogenic acid or 2 $\beta$ ,3 $\beta$ -dihydroxy-23-oxo-olean-12-en-28-oic acid based upon fragmentation  
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50 of its aglycone anion at  $m/z$  485 and a predicted molecular formula of  $C_{30}H_{46}O_5$  (Supplemental  
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52 Figure S6; <http://www.chemspider.com> and (Pollier et al. 2011). In addition, the amount of  
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54 oleanolic acid detected was lower in the *MtCYP72A68* (+) NADPH assay, indicating its  
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56 consumption as a substrate. When hederagenin was used as a substrate (Table 2B), large  
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58 quantities of gypsogenin and gypsogenic acid were detected as products in the *MtCYP72A68*  
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60 assay and not detected in the empty vector control samples (see also Supplemental Figure S6B).  
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62 The amount of hederagenin detected was lower in the *MtCYP72A68* (+) NADPH assay,  
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4 indicating its consumption as a substrate.  $\beta$ -amyrin and erythrodiol were also tested as substrates,  
5 but no products were detected via GC-MS (data not shown; a summary of all substrates that were  
6 used for MtCYP72A68 yeast assays is listed in Supplemental Table S6). Taken together, these  
7 results indicate that microsomes containing recombinant MtCYP72A68 possess the ability to  
8 catalyze the sequential oxidation of C-23 on oleanolic acid, yielding the C-23 alcohol, aldehyde  
9 and carboxylic acid derivatives. The data also indicated that recombinant MtCYP72A68 will not  
10 accept saponin with C-28 methyl or C-28 hydroxyl groups as substrates.  
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18 Next, yeast strains KM1 and KM3 were created from a sterol engineered  $\beta$ -amyrin  
19 producing yeast strain TM3 (Moses, Pollier, et al. 2014). Yeast strain KM3 expresses a self-  
20 processing polyprotein in which CYP716A12 and MtCPR1 were linked via a 2A oligopeptide.  
21 KM3 also expresses *MtCYP72A68* from a high-copy number plasmid, whereas KM1 does not.  
22 Comparison of the GC-MS chromatograms of extracts from the spent medium of KM1 and KM3  
23 cultured with M $\beta$ CD showed three unique peak in strain KM3 that correspond to hederagenin,  
24 gypsogenin and gypsogenic acid (Figure 6A) (Fukushima et al. 2013).  
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### 34 **Ectopic expression and characterization of *MtCYP72A68* planta**

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36 *M. truncatula* hairy roots were transformed and *MtCYP72A68* overexpressed to further  
37 substantiate the role of *MtCYP72A68* in triterpene saponin biosynthesis. The *MtCYP72A68*  
38 transcript levels in hairy roots were quantified by qRT-PCR, and an average 1.51-fold induction  
39 in *MtCYP72A68* expression was observed compared to the control. Saponin analyses were  
40 performed by UHPLC-(-)ESI-QToF-MS on the *MtCYP72A68-OE* (overexpression) hairy roots  
41 and compared to hairy roots transformed with an empty vector. The fold changes in saponin  
42 content that were detected in *MtCYP72A68-OE* hairy roots are presented in Figure 7, and the  
43 data revealed significantly increased levels of several aglycones, including oleanolic acid,  
44 hederagenin, polygalagenin (putative identification), soyasapogenin E and total saponins. None  
45 of the metabolites were decreased. Unexpectedly, the level of 2 $\beta$ -hydroxy oleanolic acid was  
46 also induced. As shown in Figure 7, both oleanate saponins and soyasapogenols branches were  
47 altered in the *MtCYP72A68-OE* hairy roots, which affected the total accumulation of saponins by  
48 1.53-fold. *M. truncatula* hairy roots were also transformed with *Mtcyp72a68* RNAi, a double-  
49 stranded RNA (hairpin RNA). However, transcript levels of *Mtcyp72a68*-RNAi were checked by  
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4 qRT-PCR and showed only a minor average decrease in gene expression of 16% and no  
5 significant metabolic changes. *Glycyrrhiza glabra*  $\beta$ AS, *MtCPR1* and *MtCYP716A12* were also  
6 transiently expressed with or without *MtCYP72A68* in *Nicotiana benthamiana* leaves using  
7 *Agrobacterium tumefaciens*-mediated infiltration. Like in yeast, comparison of the  
8 chromatograms of organic extracts from leaves three days after co-infiltration revealed three  
9 unique peaks corresponding to hederagenin, gypsogenin and gypsogenic acid in leaves that were  
10 co-infiltrated with *MtCYP72A68* (Figure 6B). Hence, heterologous expression of *MtCYP72A68*  
11 in yeast and tobacco all point towards the sequential oxidation of C-23 of oleanolic acid by  
12 *MtCYP72A68* and confirm similar results obtained by (Fukushima et al. 2013).  
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22 We also measured the saponin levels extracted from root tip tissues from Tnt1  
23 retrotransposon mutants (Tadege et al. 2008) in the *MtCYP72A68* gene. Two independent mutant  
24 lines with retrotransposon insertions in the *MtCYP72A68* gene were identified through PCR  
25 screening (Tadege et al. 2008). Tnt1 insertion lines NF1698-4, NF12169, and R108 (control)  
26 were germinated and the root tips (approximately 3 mm) from 25 plants (a mixed population of  
27 heterozygous and wild type, 2:1) were collected. The mRNA levels revealed that both Tnt1  
28 mutant lines possessed lower *MtCYP72A68* mRNA levels, 22 % lower for NF1698-4 and 20%  
29 lower for NF12169, compared to wildtype R108 control (Supplemental Figure S7A). This was  
30 followed by UHPLC-(-)ESI-QToF-MS analyses of saponin aglycones (Supplemental Figure  
31 S7B). These analyses showed reduction of medicagenic acid and zanhic acid of NF12169 line.  
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## 41 **DISCUSSION**

### 42 **Genomic and coexpression analyses identify genes involved in triterpene saponin** 43 **biosynthesis** 44

45 Currently, several P450s from multiple P450 families have been reported in relation to saponin  
46 biosynthesis (Augustin et al. 2011; Miettinen et al. 2018; Seki et al. 2015). Diversity of P450s  
47 are involved in triterpene saponin biosynthesis across many species, hence the prediction of  
48 specific substrate and enzymatic activity based on sequence alone is complex (Nelson and  
49 Werck-Reichhart 2011). Thus, there is a need for large-scale data approaches to identify and  
50 prioritize candidate P450s and other gene candidates involved in triterpene biosynthesis.  
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57 Germplasm collections are powerful resources for exploring the natural variation for any number  
58 of phenotypes (Ronfort et al. 2006), including saponin content. We measured the total saponin  
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4 content for 106 of *M. truncatula* ecotypes and revealed substantial differential accumulation of  
5 these specialized metabolites. Four diverse ecotypes with strong differential accumulation of  
6 saponins in root and aerial tissues were selected for further transcriptome analyses. Correlation  
7 analyses between gene-gene expression and gene-metabolite accumulation were then performed  
8 to identify genes that are likely involved in triterpene saponin biosynthesis. The correlation  
9 coefficients between microarray gene expression levels for putative P450s,  $\beta$ AS, known  
10 sapogenins and total saponins revealed a significantly correlated cluster of genes, including  
11 *MtCYP72A67*, *MtCYP72A68*, *MtCYP716A12* and *Mt $\beta$ AS*. Saponin levels, including total known  
12 saponins, total aglycones and medicagenic acid, were also clustered (Figure 3 and Supplemental  
13 Figure S3). These correlations represent discovery events that identify putative saponin  
14 biosynthetic genes. To further assess the CYP72 candidate genes, correlation coefficients for  
15 gene-gene expression values of the P450s were calculated using the *M. truncatula* Gene  
16 Expression Atlas (Supplemental Figure S3) (Benedito et al. 2008; He et al. 2009). This revealed  
17 similar clustering of *MtCYP72A67* and *MtCYP72A68* genes with *Mt $\beta$ AS* and *MtCYP716A12*  
18 (Figure 3). Both *Mt $\beta$ AS* and *MtCYP716A12* genes have been previously reported as saponin  
19 biosynthetic enzymes (Supplemental Figure S3) (Carelli et al. 2011; Fukushima et al. 2011;  
20 Miettinen et al. 2017). *MtCYP72A61*, *MtCYP72A67* and *MtCYP72A68* were also implicated  
21 based upon their coexpression with  $\beta$ -amyrin synthase in methyl jasmonate-elicited cell culture  
22 data (Naoumkina et al. 2010) and their expression is under control of the regulator of hemolytic  
23 saponin biosynthesis, the transcription factor TSAR2 (Mertens et al. 2016). Thus, *MtCYP72A67*  
24 and *MtCYP72A68* were prioritized for functional characterization.

#### 25 **MtCYP72A67 is a C-2 hydroxylase involved in the biosynthesis of oleanate sapogenins**

26 Heterologous combinatorial biosynthesis is a method that establishes novel enzyme-substrate  
27 combinations *in vivo* (Pollier et al. 2011). However, heterologous combinatorial expression of  
28  $\beta$ AS/CPR/CYP716A12/CYP72A67 genes in yeast strains did not demonstrate CYP72A67  
29 activity (Fukushima et al. 2011), whereas in this study, we were able to demonstrate the 2 $\beta$ -  
30 hydroxylase activity of MtCYP72A67 on oleanolic acid. This apparent discrepancy could be due  
31 to the difference in the yeast strains used or due to different culturing conditions. In the study of  
32 Fukushima et al. (2013), the non-engineered yeast strain INVSc1 was used, whereas for this  
33 study we used the sterol-engineered yeast strain TM3. In strain TM3, the native yeast gene

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4 ERG7 is under control of a methionine-repressible promoter allowing for increased accumulation  
5 of 2,3-oxidosqualene, the substrate for  $\beta$ -amyrin synthase. In addition, strain TM3 also expresses  
6 a truncated feedback-insensitive copy of isoform 1 of the *S. cerevisiae* 3-hydroxy-3-  
7 methylglutaryl-CoA reductase (tHMG1) gene, allowing for an increased accumulation of 2,3-  
8 oxidosqualene (Kirby et al. 2008; Moses, Pollier, et al. 2014). Furthermore, the use of M $\beta$ CD in  
9 the cultivation process improves the catalytic efficiency of the P450s, likely due to removal of  
10 feedback inhibition on the P450 activity or removal of toxicity due to lower intracellular  
11 accumulation of sapogenins (Moses, Pollier, et al. 2014).  
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19 Heterologous MtCYP72A67 catalyzed the oxidation of C-2 of both oleanolic acid and  
20 hederagenin, yielding the products 2 $\beta$ -hydroxy oleanolic acid and bayogenin, respectively (Table  
21 1). The hydroxylation of hederagenin indicates that compounds with C-23 hydroxyl substitution  
22 are also substrates for MtCYP72A67-mediated C-2 oxidation. MtCYP72A67 P450-mediated C-2  
23 oxidation activity is also supported by the lack of product accumulation in assays deficient in  
24 NADPH (Supplemental Figure S4). Lack of product accumulation in these assays indicates that  
25 the cytochrome P450 reductase/MtCYP72A67 requires NADPH as an electron donor for activity  
26 (Liu et al. 2003; Seki et al. 2008). No products were detected when MtCYP72A67 was assayed  
27 with  $\beta$ -amyrin or erythrodiol, which implies that compounds with a C-28 methyl ( $\beta$ -amyrin) or  
28 C-28 hydroxyl group (erythrodiol) are not suitable substrates for MtCYP72A67-mediated C-2  
29 oxidation. Functional genomics studies of *Mtcyp72a67* RNAi and *MtCYP72A67* overexpression  
30 in *M. truncatula* hairy roots further demonstrated that MtCYP72A67 possessed *in planta*  
31 oxidation activity for C-2 of oleanolic acid and hederagenin, yielding the C-2 alcohols in  
32 catalyzed substrates of the oleanate sapogenin branch of the triterpene saponins (Figure 5 and  
33 Supplemental Figure S5).  
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#### 48 **MtCYP72A68 is a multifunctional oxidase involved in the oleanate sapogenin** 49 **biosynthesis** 50

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52 Previous studies have shown that an individual cytochrome P450 can catalyze the oxidation of a  
53 given carbon, yielding the hydroxyl, carbonyl and carboxylic acid products related to diterpenes  
54 in loblolly pine and Arabidopsis (He et al. 2009; Ro et al. 2005). More recently, a  
55 multifunctional oxidase involved in triterpene saponin biosynthesis (MtCYP716A12) has also  
56 been identified in *M. truncatula* (Carelli et al. 2011). Both heterologous expression assays in  
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4 yeast and tobacco demonstrate that MtCYP72A68 catalyzes the initial oxidation of C-23 of  
5 oleanolic acid, yielding the C-23 hydroxyl product hederagenin (Table 2, Supplemental Figure  
6 S6). Additional products with  $m/z$  values of 469.35 and 485.35 were also detected in the  
7  
8 MtCYP72A68 oleanolic acid assays. These have been tentatively identified as gypsogenin, the  
9 C-23 aldehyde derivative of oleanolic acid, and gypsogenic acid, the C-23 carboxy derivative of  
10 oleanolic acid, based upon accurate mass and literature information. Assays of yeast expressing  
11 *MtCYP72A68* tested with hederagenin also showed production of gypsogenin and gypsogenic  
12 acid products. No products were detected when MtCYP72A68 was assayed with  $\beta$ -amyrin or  
13 erythrodiol, which implies that compounds with a methyl C-28 ( $\beta$ -amyrin) or C-28 hydroxyl  
14 group (erythrodiol) are not suitable substrates for MtCYP72A68-mediated C-23 oxidation.  
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24 The *in planta* study of *MtCYP72A68* overexpressed in *M. truncatula* hairy roots  
25 supported the oxidase activity of CYP72A68 for oleanolic acid, yielding the C-23 alcohol in  
26 catalyzed substrates of the oleanate sapogenin branch of the triterpene saponins (Figures 8). Tnt1  
27 mutant lines with an insertion in the *MtCYP72A68* gene accumulated high levels of 2 $\beta$ -hydroxy  
28 oleanolic acid due to the reduced level of *MtCYP72A68* (Supplemental Figure S7). However, the  
29 reason for high induction of 2 $\beta$ -hydroxy oleanolic acid in *MtCYP72A68* overexpressing lines is  
30 not clear and may be due to more complex regulation of the pathway or regulatory function of  
31 the *CYP72A68* gene. Taken together, these results indicate that MtCYP72A68 catalyzes the  
32 oxidation of C-23 of oleanate sapogenins, yielding the alcohol (hederagenin), and likely also  
33 catalyzes the further oxidation towards the aldehyde (gypsogenin) and carboxy acid (gypsogenic  
34 acid).  
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#### 44 **Multisubstrate enzymes enhance chemical diversity**

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46 *MtCYP72A67* and *MtCYP72A68* are novel enzymes that catalyze the sapogenin biosynthesis in  
47 *M. truncatula*. MtCYP72A67 is a multisubstrate C-2 oxidase yielding 2 $\beta$ -hydroxy oleanolic acid  
48 and bayogenin, and MtCYP72A68 is a multisubstrate C-23 multifunctional oxidase yielding  
49 hederagenin, gypsogenin and gypsogenic acid. Although gypsogenin was not detected *in planta*,  
50 it was detected in yeast heterologously producing MtCYP72A68 and supplied *in vivo* or *in vitro*  
51 with oleanolic acid and hederagenin acid (Table 2, Figures 7-8 and Supplemental Figure S6-S7).  
52 We suggest that *MtCYP72A68* has the potential to convert the C-23 alcohol of hederagenin to a  
53 carbonyl group leading to gypsogenin. Together these genes have the potential to synthesize at  
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4 least four different saponin precursors from a common precursor, oleanolic acid. Biosynthesis of  
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6 bayogenin, for instance, requires both genes for oxidation of C-2 and C-23 (Figure 1). However,  
7  
8 the order of these reactions has not yet been determined. Evidence for cytochrome P450 enzymes  
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10 as multisubstrate enzymes in biochemistry has been accumulating for some time (Carelli et al.  
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12 2011) Siminszky et al., 1999; Schmidt et al., 2003; Ro et al., 2005). Enzymes with broad  
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14 substrate tolerance are also commonly found in natural product biosynthesis. Enzymes that can  
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16 act on more than one substrate to give multiple products is a mechanism that generates chemical  
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18 diversity, and, as long as one of the products enhances the fitness of the producer, the genes  
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20 coding for the overall process will be favored by selection, and chemical diversity will be  
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22 retained (Firn and Jones, 2003; Weissman and Leadley, 2005; Gershenzon and Dudareva, 2007).

## 23 CONCLUSIONS

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25 We exploited the genetic and biochemical diversity of a *M. truncatula* population and used  
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27 integrated metabolomics and transcriptomics to identify novel genes involved in saponin  
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29 biosynthesis. This was achieved through UHPLC(-)ESI-QToF-MS metabolite profiling of a  
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31 diverse collection of *M. truncatula* ecotypes, which further resulted in the identification of four  
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33 specific ecotypes with substantial differential saponin accumulation. Correlated gene-to-gene  
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35 expression and gene-to-metabolite accumulation data identified *MtCYP72A67* and *MtCYP72A68*  
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37 as potential oxidative enzymes associated with saponin biosynthesis. These genes were then  
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39 functionally characterized using traditional *in vitro*, *in vivo* and *in planta* biochemical assays  
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41 along with genetic approaches to prove function. The data provide evidence that *MtCYP72A67*  
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43 is a C-2  $\beta$ -hydroxylase of oleanolic acid and bayogenin, and that *MtCYP72A68* is a  
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45 multisubstrate, C-23 multifunctional oxidase of hederagenin, gypsogenin, and gypsogenic acid.  
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47 The successful integration of metabolomics and transcriptomics illustrated here provides  
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49 additional evidence of the value of these exciting new technologies in the discovery and  
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51 characterization of novel specialized metabolism genes.

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53  
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55  
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57  
58 thank Joseph Chappell for providing the WAT11 yeast strain. We thank Dr. Qiao Zhao for  
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4 **SUPPLEMENTAL DATA**  
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6 **Supplemental Table S1:** Metabolomics Standards Initiative Compliant Metadata supporting the  
7 experiments reported here (Fiehn et al. 2007).  
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10 **Supplemental Table S2:** Full ecotype UHPLC-ESI(-)-QToFMS data of *Medicago truncatula*  
11 aerial tissue  
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13 **Supplemental Table S3:** Full ecotype UHPLC-ESI(-)-QToF-MS data for *Medicago truncatula*  
14 root tissue  
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17 **Supplemental Table S4:** Relative saponin accumulation levels and gene expression data for  
18 selected *M. truncatula* lines (Log10).  
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21 **Supplemental Table S5:** Expression values from Medicago gene expression atlas of terpene  
22 P450s genes, unknown P450s and other genes from the terpenes biosynthesis which were used  
23 for CE heat map.  
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26 **Supplemental Table S6:** Substrates that were tested in CYP72A67 and CYP72A68 activity  
27 assay  
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30 **Supplemental Table S7:** Primer list used for qRT-PCR.  
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32 **Supplemental Table S8:** Yeast strains generated in this study and primers used for cloning.  
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34 **Supplemental Table S9:** Primer list of *MtCYP72A67* and *MtCYP72A68* cloning to hairy roots  
35 transformation.  
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38 **Supplemental Figure S1.** Relative saponin accumulation relative to gene expression levels of  
39 *MtCYP72A67*, *MtCYP72A68*, *MtCYP716A12* and  $\beta AS$  genes. Root and aerial tissues were  
40 collected from four selected ecotypes: (A) A17; (B) ESP105; (C) GRC43; and (D) R108.  
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Microarray analyses and UHPLC(-)ESI-QToF-MS were performed using three biological replicates. Expression values are reported as Log10 of shoots (black bars) and root (white bars).

**Supplemental Figure S2.** Gene expression of *MtCYP72A67*, *MtCYP72A68*, *MtCYP716A12* and  $\beta AS$ . Root and aerial tissues were collected from 5-week-old *M. truncatula* ecotypes, including ESP105, GRC43, A17, and R108. Relative gene expression was measured and reported for two methods: A) qRT-PCR; and B) microarray analyses. In both methods, three biological replicates were used.

**Supplemental Figure S3.** Correlation coefficient (Pearson's r) heat map of known, unknown P450s and genes from the terpene biosynthesis using expression values from Gene Expression

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4 Atlas. Transcript levels were measured in the different tissues (microarray data were obtained  
5 from the *M. truncatula* Gene Expression Atlas database version 2, MtGEAv2,  
6 <https://mtgea.noble.org/v2/>). The P450s involved in triterpene biosynthesis were selected  
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8 according to a previous paper (Naoumkina et al. 2010). Upper left triangle matrix presents probe  
9 set correlation coefficients using Pearson's correlation (scale -0.5 to 1), and lower triangle matrix  
10 presents *P* value of the correlation coefficient test (light blue present *P* value < 0.01 and brown  
11 present non-significant). The black-bordered squares illustrated a highly correlated set of genes,  
12 including *MtCYP72A67*, *MtCYP72A68*, *MtCYP716A12*, *βAS*, *MtCYP93E2*, *isopentenyl*  
13 *pyrophosphate isomerase*, *phosphomevalonate kinase* and *mevalonate kinase*.  
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21 **Supplemental Figure S4.** UHPLC-(-)ESI-QToF-MS chromatograms of the MtCYP72A67  
22 product generated yeast *in vitro* enzymatic assays. (A) Chromatogram of activity assay using  
23 oleanolic acid and (B) hederagenin. The full-length MtCYP72A67 was tested with cofactor  
24 (MtCYP72A67 (+) NADPH), without cofactor (MtCYP72A67 (-) NADPH) and empty vector  
25 with cofactor (empty vector (+) NADPH). The results illustrate that MtCYP72A67 hydroxylates  
26 the C-2 position of oleanolic acid and hederagenin to produce 2-hydroxyoleanolic acid and  
27 bayogenin.  
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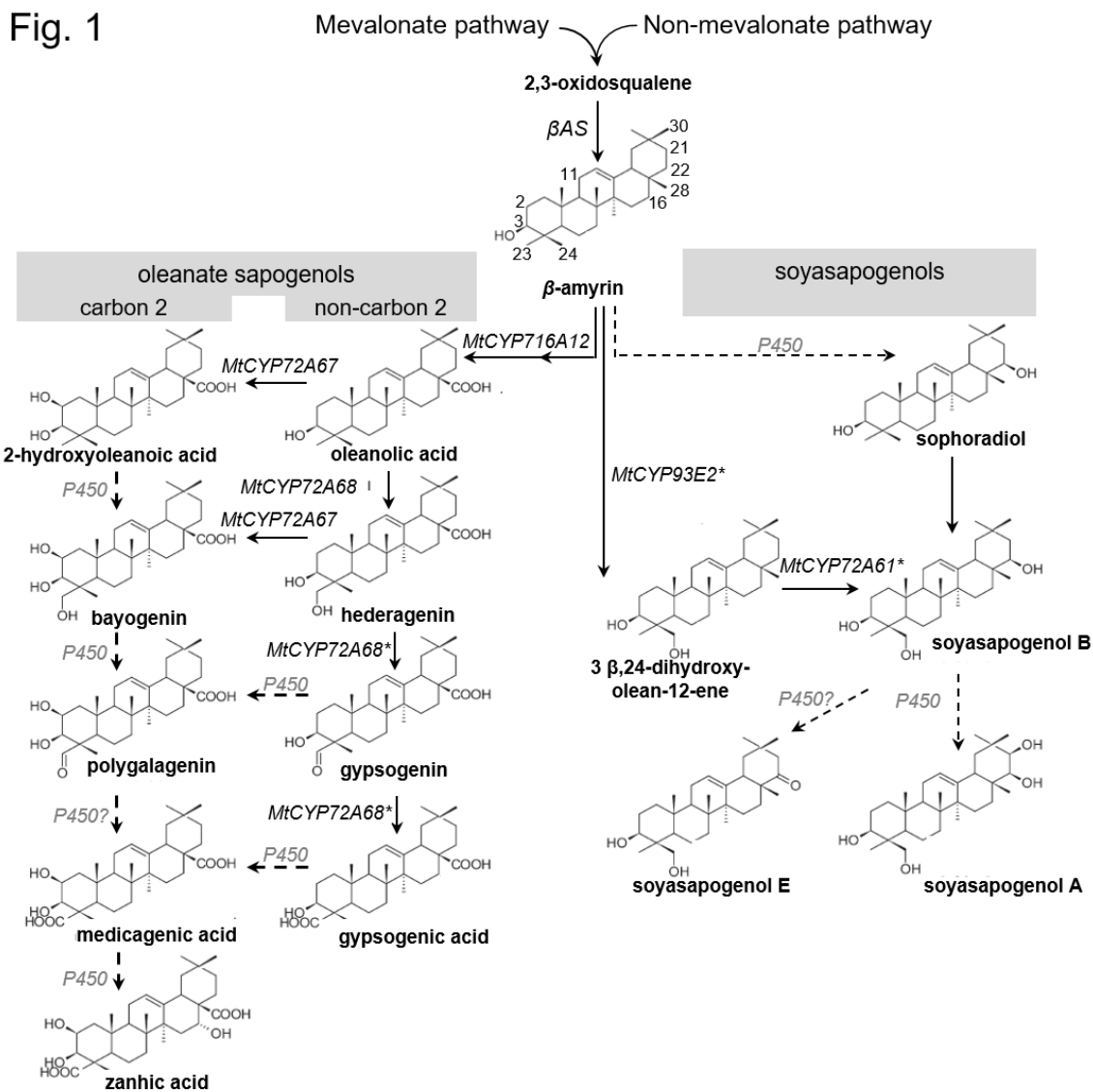
35 **Supplemental Figure S5.** Results of overexpression (OE) of *MtCYP72A67* full gene in *M.*  
36 *truncatula* hairy roots. (A) Proposed biosynthetic pathway of the saponin with observed  
37 metabolite fold changes in *M. truncatula* hairy roots. The saponin names are marked in  
38 different colors according to the fold changes: red – saponin significant fold increase; black – no  
39 change; and gray – not detected. The observed saponin fold changes are noted in brackets.  
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44 \*Gypsogenin and polygypsogenin were putatively identified using tandem mass (no authentic  
45 standard).  
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49 **Supplemental Figure S6.** *In vitro* enzymatic assays of recombinant MtCYP72A68 in yeast  
50 WAT11 cells. (A) UHPLC-(-)ESI-QToF-MS chromatograms of activity assays using oleanolic  
51 acid and (B) hederagenin as substrates. The full-length CYP72A68 was tested with cofactor  
52 (CYP72A68 (+) NADPH) and empty vector with cofactor (empty vector (+) NADPH).  
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54 Gypsogenin (3-hydroxy-23-oxoolean-12-en-28-oic acid) was tentatively identified here based  
55 upon literature information including accurate mass, aglycone anion at *m/z* 469 and a predicted  
56 molecular formula of C<sub>30</sub>H<sub>46</sub>O<sub>4</sub>. Gypsogenic acid, ((3β)-3-hydroxyolean-12-ene-23,28-dioic  
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4 acid) was previously identified from fragmentation of saponins yielding an aglycone anion at  $m/z$   
5 485 and a predicted molecular formula of  $C_{30}H_{46}O_5$  (<http://www.chemspider.com> and (Pollier et  
6 al. 2011)).  
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10 **Supplemental Figure S7.** Metabolite accumulation and qRT-PCR levels in *cyp72a68* Tnt1  
11 mutant lines. Metabolites and mRNA were extracted from 3mm root tips of 3-day-old seedlings  
12 with three biological repeats from two Tnt1 mutant lines, NF1698-4 and NF12169. (A) The  
13 relative level of *CYP72A68* gene expression detected by qRT-PCR in NF1698-4 and NF12169.  
14 Expression values were normalized relative to the endogenous ubiquitin control gene. (B) Fold  
15 changes in saponin content observed in the two mutant lines NF1698-4 and NF12169. In bold,  
16 Student's *t*-test with *P* value < 0.05.  
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FIGURES and LEGENDS



**Figure 1.** Proposed biosynthetic pathway of triterpene sapogenins reported in *Medicago* spp. (modified from Pollier et al., (2011) *J. Natural Products*). Solid arrows mark the enzymatic steps which have been characterized: *MtCYP72A68* and *MtCYP72A67* in the current paper in addition to *Mt* $\beta$ AS (Suzuki et al. 2002), *MtCYP716A12* (Carelli et al. 2011)(Fukushima et al. 2011); Seki et al., 2011), *MtCYP93E2* (Fukushima et al. 2011), *MtCYP72A68\** (Fukushima et al. 2013). Dashed arrows mark the unknown enzymes. Asterisk marks enzyme activity that has been tested in yeast system only (not *in planta*).

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Fig. 2

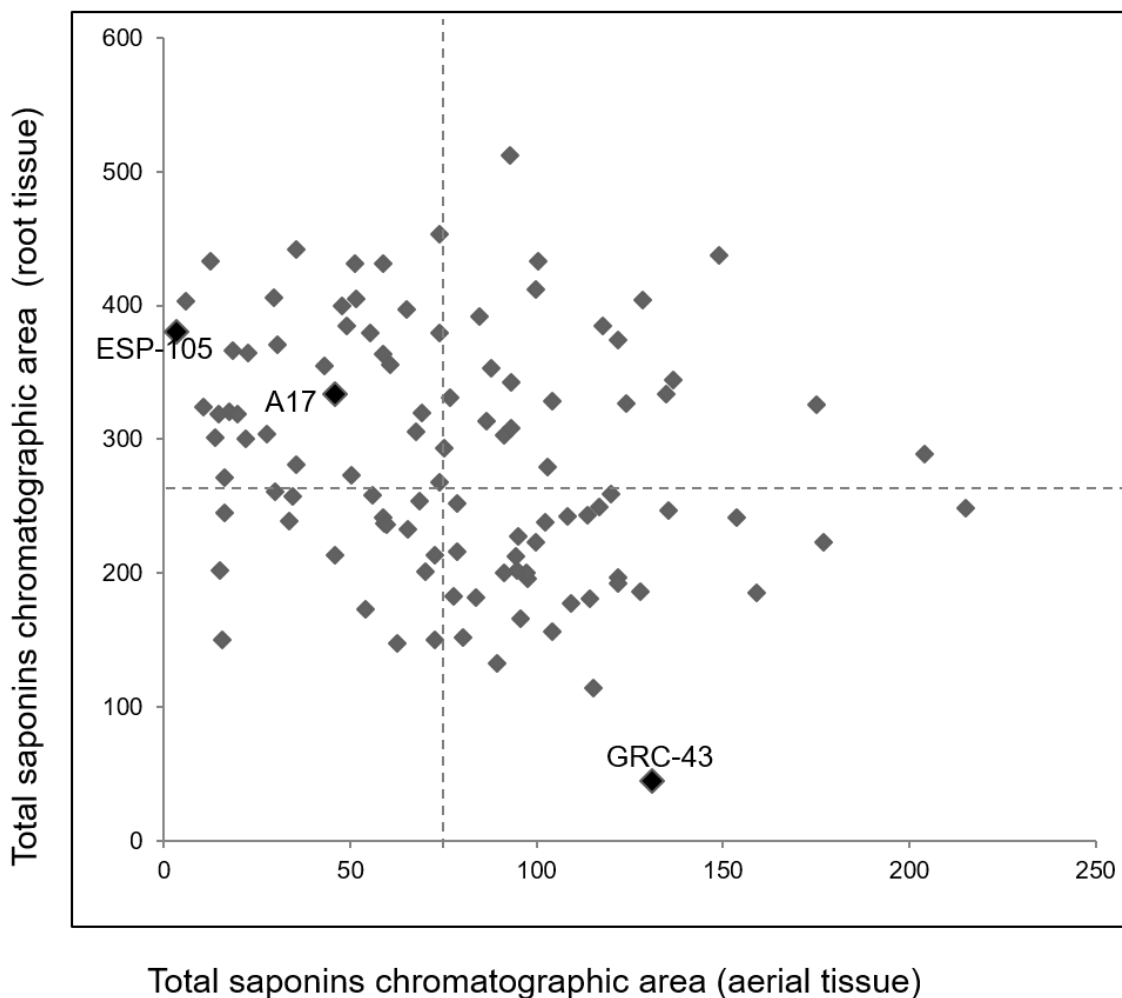
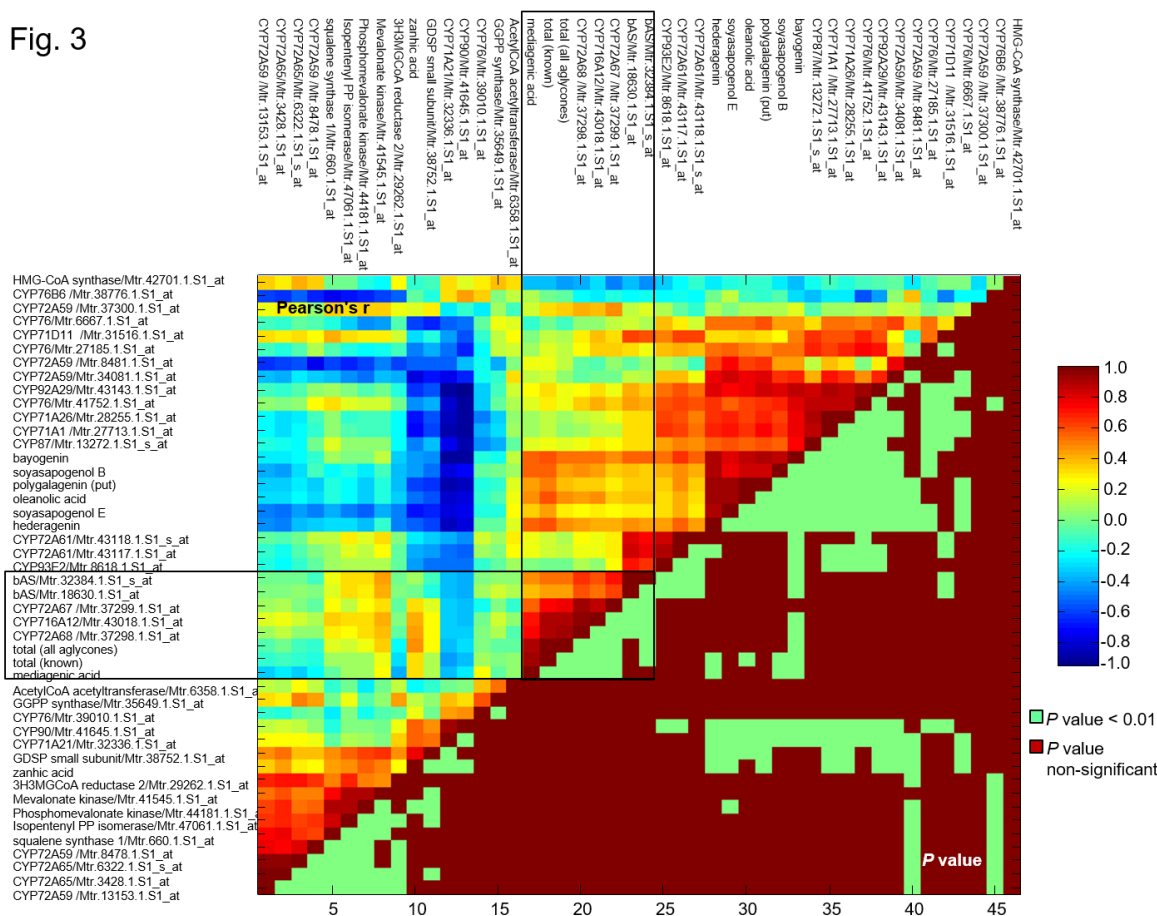


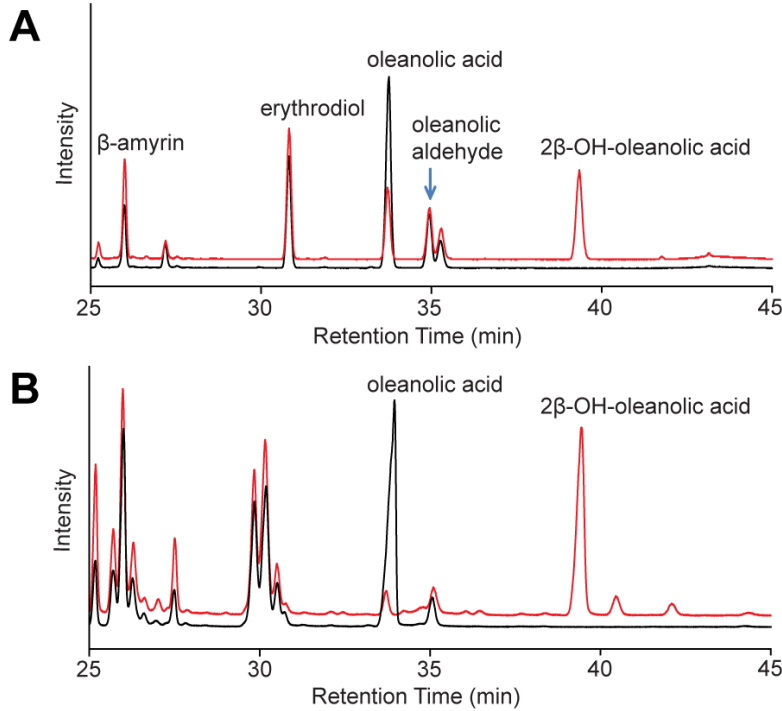
Figure 2. Scatter plot of total saponin content in root and aerial tissue measured for a germplasm collection of 106 *M. truncatula* ecotypes. Saponin content was quantified using UHPLC(-)ESI-QToF-MS total chromatographic peak area and measured for 5-week-old root and aerial tissues (three to four biological replicates for each ecotype tissue). Dashed line, average values of the X = 78.32 and Y = 285.35 total saponin chromatogram peak areas. ESP-105 had relatively high levels of saponins in roots but low levels in aerial tissues, and vice versa for GRC-43. Ecotype A-17 was used as a reference line.



Fig. 3

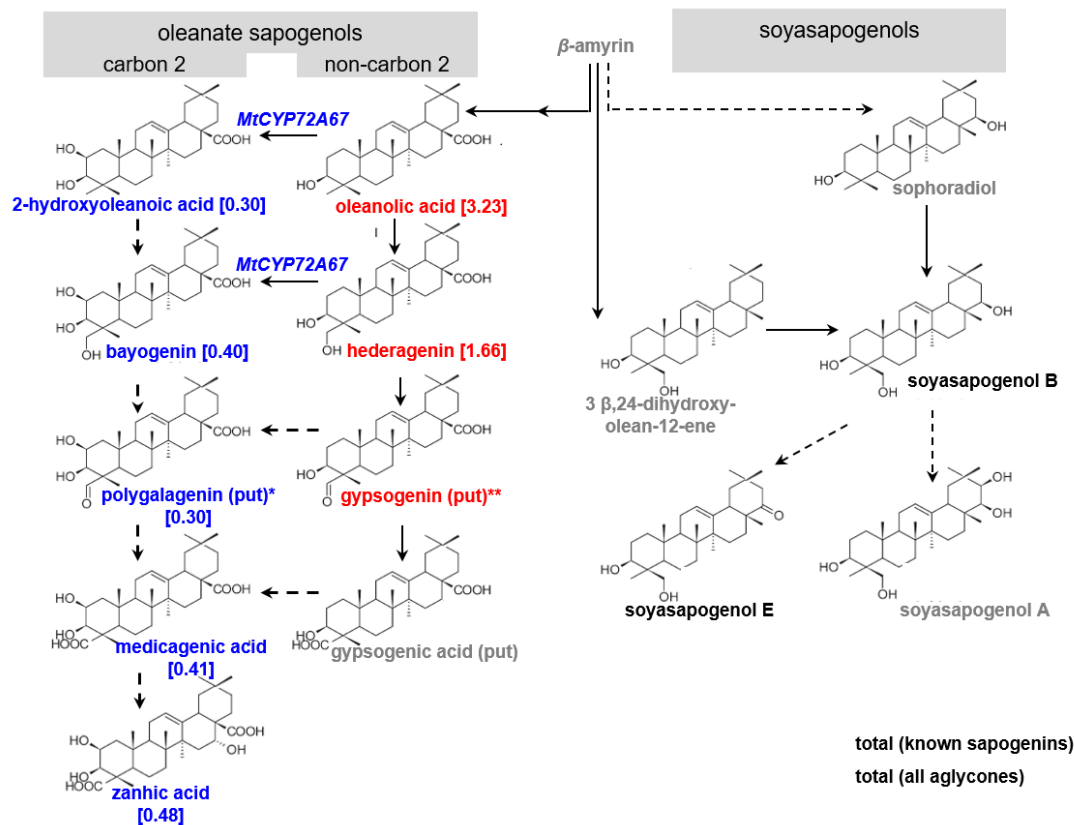


**Figure 3.** Correlation coefficient Pearson's  $r$  heat map of sapogenin metabolites and putative triterpene P450 transcripts. Metabolites and transcript levels were measured in root and aerial tissues of four selected *M. truncatula* ecotypes, A17, ESP105, GRC43 and R108 (three biological replicates each). Characterized and putative P450s involved in terpene biosynthesis were selected according to a previous report (Naoumkina et al. 2010) in addition to  $\beta$ AS and nine other known *M. truncatula* genes involved in early terpene biosynthesis. Upper left triangle matrix presents correlation coefficient Pearson's  $r$  as determined using Pearson's correlation (scale -0.8 to 1), and lower right triangle matrix presents  $P$ -values of the correlation coefficient test (light green highlights  $P$ -values  $< 0.01$ , and brown highlights non-significant correlation). The black-bordered boxes encapsulate highly correlated probe sets including CYP72A67, CYPA68, CYP716A12 and  $\beta$ AS, and metabolites medicagenic acid, total saponins, and total aglycones.



**Figure 4.** *In vivo* functional analysis of CYP72A67. Overlay of GC-MS chromatograms showing accumulation of trimethylsilylated enzymatic products. (A) *S. cerevisiae* strain KM1 expressing Gg $\beta$ AS, MtCPR1 and CYP716A12 (black) and *S. cerevisiae* strain KM2 expressing Gg $\beta$ AS, MtCPR1, CYP716A12 and CYP72A67 (red). (B) *N. benthamiana* co-infiltrated with *A. tumefaciens* strains armed with gene silencing suppressor p19, Gg $\beta$ AS and CYP716A12 (black) and *N. benthamiana* co-infiltrated with *A. tumefaciens* strains armed with gene silencing suppressor p19, Gg $\beta$ AS, CYP716A12 and CYP72A67 (red). CYP72A67v2 (version 2) was used.

Fig. 5



**Figure 5.** Characterization of *Mtcyp72a67*-RNAi in *M. truncatula* hairy roots. (A) Proposed sapogenin biosynthesis pathway in *M. truncatula* hairy root and observed fold changes noted in brackets. The sapogenin names are marked in different colors, according to the fold changes: red – saponin significant fold increase; blue – significant fold decrease; black – no change; and gray – not detected. Sapogenins were identified based upon authentic standards, except gypsogenin, gypsogenic acid and polygalagenin which were putatively identified using tandem mass. Gypsogenin (put) was detected only in the *Mtcyp72a67*-RNAi hairy root transgenic lines but not in the wild type.

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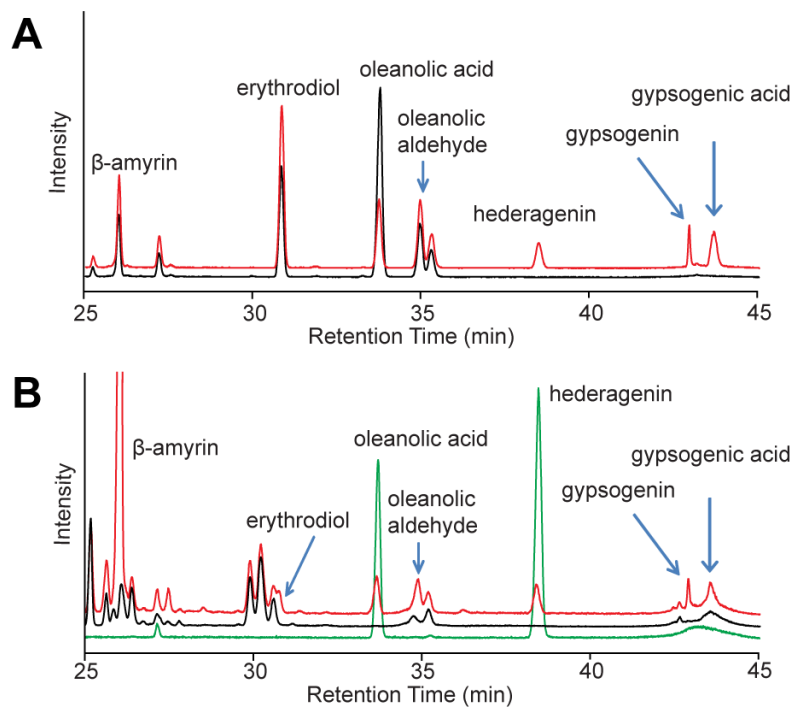
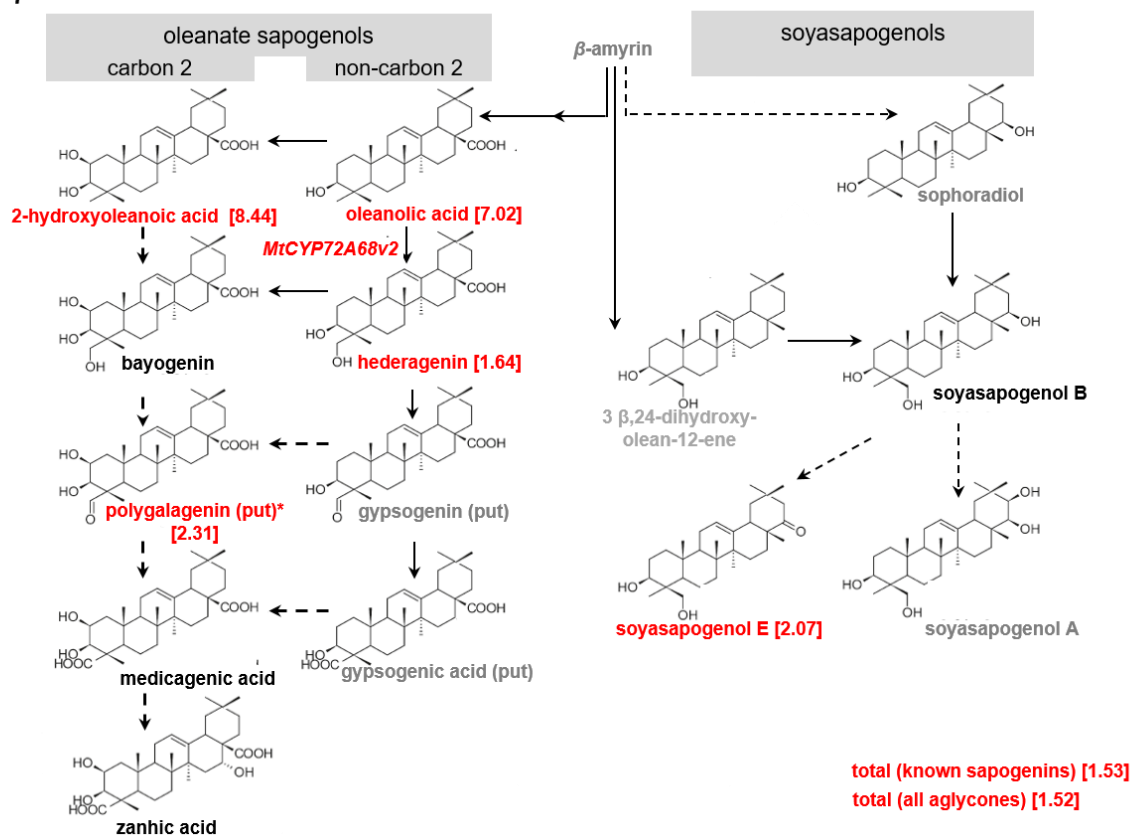


Figure 6. In vivo functional analysis of CYP72A68. Overlay of GC-MS chromatograms showing accumulation of trimethylsilylated enzymatic products in (A) *S. cerevisiae* strains KM1 expressing Gg $\beta$ AS, MtCPR1 and CYP716A12 (black) and KM3 expressing Gg $\beta$ AS, MtCPR1, CYP716A12 and CYP72A68 (red). (B) *N. benthamiana* infiltrated with an *A. tumefaciens* strain armed with gene silencing suppressor p19 (black) and *N. benthamiana* co-infiltrated with *A. tumefaciens* strains armed with gene silencing suppressor p19, Gg $\beta$ AS, CYP716A12 and MtCYP72A68. Also shown are oleanolic acid and hederagenin standards (green). CYP72A68v2 (version 2) was used.

Fig. 7



**Figure 7.** Characterization of *MtCYP72A68* overexpression in *M. truncatula* hairy roots. (A) Proposed sapogenin biosynthesis pathway in *M. truncatula* hairy root and observed fold changes noted in brackets. The sapogenin names are marked in different colors, according to the fold changes: red – saponin significant fold increase; black – no change; and gray – not detected. Sapogenins were identified based upon authentic standards, except gypsogenin, gypsogenic acid, and polygalagenin which were putatively identified using tandem mass.

**Table 1.** *In vitro* enzymatic assays of CYP72A67. Values in the table represent the mean, normalized chromatogram peak areas and standard error for each of the metabolites (three biological replicates per assay condition). The substrates that were used for CYP72A67 protein were (A) oleanolic acid and (B) hederagenin.

A) Substrate: oleanolic acid

Name	RT	m/z	CYP72A67 (+) NADPH			empty vector			fold change CYP72A67 / empty vector	P value
				±			±			
oleanolic acid	28.63	455.35	44866.7	±	2321.2	49566.7	±	1026.9	0.91	1.4E-01
2-hydroxyoleanoic	26.36	471.35	97000.0	±	3507.6	N.D.	±		high	

B) Substrate: hederagenin

Name	RT	m/z	CYP72A67 (+) NADPH			empty vector			fold change CYP72A67 / empty vector	P value
				±			±			
hederagenin	22.76	471.35	42033.3	±	1299.1	47966.7	±	1026.9	0.88	2.4E-02
bayogenin	20.28	487.34	45433.3	±	448.5	N.D.	±		high	

**Table 2.** *In vitro* enzymatic assay of CYP72A68 protein. Values in the table represent the mean, normalized chromatogram peak areas and standard error for each of the metabolites (three biological replicates per assay condition). The substrates that were used for CYP72A68 protein were (A) oleanolic acid and (B) hederagenin.

A) Substrate: oleanolic acid

Name	RT	m/z	CYP72A68 (+) NADPH		empty vector			fold change CYP72A68 / empty vector	P value
oleanolic acid	28.80	455.35	218673.0	± 12616.4	328189.4	±	6112.1	0.67	1.7E-04
hederagenin	22.96	471.35	175971.8	± 10435.2	N.D.			high	N.D.
gypsogenin (put)	24.91	469.33	258803.0	± 18852.2	N.D.			high	N.D.
gypsogenic acid (put)	21.88	485.33	156.8	± 39.1	N.D.			high	N.D.

B) Substrate: hederagenin

Name	RT	m/z	CYP72A68 (+) NADPH		empty vector			fold change CYP72A68 / empty vector	P value
hederagenin	22.96	471.35	227765.9	± 12260.6	250615.2	±	4084.8	0.91	3.8E-02
gypsogenin (put)	24.91	469.33	119954.8	± 16918.1	N.D.			high	N.D.
gypsogenic acid (put)	21.89	485.55	762.9	± 221.0	N.D.			high	N.D.

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**Compliance with Ethical Standards:** All authors including Vered Tzin, John H. Snyder, Dong Sik Yang, David V. Huhman, Bonnie S. Watson, Stacy N. Allen, Yuhong Tang, Karel Miettinen, Philipp Arendt, Jacob Pollier, Alain Goossens, and Lloyd W. Sumner declare no conflict of interest that could have direct or potential influence or impart bias on the reported work. All authors also declare that no animal or human test subjects were involved in research reported here. Thus, no informed consent is necessary. This project was financially supported by the Oklahoma Center for the Advancement of Science and Technology (OCAST Award#PSB10-027), the National Science Foundation Molecular and Cellular Biosciences Award#1024976, The Samuel Roberts Noble Foundation, Oklahoma EPSCoR Research Experience for Undergraduates subaward #EPSCoR-2011-15, the VIB International PhD Fellowship Program (predoctoral fellowship to P.A.), the Research Foundation Flanders (postdoctoral fellowship to J.P.), and the European Union Seventh Framework Program FP7/2007–2013 under grant agreement number 613692–TriForC.



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