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Towards take-all control: a C-21 β oxidase required for acylation of triterpene defence compounds in oat

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Summary

- Oats produce avenacins, antifungal triterpenes that are synthesized in the roots and provide protection against take-all and other soilborne diseases. Avenacins are acylated at the carbon-21 position of the triterpene scaffold, a modification critical for antifungal activity. We have previously characterized several steps in the avenacin pathway, including those required for acylation. However, transfer of the acyl group to the scaffold requires the C-21 β position to be oxidized first, by an as yet uncharacterized enzyme.
- We mined oat transcriptome data to identify candidate cytochrome P450 enzymes that may catalyse C-21 β oxidation. Candidates were screened for activity by transient expression in *Nicotiana benthamiana*.
- We identified a cytochrome P450 enzyme AsCYP72A475 as a triterpene C-21 β hydroxylase, and showed that expression of this enzyme together with early pathway steps yields C-21 β oxidized avenacin intermediates. We further demonstrate that AsCYP72A475 is synonymous with *Sad6*, a previously uncharacterized locus required for avenacin biosynthesis. *sad6* mutants are compromised in avenacin acylation and have enhanced disease susceptibility.
- The discovery of AsCYP72A475 represents an important advance in the understanding of triterpene biosynthesis and paves the way for engineering the avenacin pathway into wheat and other cereals for control of take-all and other diseases.

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Introduction

Plants produce a wealth of diverse natural products. These compounds have important ecological roles, providing protection against pests and diseases. Oats (*Avena* spp.) are unusual amongst the cereals and grasses in that they produce antimicrobial triterpene glycosides (saponins) (Turner, 1960). Saponins are one of the largest families of plant natural products and are produced primarily by eudicots (Hostettmann & Marston, 1995). Previously we have shown that saponins produced in oat roots (avenacins) provide protection against soilborne pathogens such as *Gaeumannomyces graminis* var. *tritici*, causal agent of take-all disease (Osbourn *et al.*, 1994; Papadopolou *et al.*, 1999). Take-all causes major yield losses in wheat, with complete crop failure occurring under severe disease conditions. Disease severity increases with successive wheat cropping, and growth of second and third wheat crops in the same field can become economically unviable (Asher & Shipton, 1981; Hornby & Bateman, 1998). The problem is confounded by the fact that sources of genetic resistance to take-all disease have not yet been identified in hexaploid wheat germplasm (Scott *et al.*, 1989; Freeman & Ward, 2004).

The most effective way to achieve simple, economic and sustainable control of take-all disease would be through genetic resistance. Oats have extreme resistance to *G. graminis* var. *tritici* and are used in rotation for take-all control. The major avenacin, A-1, fluoresces bright blue under ultraviolet illumination as a result of the presence of *N*-methyl anthranilate, which is attached at the carbon-21 position of the triterpene scaffold (Fig. 1a). Previously we exploited this property to screen for mutants of diploid oat (*Avena strigosa*) with reduced root fluorescence, and identified *c.* 100 avenacin-deficient mutants (Papadopolou *et al.*, 1999; Qi *et al.*, 2006). These mutants have enhanced susceptibility to *G. graminis* var. *tritici* and other soilborne fungal pathogens, indicating that avenacins confer disease resistance (Papadopolou *et al.*, 1999; Mugford *et al.*, 2013). We have subsequently characterized six steps in the avenacin pathway and have shown that the corresponding genes form part of a biosynthetic gene cluster (Haralampidis *et al.*, 2001; Qi *et al.*, 2004, 2006; Geisler *et al.*, 2013; Mugford *et al.*, 2013; Owatworakit *et al.*, 2013; T. Louveau & A. Osbourn, unpublished). The avenacin gene cluster represents the only source of genetic resistance to take-all to be characterized from any cereal or grass species. Wheat and other cereals do not make avenacins, nor do they appear to make other triterpene glycosides. Characterization of

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the complete avenacin biosynthetic pathway would open up new opportunities for take-all control in wheat and other cereals using metabolic engineering approaches.

Oats produce four structurally related avenacins, A-1, B-1, A-2 and B-2 (Fig. 1a). The major avenacin found in oat roots is avenacin A-1. The first committed step in avenacin biosynthesis involves the cyclization of the linear isoprenoid precursor 2,3-oxidosqualene to the pentacyclic triterpene scaffold, β -amyrin. Conversion of β -amyrin to avenacins involves a series of oxygenation steps, addition of a trisaccharide chain at the C-3 position, and acylation at the C-21 position, shown for avenacin A-1 in Fig. 1(b). Acylation of avenacin A-1 is carried out by the serine carboxypeptidase-like acyltransferase *AsSCPL1* (*SAD7*), which uses *N*-methylanthranilate glucoside generated via the shikimate pathway as the acyl donor (Mugford *et al.*, 2009). We have previously shown that acylation is critical for antifungal activity (Mugford *et al.*, 2013). The *AsSCPL1/Sad7* acyltransferase gene is adjacent in the avenacin biosynthetic gene cluster to *AsMT1/Sad9* and *AsUGT74H5/Sad10*, genes that together encode the enzymes necessary for the biosynthesis of the *N*-methylanthranilate glucoside acyl donor (Mugford *et al.*, 2013). However, addition of the acyl group will depend on prior functionalization of the C-21 position of the triterpene scaffold by oxygenation. The enzyme that carries out this key step has not yet been characterized.

Here we identify the missing enzyme that carries out this important functionalization step, so paving the way for the union of the isoprenoid pathway-derived triterpene backbone and the shikimate acid pathway-derived acyl group (Fig. 1b). We show that this cytochrome P450 (CYP) (*AsCYP72A475*), catalyses C-21 β hydroxylation of the avenacin triterpenoid backbone. To our knowledge, this is the first such triterpene C-21 oxidase to be described from monocots. We further demonstrate that *AsCYP72A475* is encoded by the *Sad6* locus, which forms part of the avenacin biosynthetic gene cluster, and show that *AsCYP72A475* is critical for avenacin acylation and plant defence.

Materials and Methods

Oat material, metabolic profiling and disease assays

The wild-type diploid oat accession used in this study was *Avena strigosa* accession S75 (Papadopolou *et al.*, 1999). The *A. strigosa* mutant lines used in this study are described in Papadopolou *et al.* (1999), Haralampidis *et al.* (2001), Qi *et al.* (2004) and Mugford *et al.* (2009). The LC-MS profiling methods used for analysis of root extracts of seedlings of wild-type and mutant oat lines are detailed in Supporting Information Methods S1. Take-all disease assays were performed as described previously, using *Gaeumannomyces graminis* var. *tritici* strain T5 (Papadopolou *et al.*, 1999).

Genetic linkage analysis

Avena atlantica accession Cc7277 (Institute of Biology, Environmental and Rural Sciences (IBERS) collection, Aberystwyth

University, UK) was sequenced by Illumina technology to *c.* 38-fold coverage using paired-end and mate-pair libraries. Assembled contigs were then mapped by survey sequencing of recombinant inbred lines from a cross between Cc7277 and *A. strigosa* accession Cc7651 (IBERS) (R. Vickerstaff & T. Langdon, unpublished). Annotations of contigs linked to the previously identified avenacin biosynthetic genes were used to identify candidate CYP genes.

Generation of Gateway entry and expression constructs

Oligonucleotide primers were purchased from Sigma-Aldrich and all PCR steps were performed using iProofTM High-Fidelity DNA Polymerase (Bio-Rad). Candidate CYP genes were identified by mining (using BLAST searches) a root tip transcriptome database for *A. strigosa* accession S75 that we generated previously (NCBI Sequence Read Archive (SRA) accession ERA148431 (Kemen *et al.*, 2014)). *A. strigosa* root tip RNA was extracted from 5-d-old seedlings and cDNA synthesized as described previously (Kemen *et al.*, 2014). PCR amplification of CYP candidate genes from root tip cDNA was carried out using the primers listed in Table S1 and the attB adapter PCR strategy described in the Gateway Technology manual (Invitrogen). The GenBank accession numbers for the full-length coding sequences of the oat candidate CYPs used in this work are CYP71E22 (MH539811), CYP72A475 (MH539812), CYP72A476 (MH539813), CYP88A75 (MH539814), CYP89N1 (MH539815), CYP706C45 (MH539816), and CYP711A54 (MH539817). The soybean *GmCYP72A69* gene (accession NM_001354946) was synthesized commercially (Integrated DNA Technologies, Leuven, Belgium) with flanking Gateway attB sites.

Gateway Entry constructs were generated by cloning the amplified or synthetic gene products into the pDONR207 vector using BP clonase II enzyme mix (Invitrogen) according to the manufacturer's instructions. The integrity of the Entry clones was checked by Sanger sequencing (performed by GATC Biotech, Konstanz, Germany). Genes were cloned from the relevant Entry vectors into pEAQ-HT-DEST1 (Sainsbury *et al.*, 2009) using the LR clonase II enzyme mix (Invitrogen) according to the manufacturer's instructions. The Lomonosoff laboratory (John Innes Centre, Norwich, UK) kindly provided us with an expression construct containing the green fluorescent protein (GFP) coding sequence in pEAQ-HT-DEST1 (Sainsbury *et al.*, 2009). Expression constructs were transformed into chemically competent cells of the *Agrobacterium tumefaciens* strain LBA4404 by flash-freezing in liquid nitrogen.

Transient expression in *N. benthamiana*

Transformed *A. tumefaciens* strains carrying the relevant expression constructs were cultured and infiltrated into *N. benthamiana* using a syringe without a needle, as previously described (Sainsbury *et al.*, 2012; Reed *et al.*, 2017). Extraction of leaves and GC-MS analysis were performed as described previously (Reed *et al.*, 2017). Methods used for LC analysis can be found in Supporting Information Methods S2.

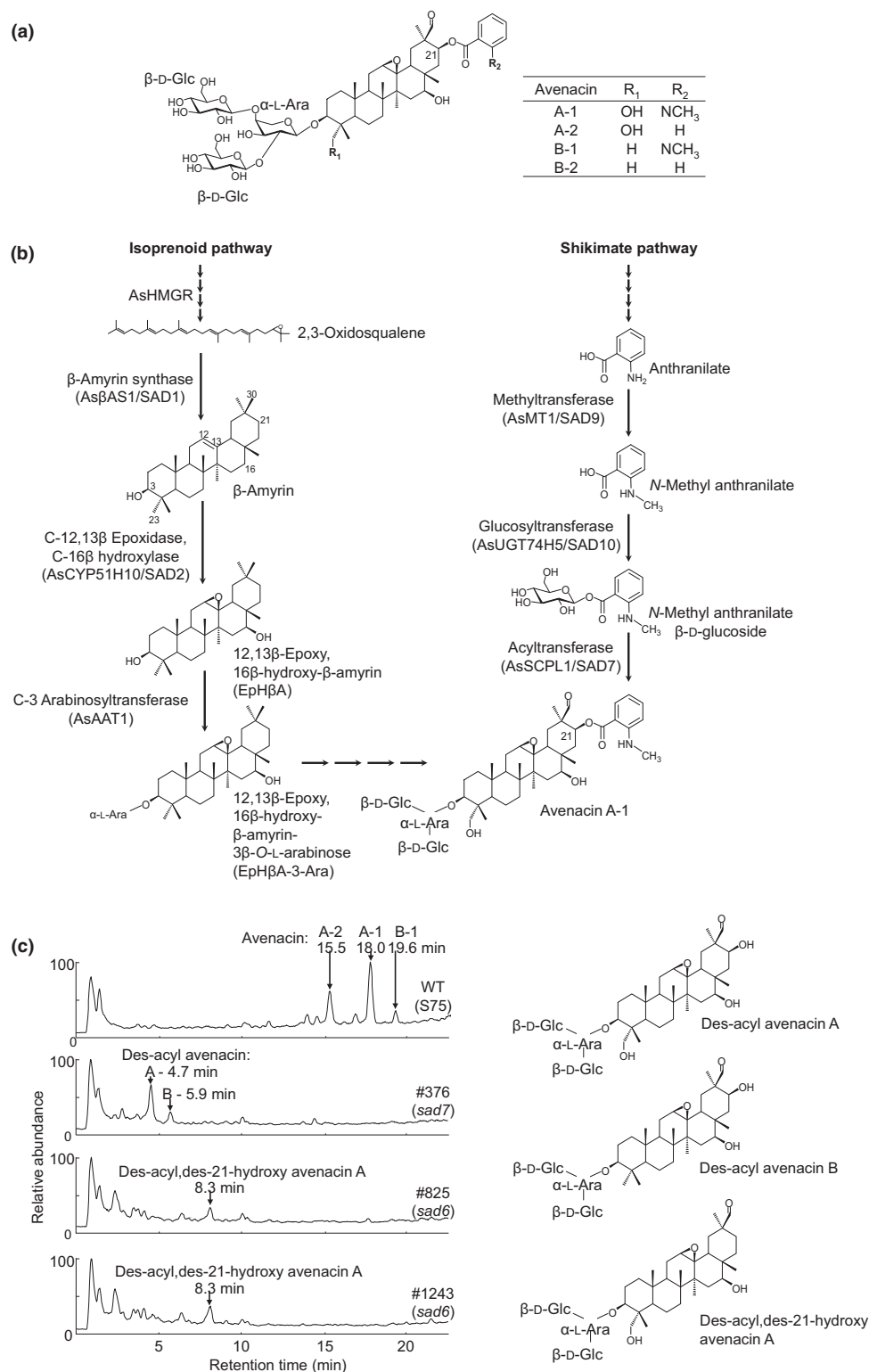


Fig. 1 Identification of C-21 β oxidase-deficient (*sad6*) oat mutants. (a) Structures of the four avenacins. Avenacin A-1 is the major avenacin found in oat roots. Avenacins A and B are differentiated by the presence (A) or absence (B) of a hydroxyl group at C-23 (R₁). Avenacins A-1 and B-1 are acylated with *N*-methyl anthranilate at the C-21 position, and avenacins A-2 and B-2 have benzoate at this position (R₂). AsHMGR, *Avena strigosa* HMG-CoA reductase. (b) Current understanding of the pathway for the biosynthesis of avenacin A-1. The triterpene scaffold originates from the isoprenoid pathway (left), and the *N*-methyl anthranilate acyl group from the shikimate pathway (right). (c) LC-MS analysis of root extracts of seedlings of the wild-type oat accession and *sad6* and *sad7* mutant lines. The major avenacin peaks (A-1, A-2 and B-2, 15.5–19.6 min) observed in the wild-type are absent in *sad6* and *sad7* mutants (the least abundant, B-2, was below the detection limit in these analyses). The *sad7* mutants accumulate nonacylated avenacins (des-acyl-avenacins A and B) as previously reported (Mugford *et al.*, 2009), owing to lack of the acyltransferase, AsSCPL1/SAD7. The *sad6* mutants #825 and #1243 accumulate a form of nonacylated avenacin, putatively designated des-acyl, des-21-hydroxy-avenacin A (Supporting Information Fig. S1).

AsCYP72A475 sequencing and gene expression analysis

Avena strigosa seedlings were grown as previously described (Papadopoulou *et al.*, 1999). Genomic DNA was extracted from *A. strigosa* leaves using a DNeasy Plant Mini Kit (Qiagen). Primers used for sequencing the *AsCYP72A475* gene are listed in Table S2. For gene expression profiling by reverse transcription polymerase chain reaction (RT-PCR) and quantitative polymerase chain (qPCR), shoots, whole roots, root tips (terminal 2–3 mm) and elongation zones (section of roots directly above the root tip, *c.* 4 mm long) of 5-d-old seedlings were harvested and RNA extracted using an RNeasy Plant Mini Kit (Qiagen) with on-column DNase digestion (RG1; Promega). cDNA was synthesized using Superscript III (ThermoFisher, Waltham, MA, USA) with oligo dT primers. One microgram of total RNA from each tissue type was used per cDNA synthesis reaction. Primers for RT-PCR (Table S3) and qPCR analysis (Table S4) were designed using Primer3 (Untergasser *et al.*, 2012). RT-PCR was performed using GoTaq Green polymerase (Promega) with a 55°C annealing temperature and 40 s elongation time over 30 cycles. PCR products were analysed by agarose gel electrophoresis. For qPCR, samples were analysed in triplicate using the DYNAMO Flash SYBR Green kit (ThermoFisher) according to the manufacturer's instructions. qPCR thermal cycling and analysis were performed using a Bio-Rad CFX96 Touch Real-Time PCR Detection system. For all experiments, target amplification was measured using the $\Delta\Delta C_q$ method (Livak & Schmittgen, 2001) relative to the *A. strigosa* *EF1- α* gene (Kemen *et al.*, 2014).

Phylogenetic analysis

CYP72A protein sequences used for phylogenetic analysis were from Prall *et al.* (2016), with additional inclusion of KpCYP72A397 from *Kalopanax septemlobus* (Han *et al.*, 2018), OsCYP72A31 from rice (Saika *et al.*, 2014), and the oat *AsCYP72A475* CYP reported here. Phylogenetic analysis was carried out using MEGA 6 (Tamura *et al.*, 2013). Sequence alignment was performed using MUSCLE (Edgar, 2004) and neighbour joining (Saitou & Nei, 1987) was carried out using the Jones–Taylor–Thornton method (Jones *et al.*, 1992) (with 1000 bootstrap replicates).

Purification and structural elucidation of 12,13-epoxy, 16,21-dihydroxy- β -amyrin-3-O-L-arabinose

Details of the purification process and nuclear magnetic resonance (NMR) experiments are provided in Supporting Information Methods S3.

Results

Identification of C-21 β oxidase-deficient (*sad6*) oat mutants

Previously, we reported the identification of two avenacin-deficient *A. strigosa* mutants #825 and #1243 that were initially

assigned to separate loci (Papadopoulou *et al.*, 1999; Qi *et al.*, 2004) but that we have subsequently shown to be allelic. These mutants represent independent mutant alleles of *Sad6*, a locus that we have defined genetically but that has not yet been cloned and characterized. The *Sad6* locus cosegregates with the other characterized avenacin biosynthetic genes and so is anticipated to be part of the biosynthetic gene cluster (Qi *et al.*, 2004). Metabolite analysis of oat root extracts by LC-MS indicates that the avenacin-deficient *sad6* mutants #825 and #1243 accumulate a new compound with a retention time of 8.3 min (Fig. 1c). *sad7* mutants lack the acyltransferase needed for addition of the acyl group at the C-21 β position (Fig. 1b) and so accumulate the nonacylated avenacins, des-acyl avenacins A and B (retention times 4.7 and 5.9 min, respectively) (Fig. 1c). The mass spectrum of the product observed for the *sad6* mutants was similar to that of des-acyl avenacin B (Fig. S1), suggesting that the acyl group is also absent in mutants #825 and #1243. However, in-chamber fragmentation of the new compound revealed that it contained only three of the four hydroxyl groups found in avenacin A (Fig. S1). The absence of the C-21 β hydroxyl group was considered most likely, given that this would prevent subsequent acylation and account for the observed masses. Hence the product was putatively designated des-acyl, des-21-hydroxy-avenacin A (Fig. S1).

Identification of candidate genes by transcriptome mining

Accumulation of des-acyl, des-21-hydroxy-avenacin A in *sad6* mutants #825 and #1243 suggests that these lines may have undergone mutations in a gene encoding a β -amyrin C-21 β oxidase, most likely a CYP. We therefore mined an oat root tip transcriptome database that we generated previously (SRA accession ERA148431; Kemen *et al.*, 2014) (within which all six previously characterized avenacin genes are represented) for all predicted CYPs. Over 100 CYP sequences were identified. To prioritize these for functional analysis, we exploited the fact that the *Sad6* locus is known to be genetically linked to the characterized avenacin genes (Qi *et al.*, 2004). We used the sequences of the six previously characterized avenacin biosynthesis genes to mine genomic sequence data generated by survey resequencing of a diploid oat mapping population derived from a cross between two avenacin-producing diploids, *A. strigosa* (IBERS Cc7651) and *A. atlantica* (IBERS Cc7277). Within the contigs that we recovered, we identified a total of eight CYPs genes, including the previously characterized *AsCYP51H10* (*Sad2*) gene (Qi *et al.*, 2006; Geisler *et al.*, 2013). The seven new *A. strigosa* CYP sequences were assigned to CYP clans and named by the Cytochrome P450 Nomenclature Committee following established convention (Nelson, 2006) (Fig. 2a).

Expression of the previously characterized avenacin biosynthetic genes is restricted to the root tips, with little/no expression in other tissues (Haralampidis *et al.*, 2001; Qi *et al.*, 2006; Mugford *et al.*, 2009, 2013; Owatworakit *et al.*, 2013; Kemen *et al.*, 2014). We therefore investigated the expression profiles of our candidate CYP genes in different oat tissues by RT-PCR. Two of the previously characterized avenacin biosynthetic genes,

(a)

CYP	Clan	Family (subfamily)	Scaffold no.	Length (bp)
<i>AsCYP51H10/Sad2</i>	CYP51	CYP51(H)	17168	1473
<i>AsCYP72A475</i>	CYP72	CYP72(A)	18622/00941/00942	1608
<i>AsCYP72A476</i>	CYP72	CYP72(A)	9269	1548
<i>AsCYP706C45</i>	CYP71	CYP706(C)	19837	1764
<i>AsCYP71E22</i>	CYP71	CYP71(E)	23466/23468	1518
<i>AsCYP89N1</i>	CYP71	CYP89(N)	18440	1485
<i>AsCYP88A75</i>	CYP85	CYP88(A)	13408/06752	1482
<i>AsCYP711A54</i>	CYP711	CYP711(A)	06108/19967	1605

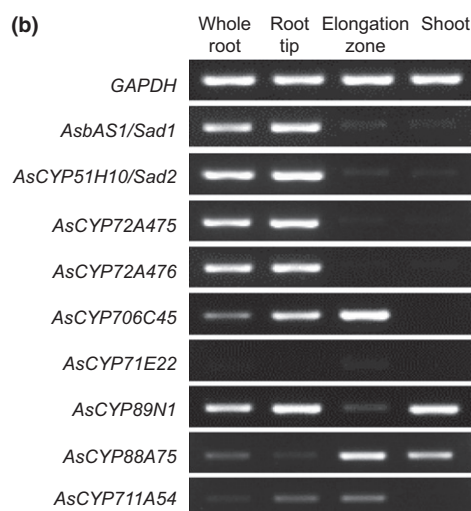


Fig. 2 Candidate CYPs identified by transcriptome mining. (a) Root-expressed candidate CYPs identified in *Avena strigosa*. The corresponding scaffold names from the oat root transcriptome database (Kemen *et al.*, 2014) are shown. (b) RT-PCR analysis of the transcript abundances of candidate CYP genes in different tissues from 3-d-old oat seedlings. Two previously characterized avenacin biosynthetic genes (*AsbAS1/Sad1* and *AsCYP51H10/Sad2*) and the housekeeping gene glyceraldehyde-3-phosphate dehydrogenase gene (*GAPDH*) are included as controls.

AsbAS1/Sad1 and *AsCYP51H10/Sad2*, were included for comparison, along with the housekeeping gene glyceraldehyde 3-phosphate dehydrogenase (*GAPDH*). Two of the seven new CYP genes (*AsCYP72A475* and *AsCYP72A476*) had expression profiles that closely resembled those of the characterized avenacin biosynthetic genes (Fig. 2b), making them the most promising candidates for the C-21 β hydroxylation step in avenacin biosynthesis.

AsCYP72A475 catalyses C-21 β hydroxylation of the β -amyryn scaffold

Previously we showed that we could coexpress the first two steps in the avenacin pathway (β -amyryn synthase (*AsbAS1/SAD1*) and the CYP *AsCYP51H10/SAD2*) in *N. benthamiana* using transient plant expression and obtain the oxygenated avenacin pathway intermediate 12,13 β -epoxy,16 β -hydroxy- β -amyryn (EpH β A) (Geisler *et al.*, 2013). Introduction of a third pathway enzyme (the oat arabinosyltransferase *AsAAT1*) into our coexpression experiments yields the C-3 arabinosylated form of EpH β A (EpH β A-3-Ara) (Fig. 1b) (T. Louveau and A. Osbourn,

unpublished). Enhanced production of these compounds can be achieved using a feedback-insensitive version of the mevalonate pathway enzyme 3-hydroxy,3-methylglutaryl-CoA reductase (*AstHMGR*) (Reed *et al.*, 2017). We therefore took advantage of this quick and efficient transient plant expression system to investigate whether any of the new candidate oat CYP enzymes were able to modify early avenacin pathway intermediates. The predicted full-length coding sequences of *AsCYP72A475*, *AsCYP72A476* and three of the other new CYP genes (*AsCYP706C45*, *AsCYP89N1* and *AsCYP711A54*) were amplified from an oat root tip cDNA library and cloned into the pEAQ-HT-DEST1 expression vector (Sainsbury *et al.*, 2009). We were unable to clone the remaining two coding sequences, *AsCYP71E22* and *AsCYP88A75*. The failure to amplify these genes is in accordance with our RT-PCR profile, which suggests low expression in root tips. Each of the five cloned candidates was evaluated for activity towards β -amyryn (by coexpression with *AstHMGR* and *AsbAS1/Sad1*), and EpH β A (by coexpression with *AstHMGR*, *AsbAS1/Sad1* and *AsCYP51H10/SAD2*). *N. benthamiana* leaves were harvested 5 d after infiltration and leaf extracts were analysed by GC-MS. Of the five candidates tested, none had detectable activity towards β -amyryn (Fig. S2a). However, *AsCYP72A475* was able to modify EpH β A when expressed in combination with *AstHMGR*, *AsbAS1/SAD1* and *AsCYP51H10/SAD2* to give a new peak at 14.9 min (Fig. 3a). No products were observed for the other CYP candidates (Fig. S2b). Analysis of the fragmentation pattern of the new compound was consistent with a putative identity of 12,13 β -epoxy, 16 β ,21 β -dihydroxy- β -amyryn (EpdiH β A) (Figs 3b, S3).

We then attempted to isolate the product generated by *AsCYP72A475* by scaling up our *N. benthamiana* experiments using vacuum infiltration (Reed *et al.*, 2017). Unfortunately, despite repeated attempts, we were unable to isolate this product with a purity suitable for NMR analysis. However, inclusion of the avenacin C-3 arabinosyltransferase (*AsAAT1*) (T. Louveau & A. Osbourn, unpublished) in our coexpression experiments led to accumulation of a new compound with a retention time of 6.2 min (Fig. 3c) that proved to be more amenable to purification. Consequently, we chose this combination for large-scale infiltration. A total of 81.4 mg of this new product was purified. Subsequent NMR analysis confirmed that the structure of this product was consistent with it being 12,13 β -epoxy,16 β ,21 β -dihydroxy- β -amyryn-3 β -O-L-arabinose (EpdiH β A-3-Ara) (Figs 3d, S4). Collectively, our data indicate that *AsCYP72A475* is a C-21 β hydroxylase.

Transient coexpression of the three genes required for avenacin A-1 acylation (*AsSCPL1*, *AsMT1* and *AsUGT74H5*) together with *AstHMGR*, *AsbAS1/SAD1*, *AsCYP51H10/SAD2*, *AsCYP72A475* and *AsUGT99D1* in *N. benthamiana* yielded EpdiH β A-3-Ara and the acyl donor *N*-methyl anthranilate β -D-glucoside, but we were unable to detect any acylated triterpene. We have previously shown that the *AsSCPL1*, *AsMT1* and *AsUGT74H5* proteins are correctly processed and functional when transiently expressed in *N. benthamiana* (Mugford *et al.*, 2009, 2013). Hence this may suggest that additional modifications to the triterpene scaffold are required before acylation can

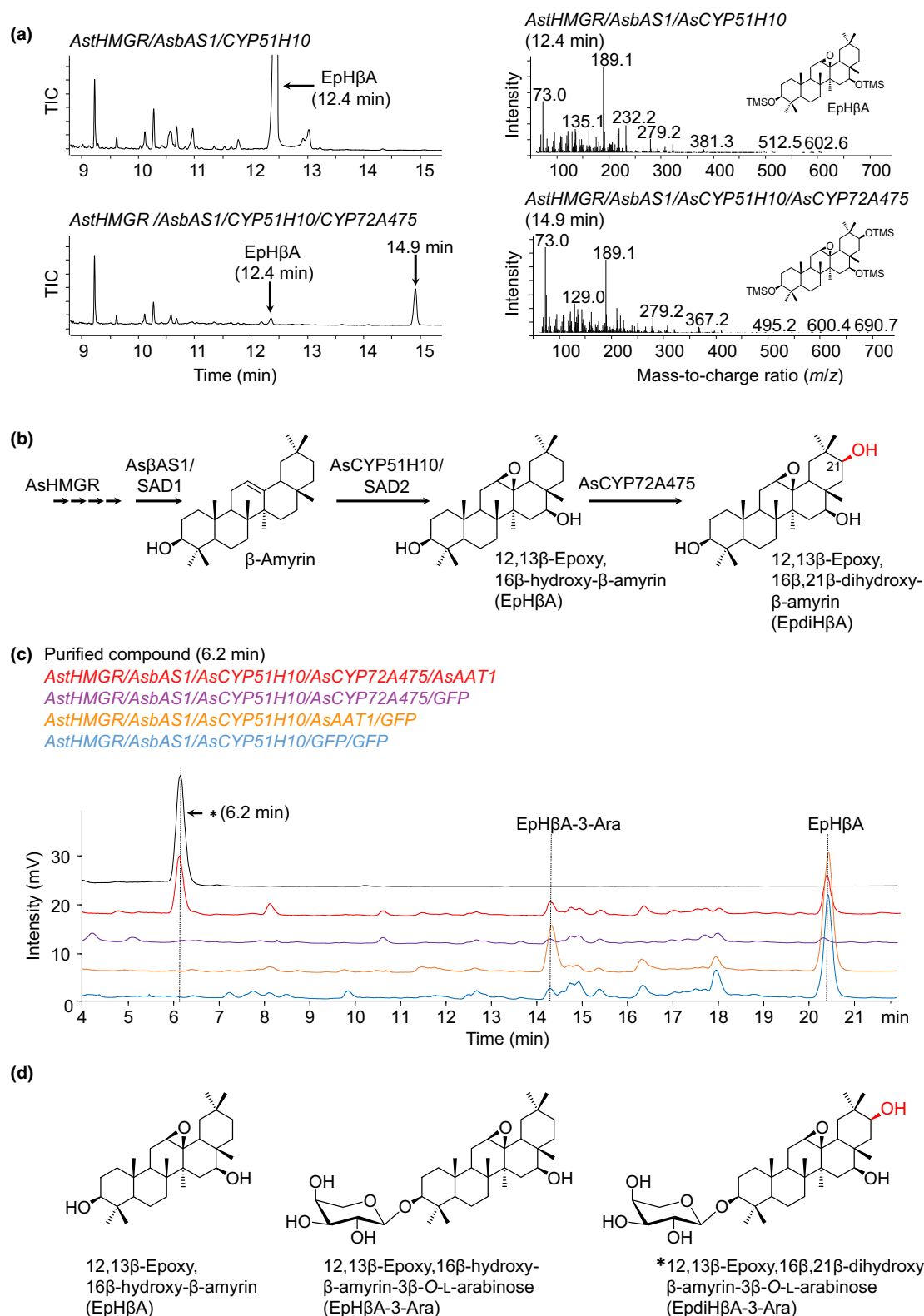


Fig. 3 Production of Epdih β A by transient expression of *AsCYP72A475* in *Nicotiana benthamiana*. (a) GC-MS total ion chromatograms (TIC) of extracts from leaves expressing *AstHMGR/AsbAS1/AsCYP51H10* alone or with *AsCYP72A475*. Mass spectra for 12,13 β -epoxy-16 β -hydroxy- β -amyrin (Eph β A, 12.4 min) and the novel peak putatively identified as 12,13 β -epoxy, 16 β , 21 β -dihydroxy- β -amyrin (Epdih β A, 14.9 min) are shown on the right. Spectra are provided in Supporting Information Fig. S3. (b) Proposed pathway for biosynthesis of Epdih β A from β -amyrin in *N. benthamiana*. (c) LC-charged aerosol detection chromatograms of extracts from leaves transiently expressing *AstHMGR*, *AsbAS1* and *AsCYP51H10* with different combinations of *AsCYP72A475* and the C-3 arabinosyltransferase, *AsAAT1*. Upon coexpression of all enzymes, a novel peak was observed at 6.2 min (marked with an asterisk). (d) Structure of the purified compound with retention time 6.2 min (c) as determined by nuclear magnetic resonance (further details provided in Fig. S4). *AsHMGR* and *AstHMGR*, *Avena strigosa* wild-type and feedback-insensitive HMG-CoA reductase, respectively (Reed *et al.*, 2017).

occur. Alternatively it is possible that any acylated product is also further modified by *N. benthamiana* endogenous enzymes to another product that we could not identify.

Phylogenetic and comparative analysis of CYP72A family members

AsCYP72A475 belongs to the CYP72 clan of the CYP superfamily (Nelson, 2006). Although the CYP72 clan is one of the largest groups of plant CYPs (Nelson & Werck-Reichhart, 2011; Hamburger & Bak, 2013; Prall *et al.*, 2016), relatively little is known about the functions of CYP72 enzymes (Irmeler *et al.*, 2000; Seki *et al.*, 2011; Fukushima *et al.*, 2013; Itkin *et al.*, 2013; Miettinen *et al.*, 2014; Saika *et al.*, 2014; Biazzi *et al.*, 2015; Umemoto *et al.*, 2016; Yano *et al.*, 2017; Han *et al.*, 2018). The only previously characterized CYP72A enzyme from monocots is CYP72A31 from rice, which functions in inactivation of an acetolactate synthase-inhibiting herbicide (Saika *et al.*, 2014). Characterized CYP72A enzymes from eudicots include those required for secaloganin biosynthesis in *Catharanthus roseus* (CYP72A1 and CYP72A224) (Irmeler *et al.*, 2000; Miettinen *et al.*, 2014), steroidal glycoalkaloid biosynthesis in tomato and potato (CYP72A186, CYP72A188 and CYP72A208) (Itkin *et al.*, 2013; Umemoto *et al.*, 2016) and several triterpene oxidases that modify various positions around the triterpene scaffold (Seki *et al.*, 2011; Fukushima *et al.*, 2013; Biazzi *et al.*, 2015; Yano *et al.*, 2017; Han *et al.*, 2018). One of these (GmCYP72A69 from soybean) has recently been reported to oxygenate the triterpene soyasapogenol B at the C-21 β position (Fig. 4a) (Yano *et al.*, 2017). The oat AsCYP72A475 CYP groups with the rice CYP72A31 enzyme in phylogenetic analysis in a subclade that is distinct from the characterized eudicot CYP72As (Fig. 4b). When the soybean enzyme GmCYP72A69 was coexpressed with AsHMGR and AsBAS1/SAD1 in *N. benthamiana*, a product with a retention time of 12.37 min and a mass spectrum consistent with that of 21-hydroxy- β -amyrin was detected (Fig. 4c). Hence GmCYP72A69, unlike AsCYP72A475, is able to oxygenate β -amyrin directly. In contrast, coexpression of GmCYP72A69 with AsHMGR, AsBAS1/SAD1 and AsCYP51H10/SAD2 yielded only low levels of EpdiH β A relative to EpH β A when compared with AsCYP72A475 (Fig. 4d).

AsCYP72A475 is synonymous with Sad6

AsCYP72A475, like the other characterized avenacin biosynthesis genes, is expressed primarily in the root tips of wild-type *A. strigosa* seedlings, with little or no expression in other tissues (Figs 2b, 5a). The transcript abundance of this gene was substantially reduced/undetectable in the roots of seedlings of the two *sad6* mutants, #1243 and #825 (Fig. 5b). DNA sequence analysis revealed a predicted stop codon in AsCYP72A475 at nucleotide 2772 in mutant #1243, suggesting that the reduced transcript abundances in this line may be attributable to nonsense-mediated RNA decay (Chiba & Green, 2009; Christie *et al.*, 2011). Despite numerous attempts, we were unable to amplify the AsCYP72A475 gene from genomic DNA of mutant #825. This

is consistent with our failure to detect any transcript for AsCYP72A475 in this line by qPCR (Fig. 5b), possibly owing to rearrangement or deletion in this region.

Analysis of a further 55 additional uncharacterized avenacin-deficient mutants from our collection by LC-MS coupled with fluorescence detection (LC-MS-Fluo) identified 12 more mutants that, like #1243 and #825, accumulate des-acyl,des-21-hydroxy avenacin A-1 and B-1 (Fig. S6). DNA sequence analysis revealed that these lines also have mutations in the AsCYP72A475 gene (Fig. S7; Supporting Information Methods S4). These lines, like #1243 and #825 (Papadopoulou *et al.*, 1999), also have increased susceptibility to take-all disease (Fig. 5c). Collectively our data show that AsCYP72A475 is synonymous with *Sad6*, and that this CYP is required for C-21 β hydroxylation of the avenacin scaffold. This modification is required for acylation and hence for plant defence.

Discussion

Biosynthesis of the major avenacin, A-1, requires the concerted action of two pathways: the isoprenoid pathway, which supplies the triterpene backbone, and the shikimate pathway, which supplies the acyl moiety (Fig. 1b). Here we report the discovery and characterization of AsCYP72A475, a CYP that oxygenates the C-21 position of the triterpene scaffold, so paving the way for avenacin acylation. The AsCYP72A475 gene is synonymous with *Sad6*, a locus that we previously identified in a forward screen for avenacin-deficient mutants. In earlier work we showed that transient expression of the anthranilate *N*-methyltransferase AsMT1 (encoded by *Sad9*) with the glucosyltransferase AsUGT74H5 (*Sad10*) in *N. benthamiana* leads to accumulation of *N*-methyl anthraniloyl-*O*-Glucose (NMA-glc), the acyl donor needed for addition of the *N*-methyl anthranilate group (Mugford *et al.*, 2013). We also showed that the acyl transferase that mediates this process, the serine carboxypeptidase-like acyltransferase AsSCPL1 (*SAD7*), can be expressed in *N. benthamiana* and that this heterologously expressed enzyme is functional (Mugford *et al.*, 2009). However, when we coexpressed AsHMGR, AsBAS1/SAD1, AsCYP51H10/SAD2, AsAAT1 and AsCYP72A475 together with these three enzymes, we were able to detect EpdiH β A-3-Ara and the acyl donor *N*-methyl anthraniloyl-*O*-Glc, but not acylated EpdiH β -3-Ara. This is not surprising, given that AsSCPL1 is localized in the vacuole (the site of avenacin acylation), while the other characterized pathway enzymes are either known or predicted to be cytosolic (Mugford *et al.*, 2013). Thus, further modification of the avenacin backbone by oxygenation and glycosylation is likely to be needed before transport to the vacuole, and potentially also one or more vacuolar transporter proteins. Future investigations will focus on identification of the missing steps in the avenacin biosynthetic pathway with the aim of reconstituting the entire pathway in *N. benthamiana*, and ultimately in major crops such as wheat.

Oxidation plays a critical role in natural product biosynthesis, furnishing functional groups that allow further elaboration of scaffold molecules through processes such as glycosylation and acylation. CYPs are therefore key drivers of metabolic

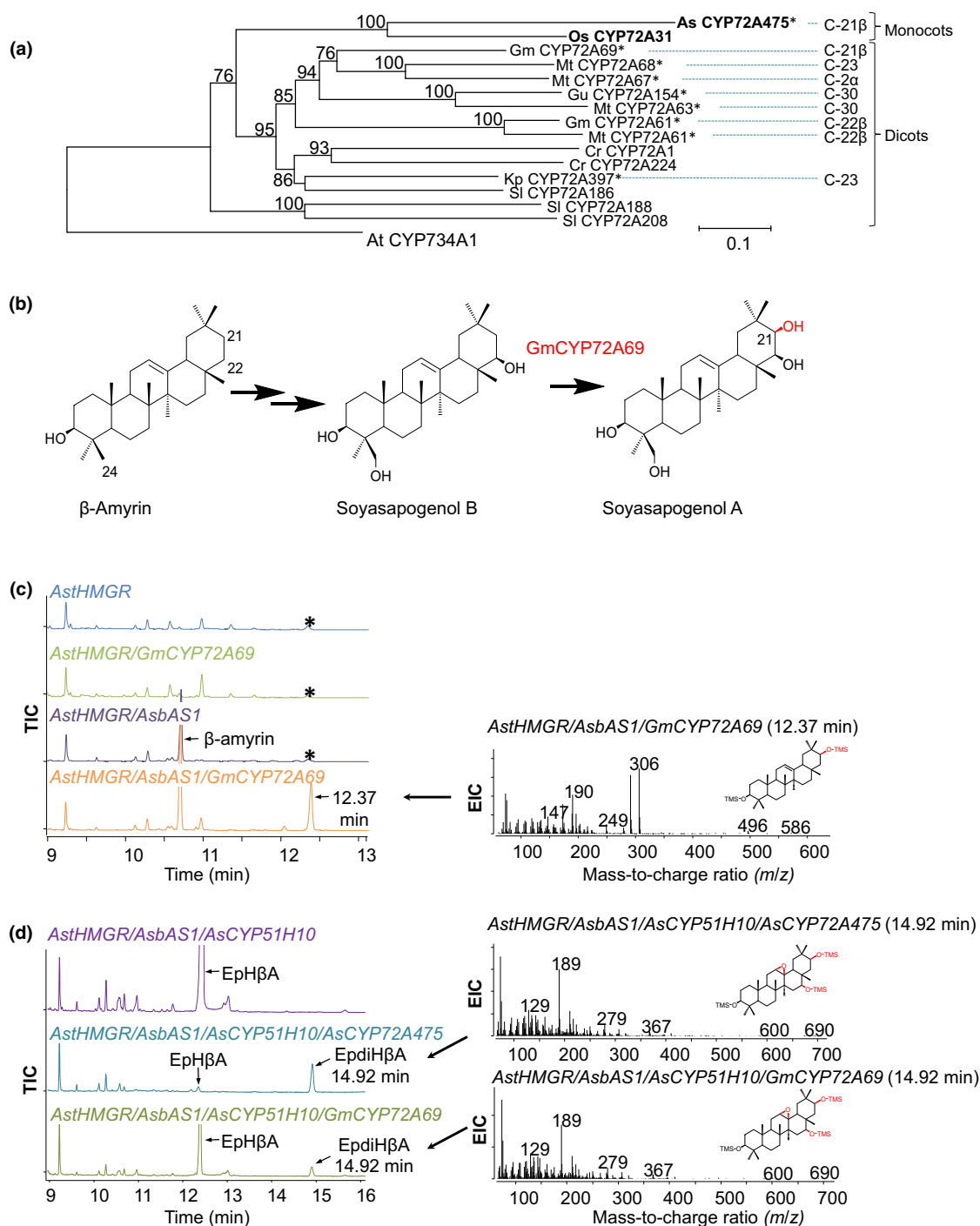


Fig. 4 Phylogenetic and comparative analysis. (a) Neighbour-joining phylogenetic tree of 16 functionally characterized angiosperm CYP72A enzymes (Irmiler *et al.*, 2000; Seki *et al.*, 2011; Fukushima *et al.*, 2013; Itkin *et al.*, 2013; Miettinen *et al.*, 2014; Saika *et al.*, 2014; Biazzi *et al.*, 2015; Umemoto *et al.*, 2016; Han *et al.*, 2018; Yano *et al.*, 2017; this work). Branch lengths represent number of amino acid substitutions per site (scale bar shown at bottom). Bootstrap support is given as percentages (1000 replicates) next to the branches. The tree is rooted using the (nonCYP72) *Arabidopsis thaliana* CYP734A1 enzyme, as previously described (Prall *et al.*, 2016). Monocot sequences are indicated in bold. The eudicot CYP72A family members known to oxidize β -amyrin (or derivatives of β -amyrin) are marked with asterisks and the carbon position that these enzymes modify are shown on the right. As, *Avena strigosa*; At, *A. thaliana*; Cr, *Catharanthus roseus*; Gm, *Glycine max*; Gu, *Glycyrrhiza uralensis*; Kp, *Kalopanax septemlobus*; Mt, *Medicago truncatula*; Os, *Oryza sativa*; Sl, *Solanum lycopersicum*. (b) The soybean *GmCYP72A69* has previously been reported to oxidize the β -amyrin-derived scaffold soyasapogenol B at the C-21 β position to produce soyasapogenol A (Yano *et al.*, 2017). (c) Coexpression of soybean *GmCYP72A69* with *AstHMGR* and *AsbAS1* results in a new peak (12.37 min), which was putatively identified as 21 β -hydroxy- β -amyrin. A representative mass spectrum for the putative 21 β -hydroxy- β -amyrin is given on the right. The mass spectrum for a minor coeluting product (labelled * in the controls) is shown in Supporting Information Fig. S5. (d) Coexpression of *GmCYP72A69* with *AstHMGR*, *AsbAS1* and *AsCYP51H10* results in accumulation of a novel product (14.92 min) with a matching retention time and mass spectrum to *EpdiH β A*, as produced by coexpression of *AstHMGR*, *AsbAS1*, *AsCYP51H10* and *AsCYP72A475*. *AstHMGR*, feedback-insensitive *A. strigosa* HMG-CoA reductase (Reed *et al.*, 2017).

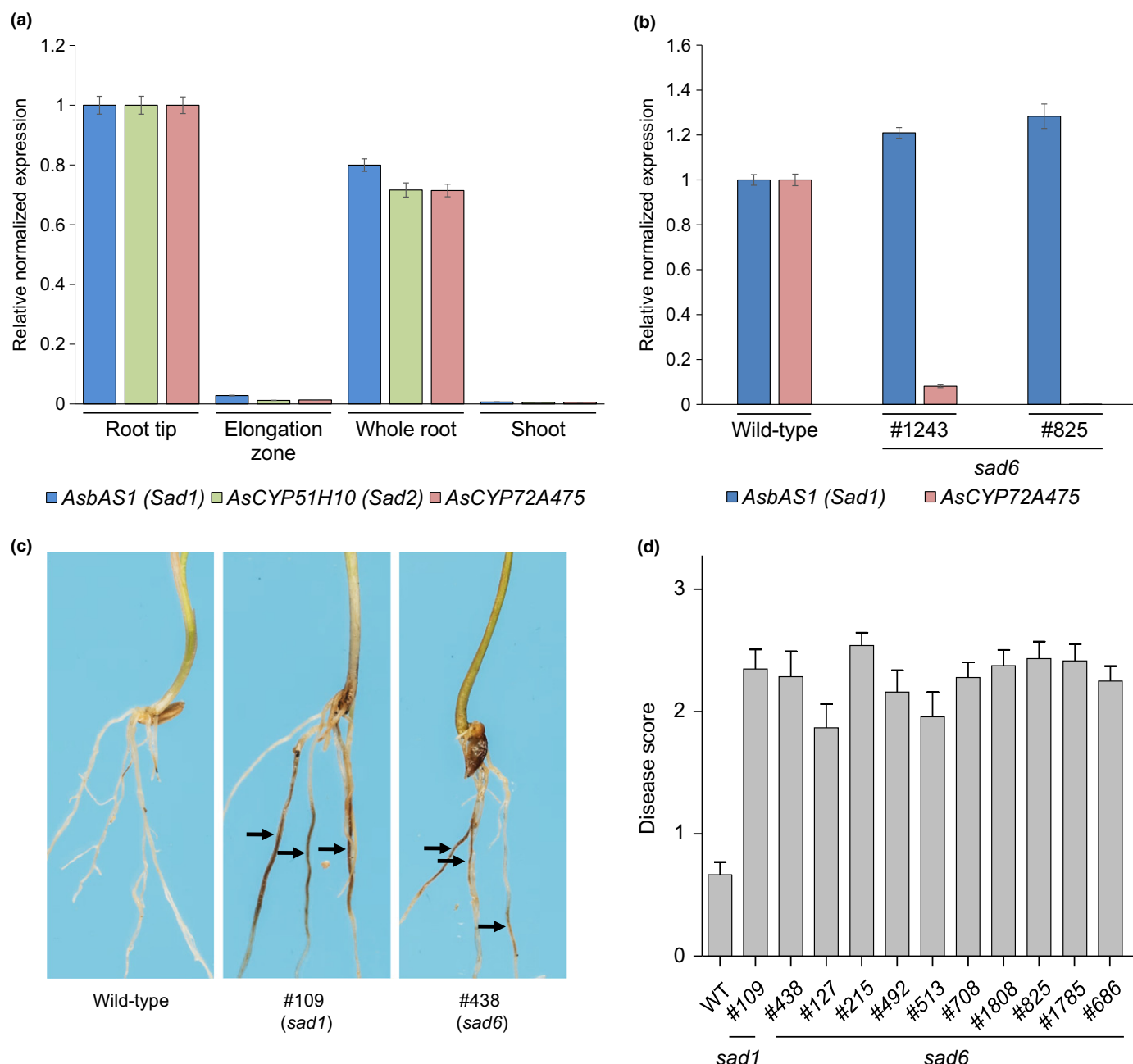


Fig. 5 Analysis of *AsCYP72A475* (*sad6*) mutants. (a) Quantitative reverse transcription polymerase chain reaction (qRT-PCR) analysis of transcript abundances of *AsbAS1/Sad1*, *AsCYP51H10/Sad2* and *AsCYP72A475* in RNA extracted from 3-d-old *Avena strigosa* seedlings. (b) qRT-PCR analysis of *AsCYP72A475* transcript abundances in RNA extracted from roots of wild-type and *sad6* mutant lines. Expression levels are shown relative to the wild-type. All qPCR data transcript abundances were normalized to those for the elongation factor 1 (*EF1- α*) housekeeping gene, using the $\Delta\Delta C_q$ method (Livak & Schmittgen, 2001). Values are means $\pm$ SE (three technical replicates). (c) Representative roots of wild-type, *sad1* and *sad6* *A. strigosa* lines following inoculation with the take-all fungus (*Gaeumannomyces graminis* var. *tritici* strain T5). Roots were scored based on presence and extent of lesions (arrows) as previously described (Papadopoulou *et al.*, 1999). (d) Disease scores for a suite of *sad6* mutant lines (Supporting Information Fig. S7a) compared with those of the wild-type (S75) and a susceptible *sad1* mutant line (#109). The bars represent mean disease scores (21–25 seedlings per line). Error bars represent the SE of the mean.

diversification. C-21-oxidized β -amyrin-derived (oleanane-type) triterpenes are common in the plant kingdom (Vincken *et al.*, 2007), and acylation at this position is prevalent within certain plant families (Lacaille-Dubois *et al.*, 2011). C-21 β acylation is critical for the antifungal activity of avenacins (Mugford *et al.*, 2013). Acylation of other triterpenoids at this position is also

known to determine cytotoxicity towards cancer cells (Chan, 2007). The numbers of characterized triterpene-modifying CYPs is continuing to grow (Thimmappa *et al.*, 2014; Seki *et al.*, 2015; Ghosh, 2017), and as it becomes possible to selectively oxygenate triterpene scaffolds at different carbon positions, this will create opportunities for metabolic engineering of this important family

of natural products for diverse applications. The number of characterized triterpene-modifying acyltransferases (SCPL-like enzymes and BAHD acyltransferases) is currently very limited (Mugford *et al.*, 2009; Shang *et al.*, 2014). However, this is likely to change as the rapidly increasing body of genome sequence data available for diverse plant species continues to expand. The tendency of genes for different triterpene biosynthetic pathways to be clustered in plant genomes, as is the case for the oat avenacin pathway and for pathways for the biosynthesis of triterpenes associated with bitterness in the Cucurbitaceae (Qi *et al.*, 2004; Mugford *et al.*, 2013; Shang *et al.*, 2014; Zhou *et al.*, 2016), is already facilitating the discovery of genes for new candidate triterpene-modifying CYPs, acyltransferases and other tailoring enzymes based on proximity to triterpene synthase genes (Field & Osbourn, 2008; Boutanaev *et al.*, 2015).

To our knowledge, the oat AsCYP72A475 is the first C-21 β triterpene oxidase to be reported from monocots. A second plant β -amyrin C-21 β oxidase, GmCYP72A69, was recently reported from soybean (Yano *et al.*, 2017). Both enzymes belong to the CYP72A subfamily of the CYP72 clan. Several CYP72A subfamily members have recently been characterized from eudicots (from the Fabaceae and Araliaceae families) and shown to be triterpene oxidases (Seki *et al.*, 2011; Fukushima *et al.*, 2013; Biazzi *et al.*, 2015; Yano *et al.*, 2017; Han *et al.*, 2018). A recent large-scale analysis of angiosperm CYP72As showed that the Poales CYP72As form a distinct monophyletic group (Prall *et al.*, 2016). Likewise, our analysis showed that AsCYP72A475 grouped with the rice OsCYP72A31 in a separate clade to the soybean C-21 oxidase GmCYP72A69 (Fig. 4a). Further transient expression experiments with the two C-21 oxidases suggested distinct differences in relative activity towards different substrates, β -amyrin and EpH β A. While the soybean GmCYP72A69 readily oxidized β -amyrin (Fig. 4c), the oat AsCYP72A475 appeared to be inactive towards this substrate (Fig. S2a). Conversely, GmCYP72A69 showed relatively poor activity towards EpH β A, while CYP72A475 resulted in near-total conversion to EpdiH β A (Fig. 4d). The ability of CYP72A enzymes to oxidize triterpenes is reported to have evolved multiple times in a lineage-specific manner in eudicots (Prall *et al.*, 2016). Given that AsCYP72A475 and GmCYP72A69 share only 46% amino acid identity, the ability to oxidize the C-21 β position of triterpene scaffolds may have arisen independently in oat and soybean. However, our phylogenetic analysis cannot rule out the possibility that the oat and soybean CYP72s may have shared a common ancestor with a role in triterpene biosynthesis. This might suggest that the shared C-21 oxidase activity of AsCYP72A475 and GmCYP72A69 represents a case of parallel rather than convergent evolution. Nevertheless, it is interesting to note that the enzymes catalysing the early steps of triterpene biosynthesis in oat (SAD1/AsbAS and SAD2/CYP51H10) appear to have evolved independently of those in eudicots (Haralampidis *et al.*, 2001; Qi *et al.*, 2006). Knowledge of the differences and parallels in triterpene biosynthesis between monocots and eudicots will facilitate our ability to identify additional triterpene biosynthetic enzymes. Understanding the differences in activity displayed by such enzymes will better inform strategies for optimizing

triterpene biosynthesis in heterologous hosts. Collectively our findings represent an important advance in understanding triterpene biosynthesis and will underpin strategies for metabolic engineering for crop protection, drug development, and other industrial applications.

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Author contributions

AL, JR, XQ, MJS, STM, REM, JCR, RV, TL and AO designed the research; AL, JR, XQ, MJS, STM, REM, JCR and RV performed research; RV and TL contributed new analytic tools; AL, JR, XQ, MJS, STM, REM, JCR, RV, TL and AO analysed data; AL, JR, and AO wrote the paper.

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Supporting Information

Additional Supporting Information may be found online in the Supporting Information section at the end of the article.

Fig. S1 Electrospray ionization (ESI) mass spectra for unacylated avenacins detected in roots of *Avena strigosa* avenacin-deficient mutants shown in Fig. 1(c).

Fig. S2 GC-MS total ion chromatograms (TIC) from analysis of extracts of *Nicotiana benthamiana* leaves expressing *AstHMGR* and *AsbAS1*, or *AstHMGR*, *AsbAS1* and *AsCYP51H10* with candidate CYPs.

Fig. S3 Electron-impact mass spectra for 12,13 β -epoxy-16 β -hydroxy- β -amyirin (EpH β A) and putative 12,13 β -epoxy-16,21 β -hydroxy- β -amyirin (EpdiH β A).

Fig. S4 NMR assignment for 12,13 β -epoxy,16 β ,21 β -dihydroxy- β -amyirin-3 β -O-L-arabinose.

Fig. S5 Electron-impact mass spectra for the product coeluting with 21-hydroxy- β -amyirin.

Fig. S6 Identification of additional *AsCYP72A475* (*sad6*) mutants by LC-MS metabolic profiling.

Fig. S7 Schematic of *AsCYP72A475* and mutation sites in *sad6* mutants.

Methods S1 Metabolite extraction, LC-MS-MS and LC-MS-Fluorescence analysis of oat roots.

Methods S2 Metabolite extraction and LC-MS-CAD analysis of extracts of *Nicotiana benthamiana* leaves.

Methods S3 Purification and structural elucidation of 12,13-epoxy,16,21-dihydroxy- β -amyirin-3-O-L-arabinose.

Methods S4 Structural analysis of *AsCYP72A475*.

Table S1 Oligonucleotide primer sequences used for Gateway cloning of coding sequences of the candidate CYP genes.

Table S2 Oligonucleotide primer sequences used for sequencing of the genomic DNA region encoding for the *Sad6* gene.

Table S3 Oligonucleotide primer sequences used for RT-PCR expression profiling of the candidate CYPs from *Avena strigosa*.

Table S4 Oligonucleotide primer sequences Primers used for quantitative PCR expression profiling of *Avena strigosa* CYPs.

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