UGT86C11 is a novel plant UDP-glycosyltransferase involved in labdane diterpene biosynthesis

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Glycosyltransferases constitute a large family of enzymes across all domains of life, but knowledge of their biochemical function remains largely incomplete, particularly in the context of plant specialized metabolism. The labdane diterpenes represent a large class of phytochemicals with many pharmacological benefits, such as anti-inflammatory, hepatoprotective, and anticarcinogenic. The medicinal plant kalmegh (Andrographis paniculata) produces bioactive labdane diterpenes; notably, the C19-hydroxyl diterpene (andrograpanin) is predominantly found as C19-O-glucoside (neoandrographolide), whereas diterpenes having additional hydroxylation(s) at C3 (14-deoxy-11,12-didehydroandrographolide) or C3 and C14 (andrographolide) are primarily detected as aglycones, signifying scaffold-selective C19-O-glycosylation of diterpenes in planta. Here, we analyzed UDP-glycosyltransferase (UGT) activity and diterpene levels across various developmental stages and tissues and found an apparent correlation of UGT activity with the spatiotemporal accumulation of neoandrographolide, the major diterpene C19-O-glucoside. The biochemical analysis of recombinant UGTs preferentially expressed in neoandrographolide-accumulating tissues identified a previously uncharacterized UGT86 member (ApUGT12/UGT86C11) that catalyzes C19-O-glucosylation of diterpenes with strict scaffold selectivity. ApUGT12 localized to the cytoplasm and catalyzed diterpene C19-O-glycosylation in planta. The substrate selectivity demonstrated by the recombinant ApUGT12 expressed in plant and bacterium hosts was comparable to native UGT activity. Recombinant ApUGT12 showed significantly higher catalytic efficiency using andrograpanin compared with 14-deoxy-11,12-didehydroandrographolide and trivial activity using andrographolide. Moreover, ApUGT12 silencing in plants led to a drastic reduction in neoandrographolide content and increased levels of andrograpanin. These data suggest the involvement of ApUGT12 in scaffold-selective C19-O-glycosylation of labdane diterpenes in plants. This knowledge of UGT86 function might help in developing plant chemotypes and synthesis of pharmacologically relevant diterpenes.

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single plant species and the diversity of specialized metabolites produced in plants, it is quite impossible to assign biochemical function of UGTs in plant specialized metabolism, solely based on sequence similarity. For example, Arabidopsis and rice encode about 120 and 215 UGTs, respectively, but only about 25% UGTs are biochemically characterized so far, and the native substrates of many of the UGTs remain to be known (7, 25, 26).

The labdane diterpenes form a large family of phytochemicals with a broad range of bioactivities (27). These diterpenes represent a major group of bioactive chemicals in the medicinal plant kalmegh (Andrographis paniculata) (28–32). Neoandrographolide, a major bioactive diterpene C19-O-glucoside in kalmegh, possesses anti-inflammatory, hepatoprotective, anticarcinogenic, hypolipidemic, and viricidal activities (33–37). In some studies, neoandrographolide showed superior bioactivity than the diterpene aglycones andrographolide and 14-deoxy-11,12-didehydroandrographolide, suggesting a possible role of glucosylation toward enhanced bioactivity and/or bioavailability of neoandrographolide (33, 37–39). Neoandrographolide biosynthesis in kalmegh starts with a diterpene cyclization reaction catalyzed by the diterpene synthase, converting a general diterpene precursor geranylgeranyl pyrophosphate to ent-copalyl pyrophosphate (Fig. 1). Previously, we have identified a diterpene synthase (ApCPS2) catalyzing the first committed reaction in the neoandrographolide biosynthetic pathway (40, 41). The final step in neoandrographolide biosynthesis involves C19-O-glucosylation of andrograpanin (Fig. 1). However, UGT involved in developmental and tissue-specific biosynthesis of neoandrographolide was not identified. Previously, two methyl jasmonate (MeJA)–inducible UGTs (UGT73AU1 and UGT5), which catalyzed in vitro C19-O-glucosylation of andrograpanin, were identified, but their involvement in planta biosynthesis of neoandrographolide was not understood (42, 43). To investigate the UGT involved in spatiotemporal biosynthesis of neoandrographolide, we have analyzed a large-scale transcriptome data of kalmegh and identified UGTs that preferentially expressed in neoandrographolide-accumulating tissues. Furthermore, 23 recombinant UGTs were screened in UGT assay, leading to the identification of ApUGT12 (UGT86C11) catalyzing C19-O-glucosylation of diterpenes in a scaffold-selective manner (Fig. 1). The steady-state kinetic of ApUGT12, an altered diterpene profiles in ApUGT12-silenced plants, and a strong correlation of ApUGT12 transcript expression with UGT activity and neoandrographolide accumulation patterns across various developmental stages and tissues suggested a pivotal role of ApUGT12 in the biosynthesis of diterpene C19-O-glucoside.

Results

Developmental and tissue-specific patterns of diterpene C19-O-glucosylation activity

To understand spatiotemporal biosynthesis of diterpenes, we conducted comprehensive profiling of diterpene aglycones (andrographolide, andrograpanin, and 14-deoxy-11,12-didehydroandrographolide) and diterpene C19-O-glucosides (neoandrographolide, andrographiside, and 14-deoxy-11,12-didehydroandrographiside) across five developmental stages (germinating seeds, cotyledonary leaf stage, 15-day-old plants, 30-day-old plants, and 60-day-old plants) using six tissues (root, leaf,
stem, sepal, petal, and seedpod) (Fig. S1). HPLC analysis of methanolic extracts revealed higher content of neoandrographolide, andrographolide, and 14-deoxy-11,12-didehydroandrographolide in leaves (Figs. 2A, S2, and S3, A and B). However, these diterpens were not detected in roots and germinating seeds. In addition, neoandrographolide, andrographolide, and 14-deoxy-11,12-didehydroandrographolide were also detected in considerable amounts in seedpod, sepal, and cotyledonary leaf stage seedlings, respectively. The increased amount of neoandrographolide than andrograpanin in kalmegh tissues suggested ready conversion of andrograpanin to neoandrographolide following C19-O-glucosylation, thus limiting in planta accumulation of andrograpanin (Figs. 1 and 2A). On the other hand, higher content of andrographolide and 14-deoxy-11,12-didehydroandrographolide than the corresponding C19-O-glucosides (andrographiside and 14-deoxy-11,12-didehydroandrographiside) indicated inefficient C19-O-glucosylation of andrographolide and 14-deoxy-11,12-didehydroandrographolide (Figs. 1 and S3, A and B). These results suggest that scaffold-selective C19-O-glucosylation potentially contributes to distinct patterns of diterpene aglycones and glucosides in planta. To investigate the involvement of UGT in diterpene C19-O-glucosylation, UGT assays were carried out using total protein extract of various tissues. Andrograpanin, andrographolide, and 14-deoxy-11,12-didehydroandrographolide were the sugar acceptors, whereas UDP-glucose served as sugar donor in assays. Diterpene C19-O-glucoside produced in assay was monitored by HPLC. UGT assays using andrograpanin as sugar acceptor revealed considerably higher enzyme activity in leaves of 60-day-old and 30-day-old plants followed by in leaves of 15-day-old plants and seedpod (Fig. 2B). These tissues also contained higher amount of neoandrographolide (Fig. 2A). UGT assay using total protein extract of roots and germinating seeds could not form neoandrographolide. Likewise, neoandrographolide was not detected in roots and germinating seeds. In contrast, UGT assay using 14-deoxy-11,12-didehydroandrographolide revealed considerably lower activity than using andrograpanin (Fig. S4). However, C19-O-glucosylation of andrographolide could not be achieved at a detectable level using total protein extract of various tissues. Thus, higher UGT activity using andrograpanin than andrographolide and 14-deoxy-11,12-didehydroandrographolide might have contributed to neoandrographolide biosynthesis at a higher rate than andrographiside and 14-deoxy-11,12-didehydroandrographiside biosynthesis, leading to distinct profiles of diterpene aglycones and C19-O-glucosides in planta (Figs. 2, A and B, S3, A and B, and S4). Unlike andrograpanin, the other two diterpene aglycones (andrographolide and 14-deoxy-11,12-didehydroandrographolide) bear additional hydroxyl group(s) at the C3 and/or C14 positions. Therefore, these results strongly suggest that scaffold-selective C19-O-glucosylation of diterpene aglycones by UGT potentially contributes to selective accumulation of diterpene aglycones and glucosides.

Identification of candidate UGT of the diterpene pathway

Andrograpanin C19-O-glucosylation activity and neoandrographolide level in various tissues showed a clear correlation (Fig. 2, A and B). The maximum UGT activity toward andrograpanin C19-O-glucosylation was noticed in leaves of 60-day-old plants; however, UGT activity was not detected in roots. Similarly, neoandrographolide was detected at a higher level in leaves of 60-day-old plants, whereas neoandrographolide could not be detected in roots. To investigate

Figure 2. Spatiotemporal patterns of UGT activity and diterpene C19-O-glucoside accumulation. A, the content of neoandrographolide (NAD), the major diterpene C19-O-glucoside, and its precursor andrograpanin (AGP) in various tissues as determined by HPLC analysis of methanolic extracts. B, the pattern of UGT activity in various tissues. In vitro UGT assays were done in 10 mM Tris–Cl, pH 7.5, at 30 °C using andrograpanin (100 μM) and UDP-glucose (800 μM). The UGT activity is presented as the rate of neoandrographolide formation in assays using total protein extracts of various tissues. A and B, the data are means ± SD of three biological replicates. FW, fresh weight; ND, not detected; TL, trace level; UGT, UDP-glycosyltransferase.
the enzyme involved in C19-O-glucosylation of diterpenes, a large-scale RNA-Seq data representing more than 170 million sequencing reads of leaves and roots were screened and UGTs that preferentially expressed in leaves were identified (41). The transcripts potentially encoding UGTs were retrieved based on annotation to the Carbohydrate-Active enZymes database following BlastX analysis. Among a total of 615 transcripts annotated to various GT families, 161 transcripts were categorized under the GT1/UGT family. Notably, UGT73AU1 (contig ApU2595) and UGT5 (contig ApU62177) transcript expression in leaves and roots was quite comparable (Fig. S5A). The previous study also found a similar transcript expression of UGT73AU1 in leaves and roots; however, UGT5 transcript expression in different tissues was not examined before (42, 43). To know whether UGT73AU1 and UGT5 contribute to developmental and tissue-specific C19-O-glucosylation of diterpenes, UGT73AU1 and UGT5 transcript expression was determined by quantitative RT-PCR (qRT-PCR) analysis and correlated with diterpene accumulation patterns and UGT activity (Fig. S6A). UGT73AU1 and UGT5 transcripts expressed at a higher level in roots of 60-day-old plants and sepal. However, andrograpanin C19-O-glucosylation activity and neoandrographolide content in these tissues were substantially lower or not detected (Fig. 2, A and B). Therefore, UGT73AU1 and UGT5 transcript expression patterns strongly indicated that they might not be playing a major role in developmental and tissue-specific biosynthesis of diterpene glucosides. Moreover, a similar catalytic efficiency of recombinant UGT5 using andrograpanin, andrographolide, and 14-deoxy-11,12-didehydroandrographolide could not corroborate differential activity of native UGT using these diterpene aglycones, if UGT5 is considered to be playing a major role in planta diterpene C19-O-glucosylation (Fig. S4) (42). The transcript expression of ApCPS2, which catalyzed the initial diterpene cyclization reaction in the neoandrographolide biosynthetic pathway, showed a strong correlation with developmental and tissue-specific biosynthesis of neoandrographolide (40, 41). Therefore, it could be hypothesized that UGT transcript expression might also coincide with in planta biosynthesis of neoandrographolide.

The analysis of RNA-Seq data identified 38 nonredundant UGT transcripts that expressed at a higher level in neoandrographolide-accumulating leaves than in roots (Fig. S5A and Table S1). To extract full-length coding sequences of the transcripts, transcriptome assemblies generated in other studies were also consulted (https://medplantnaseq.org/) (44). Five transcripts that either represented incomplete ORFs or encoded truncated proteins were not considered thereafter (Table S1). Furthermore, the analysis of structural motifs in protein sequences identified 32 UGTs that contained the conserved 44 amino acids long plant secondary product glycosyltransferase motif, which participates in binding of UDP-sugar to the active site of bona fide UGTs (Figs. S5B and S7) (45). Although UGTs of the diterpene pathway were not well studied, some UGTs that participate in phenylpropanoid, sterol, and phytohormone pathways were previously characterized (10, 18, 20, 21, 25, 26, 46–48). To shortlist candidate UGTs of kalmegh diterpene pathway, a phylogenetic analysis was carried out using biochemically characterized UGTs. The analysis revealed 24 UGTs, which are not closely related with well-known UGTs of phenylpropanoid, sterol, and phytohormone pathways (Fig. S8). These UGTs were further considered for biochemical characterization to know their involvement in diterpene glucosylation.

**ApUGT12 catalyzes diterpene C19-O-glucosylation**

Among the selected UGTs, the complete ORF of 23 UGTs could be amplified using leaf complementary DNA (cDNA) as template and cloned in pET28a(+) vector for the expression of N-terminally 6xHis-tagged recombinant proteins in *Escherichia coli*. To examine these UGTs for diterpene C19-O-glucosylation activity, *in vitro* assays were carried out using UDP-glucose as sugar donor whereas andrograpanin, andrographolide, and 14-deoxy-11,12-didehydroandrographolide as the sugar acceptors. To begin with, total protein extract of *E. coli* expressing recombinant UGTs and enriched fractions of recombinant UGTs (prepared following nickel–nitrilotriacetic acid [Ni–NTA] affinity chromatography) were used in UGT assays with andrograpanin as sugar acceptor. TLC and HPLC analysis of assay products revealed that only ApU56292 (thereafter described as ApUGT12) catalyzed C19-O-glucosylation of andrographolide to form neoandrographolide (Fig. S9, A and B). To confirm this result, ApUGT12 was purified to electrophoretic homogeneity following Ni–NTA affinity chromatography, representing a ~55 kDa protein band in 10% SDS-PAGE (Fig. 3A). The observed molecular mass of ApUGT12 was similar to the calculated molecular mass (57.99 kDa) of the recombinant ApUGT12. Furthermore, UGT assay was repeated using purified ApUGT12, andrograpanin, and UDP-glucose. The assay product analyzed by LC quadrupole TOF MS (LC–QTOF–MS) confirmed C19-O-glucosylation of andrographolide in neoandrographolide (Fig. 4, A and B). The QTOF mass spectrum of neoandrographolide formed in ApUGT12 assay was in accordance to the observed mass spectrum of neoandrographolide standard (Figs. 4B and S10). Besides, the recombinant ApUGT12 also catalyzed C19-O-glucosylation of 14-deoxy-11,12-didehydroandrographolide to produce 14-deoxy-11,12-didehydroandrographolide, which was identified based on mass spectrum data in LC–QTOF–MS analysis (Fig. S11, A and B). Notably, ApUGT12 showed considerably higher activity using andrograpanin than 14-deoxy-11,12-didehydroandrographolide (Fig. 3D). On the other hand, ApUGT12 exhibited trace activity using andrographolide as sugar acceptor (Figs. 3D and S12, A and B). The optimum pH and temperature for recombinant ApUGT12 were determined *in vitro* assays in a range of pH (pH 4.5–10) and temperatures (20–50 °C) using andrograpanin substrate (Fig. 3, B and C). The analysis revealed that pH 7.5 and 35 °C are the optimum pH and temperature for ApUGT12 activity. Taken together, these results confirmed that ApUGT12 catalyzes C19-O-glucosylation of andrograpanin, 14-deoxy-11,12-didehydroandrographolide, and andrographolide, although at a variable efficiency.
To understand the catalytic property of ApUGT12, steady-state kinetic parameters of recombinant ApUGT12 were determined using andrographanin, 14-deoxy-11,12-didehydroandrographolide, and UDP-glucose. As ApUGT12 showed only trivial activity using andrographanin, the kinetic parameters could not be determined accurately using andrographanin. ApUGT12 displayed quite a dissimilar affinity for andrographanin ($K_m = 137.3$ $\mu$M), 14-deoxy-11,12-didehydroandrographolide ($K_m = 506.7$ $\mu$M), and UDP-glucose ($K_m = 271.16$ $\mu$M). Moreover, $k_{cat}$ value for andrographanin ($0.231$ S$^{-1}$) was significantly higher than 14-deoxy-11,12-didehydroandrographolide ($0.016$ S$^{-1}$). Consequently, ApUGT12 exhibited about 48-fold higher catalytic efficiency ($k_{cat}/K_m$) using andrographanin ($1698.61$ M$^{-1}$ S$^{-1}$) than using 14-deoxy-11,12-didehydroandrographolide ($34.70$ M$^{-1}$ S$^{-1}$). The estimated $k_{cat}$ and $k_{cat}/K_m$ values for 14-deoxy-11,12-didehydroandrographolide were also considerably lower than $k_{cat}$ ($0.167$ S$^{-1}$) and $k_{cat}/K_m$ ($622.80$ M$^{-1}$ S$^{-1}$) values for UDP-glucose. To gain more insights into substrate specificity of ApUGT12, in vitro assays were carried out using UDP-glucose and a range of phytochemicals belonging to terpene, phenylpropanoid, and phenolic classes, such as steviol, kaempferol, gallic acid, 11-keto-β-boswellic acid, arjunic acid, oleanolic acid, maslinic acid, and corosolic acid (Fig. S13). However, ApUGT12 could not glucosylate any of these tested compounds. Overall, these results indicated strict scaffold selectivity of ApUGT12.

**ApUGT12 subcellular localization and diterpene C19-O-glucosylation in planta**

To investigate subcellular localization, ApUGT12 was expressed as enhanced yellow fluorescent protein (EYFP)–tagged...
proteins in *Nicotiana benthamiana* leaf epidermal cells, and EYFP fluorescence in cells was examined with the help of a confocal fluorescence microscope (Fig. 5A). To rule out any possibility of potential subcellular localization signal at the N- or C-terminal region of ApUGT12 that might get masked because of EYFP tagging, both N- and C-terminal EYFP tagging of ApUGT12 (designated as EYFP-ApUGT12 and ApUGT12-EYFP) were carried out. P19, an RNA silencing suppressor, was also coexpressed to attain increased expression of EYFP-ApUGT12 or ApUGT12-EYFP in *N. benthamiana* leaf (49). EYFP fluorescence was detected in the cytoplasm and nucleus of cells expressing free EYFP. However, EYFP fluorescence mostly confined in the cytoplasm of cells expressing EYFP-ApUGT12 or ApUGT12-EYFP (Fig. 5A). To substantiate these results, EYFP-ApUGT12 or ApUGT12-EYFP was coexpressed with a cytoplasmic marker monomeric red fluorescent protein (mRFP) (50). mRFP and EYFP fluorescence were clearly merged in cells coexpressing mRFP and EYFP-ApUGT12 or ApUGT12-EYFP, confirming cytoplasmic colocalization of mRFP, EYFP-ApUGT12, and ApUGT12-EYFP proteins (Fig. 5, B and C). The cytoplasmic localization of ApUGT12 was quite reasonable because plants mostly accumulate glycosylated metabolites in vacuoles, and the transport of such metabolites is potentially driven by their glycosylation in cytoplasm (6, 11).

To know whether ApUGT12 could catalyze in planta diterpene C19-O-glycosylation, *N. benthamiana* leaves expressing ApUGT12-EYFP were infiltrated with andrograpanin. One-day after infiltration of andrograpanin, methanolic extracts of leaves were prepared and analyzed by HPLC. The leaves expressing ApUGT12-EYFP produced neoandrographolide when infiltrated with andrograpanin, implying that ApUGT12 catalyzed in planta C19-O-glycosylation of andrograpanin to produce neoandrographolide (Fig. 5D). However, andrograpanin infiltration in leaves expressing free EYFP (vector control) did not form neoandrographolide, suggesting that the endogenous UGT of *N. benthamiana* could not catalyze C19-O-glycosylation of andrograpanin. To substantiate these results, in vitro C19-O-glycosylation of andrographanin, andrographolide, and 14-deoxy-11,12-didehydroandrographolide was tested using total protein extract of *N. benthamiana* leaves expressing ApUGT12. In vitro assays using protein extract of ApUGT12-expressing leaves clearly showed C19-O-glycosylation of andrographanin to form neoandrographolide (Figs. 5E and S14A). C19-O-glycosylation of 14-deoxy-11,12-didehydroandrographolide was also achieved in assays; however, at a lower rate than andrograpanin C19-O-glycosylation (Figs. 5E and S14B). In contrast, andrographolide could not be glucosylated at a detectable level in assays using total protein extract of ApUGT12-expressing *N. benthamiana* leaves. Overall, these results suggest that ApUGT12 localized in the cytoplasm and catalyzed in planta diterpene C19-O-glycosylation.

**ApUGT12 transcript expression coincides with spatiotemporal biosynthesis of diterpene C19-O-glucoside**

To understand the involvement of ApUGT12 in spatiotemporal biosynthesis of neoandrographolide, *ApUGT12* transcript
expression in kalmegh tissues was determined by qRT-PCR analysis. In accordance with increased UGT activity and neoandrographolide content, ApUGT12 transcript expressed at an increased level in leaves of 60-day-old and 30-day-old plants (Figs. 2, A and B and 6A). Besides, ApUGT12 transcript expressed at a decreased level in tissues showing reduced UGT activity and neoandrographolide content, such as in germinating seeds, roots, and petals. Moreover, MeJA-inducible expression of ApUGT12 transcript and increased UGT activity and neoandrographolide content in leaves of MeJA-treated plants signify a potential role of ApUGT12 in MeJA-induced biosynthesis of neoandrographolide (Figs. 6B and S15, A and B). Taken together, the patterns of
ApUGT12 transcript expression, UGT activity, and neoandrographolide accumulation in various tissues suggest the involvement of ApUGT12 in spatiotemporal biosynthesis of neoandrographolide.

Virus-induced gene silencing of ApUGT12

To clarify the role of ApUGT12 in planta biosynthesis of neoandrographolide, virus-induced gene silencing (VIGS) was carried out based on agroinfiltration method using tobacco rattle virus (TRV)–based vectors (51–53). The successful expression of TRV RNAs in agroinfiltrated seedlings was confirmed by RT-PCR detection of replicase (RdRp) and coat protein transcripts (Fig. S16). First, to confirm the effectiveness of VIGS, ApPDS silencing was carried out, developing the characteristic leaf photobleaching phenotype in VIGS seedlings (Fig. 7A) (53). The decreased expression of ApPDS transcript in leaves of VIGS seedlings further confirmed ApPDS silencing (Fig. 7B). Similarly, VIGS led to significant reduction (~60%) in ApUGT12 transcript level in leaves (Fig. 7C). HPLC analysis of leaf metabolites in ApUGT12-silenced seedlings revealed ~50% reduction in neoandrographolide content, suggesting an essential role of ApUGT12 in the biosynthesis of major diterpene C19-O-glucoside (Fig. 7, D and E). Interestingly, andrograpanin content in ApUGT12-silenced seedlings was increased significantly as compared with the vector control (Fig. 7, D and E). The increased accumulation of andrograpanin in VIGS seedlings could be due to the decreased rate of C19-O-glucosylation of andrograpanin to neoandrographolide because of ApUGT12 silencing. Furthermore, to examine whether reduced content of neoandrographolide in VIGS seedlings was solely because of ApUGT12 silencing, UGT73AU1 and UGT5 transcript levels were analyzed in ApUGT12-silenced seedlings (Fig. S17). However, unaltered transcript levels of UGT73AU1 and UGT5 in VIGS seedlings indicated that targeted silencing of ApUGT12 led to a reduction in neoandrographolide content in VIGS seedlings. Overall, these results suggest a major role of ApUGT12 in planta biosynthesis of neoandrographolide.

ApUGT12 is a member of the poorly characterized UGT86 family

UGTs are classified into different families and subfamilies considering about 45% and 60% sequence identities, respectively. Accordingly, the UGTs of Arabidopsis and rice were grouped into different UGT families/subfamilies such as UGT71-94, UGT96-99, and UGT701–UGT710 (https://prime.vetmed.wsu.edu/resources/udp-glucuronosyltransferase-homepage). To understand phylogenetic relationship of ApUGT12 with the classified UGTs, a phylogenetic analysis was conducted based on neighbor-joining method using Arabidopsis and rice UGTs (54). In the phylogenetic tree, ApUGT12 was grouped with the Arabidopsis and rice UGT86 family members (UGT86B1, UGT86A1, and UGT86A2) to which ApUGT12 showed 41 to 46% sequence identities (Fig. 8A). Accordingly, the UGT committee has assigned UGT86C11 nomenclature to ApUGT12. The biochemical function of Arabidopsis and rice UGT86 family members still remains unknown. So far, a single plant UGT86 family member (UGT86C10), having about 55% sequence identity with ApUGT12, has been biochemically characterized. UGT86C10 catalyzed glucosylation of C13-apocarotenoids in Mentha × piperita (55). In contrast, ApUGT12 showed only 23 to 31% sequence identity with UGT73AU1 and UGT5 (UGT74 member), which also catalyzed andrograpanin C19-O-glucosylation (42, 43). A comparison of amino acid sequences of ApUGT12, UGT73AU1, and UGT5 identified several conserved residues, besides the UDP-sugar binding plant secondary product glycosyltransferase motif and the catalytic histidine and aspartate residues, which might determine their substrate specificity (Fig. 8B) (56). Overall, the identification of ApUGT12 not only helped us to understand biosynthesis of medicinal diterpene glucosides in plant but...
also expanded our knowledge on biochemical function of a poorly characterized UGT86 family in plant specialized metabolism.

Discussion

GTs represent one of the largest enzyme families and typically account for 1 to 2% of the protein-coding genes in plants (6–8). The glycosylation of small molecules catalyzed by UGTs, the largest GT family in plants, plays crucial roles in plant development, metabolism, and stress tolerance (9, 10, 14–21, 57). In this work, we have identified a previously uncharacterized UGT86 member (ApUGT12/UGT86C11) that catalyzed C19-O-glucosylation with strict scaffold selectivity and is involved in developmental and tissue-specific biosynthesis of bioactive labdane diterpenes in the medicinal plant kalmegh. The catalytic property of recombinant ApUGT12 expressed in E. coli and N. benthamiana was very much similar to the native UGT activity detected in kalmegh (Figs. 3D, 5E, and S4). The recombinant ApUGT12 exhibited significantly higher catalytic efficiency (kcat/Km) using andrograpanin (19-hydroxy diterpene) than 14-deoxy-11,12-didehydroandrographolide (3,19-dihydroxy diterpene) (Table 1). However, ApUGT12 showed marginal activity using andrographolide (3,14,19-trihydroxy diterpene). It appears that the better catalytic efficiency of ApUGT12 using andrograpanin is likely because of the increased substrate affinity and increased rate of C19-O-glucosylation reaction. 14-Deoxy-11,12-didehydroandrographolide and andrographolide bear hydroxyl group(s) at C3 and/or C14 position, which are not found in andrograpanin (Fig. 1). Therefore, we cannot exclude the possibility that the hydroxyl group(s) at C3 and/or C14 in diterpene scaffold lead to a drastic change in rate of C19-O-glucosylation catalyzed by ApUGT12. The structure–function analysis of ApUGT12 might provide more detailed insights into scaffold-selective C19-O-glucosylation catalyzed by ApUGT12.

Two MeJA-responsive UGTs (UGT73AU1 and UGT5) were previously shown to catalyze in vitro C19-O-glucosylation of andrograpanin (42, 43). UGT73AU1 and UGT5 have marginal sequence identity (23–31%) with ApUGT12 and classified to different UGT families (UGT73 and UGT74) than ApUGT12 (UGT86). MeJA-inducible expression of UGT73AU1 and UGT5 transcripts could potentially contribute to MeJA-inducible neoandrographolide biosynthesis, but UGT73AU1 and UGT5 transcript expression patterns did not show obvious correlation with the developmental and tissue-specific biosynthesis of neoandrographolide (Figs. 2, A and B, S6, A and B, and S15, A and B). In contrast, ApUGT12 transcript expression was not only inducible by MeJA treatment but also showed a clear
correlation with the developmental and tissue-specific patterns of UGT activity and neoandrographolide content (Figs. 2, A and B, 6, A and B, and S15, A and B). Further substantiating these observations, ApUGT12 silencing led to altered profiles of diterpenes in kalmegh (Fig. 7, C–E). ApUGT12-silenced plants showed significantly depleted content of neoandrographolide and increased level of aglycone substrate andrograpanin, which was otherwise detected in a trace level, suggesting that andrograpanin is a native substrate of ApUGT12 and that in planta function of ApUGT12 cannot be compensated by the endogenous UGT73AU1 and UGT5 (Figs. 2A, 7, C–E, and S17). Moreover, the kinetic constants of recombinant ApUGT12 using various diterpene aglycones and the profiles of diterpene aglycones and C19-O-glucosides in ApUGT12-expressing tissues also suggested that andrograpanin is a preferred substrate of ApUGT12 in planta (Table 1 and Figs. 2A, 6A, and S3, A and B). Interestingly, andrograpanin showed promising bioactivities such as anti-inflammatory and antimicrobial properties, but, the development of andrograpanin-based products is so far compromised by the endogenous UGT73AU1 and UGT5 (Figs. 2A, 7, C–E, and S17).

Table 1

| Substrate/sugar donor | $K_m$ (μM) | $V_{max}$ (μmol min$^{-1}$ mg$^{-1}$) | $k_{cat}$ (S$^{-1}$) | $k_{cat}/K_m$ (M$^{-1}$ S$^{-1}$) |
|-----------------------|-----------|-------------------------------|-----------------|-------------------------------|
| Andrograpanin         | 137.3 ± 15.58 | 0.239 ± 0.007                  | 0.231 ± 0.007   | 1698.61 ± 153.91             |
| 14-Deoxy-11,12-didehydroandrographolide | 506.7 ± 197.94 | 0.017 ± 0.003                  | 0.016 ± 0.003   | 34.70 ± 6.39                 |
| UDP-glucose           | 271.16 ± 36.59 | 0.173 ± 0.006                  | 0.167 ± 0.005   | 622.80 ± 66.39               |

Bacterially expressed N-terminally 6×His-tagged ApUGT12 was purified and used for in vitro assay with sugar donor UDP-glucose and sugar acceptors (andrograpanin and 14-deoxy-11,12-didehydroandrographolide) as described in the Experimental procedures section. The data are the mean ± SD of three independent assays.
because of its unavailability in a sufficient quantity from kalmegh (58, 59). Hence, ApUGT12 silencing could be a useful approach to improve andrograpanin content in plants. Overall, these results indicated a major role of ApUGT12 in developmental and tissue-specific biosynthesis of major diterpene C19-O-glucoside, whereas multiple UGTs (ApUGT12, UGT73AU1, and UGT5) might contribute to MeJA-inducible biosynthesis of diterpene C19-O-glucoside in kalmegh. Similar to ApUGT12, ApCPS2 transcript also showed a clear correlation with tissue-specific biosynthesis of neoandrogropholide (40). ApCPS2 having chloroplast targeting peptide potentially operates in the chloroplast; however, ApUGT12 appears to function in the cytoplasm (Fig. 5, A–C). Therefore, the diterpene biosynthetic pathway of kalmegh potentially represents an example of specialized metabolic pathways, which are compartmentalized in multiple subcellular organelles and predominantly under the transcriptional regulation (2).

Labdane diterpenes with about 7000 known structures are a large family of natural products (27). The terpene synthases catalyzing the early steps in labdane diterpene biosynthesis were very well characterized, but the enzymes involved in the late-stage structural modifications such as those catalyzing regio-selective glycosylation were not well studied (60–63). Previously, UGT73 to UGT76 and UGT85 members were shown to catalyze labdane diterpene glycosylation in plants, but the role of UGT86 in diterpene glycosylation was not known before (17, 42, 43, 64). ApUGT12 is a new addition to the list of UGT families participating in diterpene glycosylation. Indeed, UGT86 represents one of the poorly studied UGT families in plants (7, 13). Until now, a single member of the UGT86 family (UGT86C10) was biochemically characterized (55). UGT86C10 catalyzing glucosylation of C13-apocarotenoids has quite a dissimilar substrate selectivity than ApUGT12, suggesting that UGT86 potentially plays diverse roles in plant specialized metabolism. It appears that ApUGT12 is very specific to the kalmegh labdane diterpene pathway and has rigid substrate selectivity because it could not glycosylate a range of other tested phytochemicals belonging to the terpene, phenylpropanoid, and phenolic classes, including steviol, another labdane diterpene (Fig. S13).

Unlike Arabidopsis and rice that encode only one to two UGT86 members, it appears that UGT86 family expanded remarkably in kalmegh, resulting in at least six UGT86 members (UGT86C7, UGT86C11, UGT86C12, UGT86E1, UGT86L1, and UGT86K1), which showed leaf-preferential transcript expression. Furthermore, studies on UGT86 family members in diverse plants could provide a more detailed information on the diversity of function they might play in plants. In conclusion, the discovery of ApUGT12 not only advanced our understanding of the biochemical function of UGTs in plant specialized metabolism but also opened up the prospect for genetic improvement of plants toward a specific diterpene chemotype.

**Experimental procedures**

**Plant materials**

Kalmegh plants (Cv. CIM-Megha) were grown in earthen pots under the natural light during the months of July to October, and samples were collected during different plant growth stages as described previously (40). MeJA (250 μM) treatment was given to 30-day-old plants as described previously (65). N. benthamiana was grown in Conviron A1000 plant growth chamber as described previously (66). Plant samples were flash frozen in liquid nitrogen and preserved at −80 °C until further use for isolation of proteins, metabolites, and RNA.

**RNA extraction and qRT-PCR analysis**

RNA isolation was done as described previously (67). RNA was cleaned up using RNeasy kit (Qiagen), DNaseI-treated, and processed for cDNA preparation using MultiScribe reverse transcriptase (Thermo Fisher Scientific). qRT–PCR was carried out in 7900HT Fast Real Time PCR using Power SYBR Green Master Mix (Thermo Fisher Scientific) with at least two technical replicates per cDNA sample. The amplification specificity was examined by analyzing melting curves at the dissociation step. The relative transcript expression was analyzed by following the 2−ΔΔct method. ApRSP4 and ApEF1-α were selected as the reference genes for normalization of qRT-PCR data based on an analysis of expression stability of the selected traditional reference genes including ApTUB, ApActin, ApPP2A, ApUBC, ApRSP4, and ApEF1-α (Fig. S18). Oligonucleotide sequences used in qRT-PCR analysis are listed in Table S2.

**Bacterial expression and purification of recombinant protein**

ApUGT coding sequences were PCR amplified from leaf cDNAs using Phusion high-fidelity DNA polymerase (Thermo Fisher Scientific) and gene-specific oligonucleotides (Table S2) and cloned into pET-28a (+). The integrity of the plasmid constructs was confirmed through sequencing using Big Dye terminator kit (Thermo Fisher Scientific). The confirmed plasmids were individually transformed into E. coli strain BL21 codon plus (DE3) RIPL. The transformants were precultured for overnight at 37 °C in LB media (5 ml) containing kanamycin (50 μg/ml), chloramphenicol (35 μg/ml), and streptomycin (100 μg/ml). The secondary culture was initiated using 0.01% of overnight grown primary culture and allowed to grow until an absorbance to 0.8 was reached at 600 nm. The culture was then incubated at 16 °C for 30 min, and after addition of 0.5 mM IPTG, it was further grown at 16 °C for overnight. Bacterial cells were harvested by centrifugation (2500g for 15 min) and washed with lysis buffer (50 mM Tris–Cl, pH 7.5, 150 mM NaCl, 10 mM MgCl2, and 10% glycerol). The cell pellet was resuspended in ice-cold lysis buffer supplemented with 10 mM imidazole, 1% Tween-20, 5 mM DTT, 0.1 mM PMSF, 1× EDTA-free protease inhibitor cocktail (Sigma–Aldrich), 0.5 mg/ml lysozyme, 2.5 units/ml benzozene (Merck Millipore), and incubated in ice for 30 min. Cell lysate was prepared by sonication for six to eight cycles (30 s ON and 99 s OFF at 50% amplitude). The soluble protein fraction was obtained after centrifugation (18,000g at 4 °C for 30 min) and subjected to affinity chromatography using Ni–NTA agarose (Qiagen). To prepare enriched fraction of recombinant UGT
for use in initial screening of UGT activity, the soluble protein of 100 ml bacterial culture was bound to pre-equilibrated Ni-NTA agarose (100 μl) for 2 to 3 h at 4 °C, passed through a 10 ml gravity column (Bio-Rad), washed with lysis buffer containing 20 mM imidazole, and eluted with lysis buffer containing 250 mM imidazole (68). For purification of ApUGT12 to electrophoretic homogeneity, soluble protein of 1 l bacterial culture was bound to pre-equilibrated Ni-NTA agarose (1 ml), passed through a 1 ml chromatography column (Bio-Rad) equipped with a peristaltic pump (Miclins India) and a fraction collector (Bio-Rad). The bound protein was washed in a step-wise gradient manner using lysis buffer containing 50 to 100 mM imidazole and eluted with lysis buffer containing 250 mM imidazole. The fractions of recombinant protein were pooled, desalted (50 mM Tris–Cl, pH 7.5, 150 mM NaCl, 10% glycerol, and 1 mM DTT), and concentrated using a 10 kDa cutoff centrifugal filter (Merck Millipore). The purity of protein was analyzed in SDS-PAGE with Coomassie blue stain, and protein concentration was determined following the Bradford method using bovine serum albumin (BSA) standard (69) and also by measuring UV absorption at an absorbance at 280 nm (Nanodrop spectro-photometer). The quantity of purified protein was also verified by densitometric analysis of SDS-PAGE gel with known quantity of BSA using ImageJ software (National Institutes of Health).

### Transient expression in N. benthamiana

ApUGT12 was cloned into pGWB441 and pGWB442 for N- and C-terminal YFP tagging. ApUGT12 was amplified using Phusion high-fidelity DNA polymerase with ORF-specific primers (Table S2), inserted into pENTR D-TOPO (Invitrogen), and finally cloned into pGWB441 and pGWB442 following LR clonase reaction (70). Empty pGWB441 and pGWB454 for expressing free YFP and mRFP were prepared by recombining empty pENTR/D-TOPO plasmid with pGWB441 and pGWB454 (68). The recombinant clones were selected in LB media containing spectinomycin (100 μg/ml). The clones were confirmed by sequencing and transformed into Agrobacterium tumefaciens (EHA105) following freeze-thaw method (71). For transient expression of recombinant proteins in N. benthamiana leaf, Agrobacterium suspension was prepared in infiltration buffer (10 mM MES, pH 5.6, 10 mM MgCl₂, and 200 μM acetoxyringone), and agro-infiltration method was followed as described previously using 5- to 6-week-old plants (66). P19, a post-transcriptional gene silencing suppressor, was coexpressed from pBin61-P19 plasmid along with recombinant ApUGT12 (49). After 36 to 48 h of agroinfiltration, leaf sections were analyzed, and images were captured under a Carl Zeiss LSM880 laser scanning confocal microscope using 63× (numerical aperture of 1.4) oil-immersion objective. YFP was excited at 514 nm and detected in a range of 525 to 562 nm, whereas RFP was excited at 561 nm and detected in a range of 570 to 652 nm (68). The simultaneous detection of RFP and YFP was done by combining the settings mentioned previously in the sequential scanning function of the microscope as per manufacturer’s instructions. Six days after agroinfiltration, leaves were infiltrated with andrograpanin for in planta UGT assay or processed for protein isolation.

### VIGS

ApUGT12 cDNA fragments from the 3’ end (472 bp) and 5’ end (419 bp) were PCR amplified using gene-specific primers and inserted in pTRV2 vector (Table S2). pTRV2-ApPDS constructs for VIGS were described previously (53). pTRV2 constructs were individually transformed into A. tumefaciens (GV3101). VIGS was performed in kalmegh seedlings at cotyledonary leaf stage (7–10 days old) as described previously with few modifications (53). Agrobacterium carrying pTRV1 and pTRV2 plasmids were cultured in 50 ml LB containing 10 mM MES, 20 μM acetoxyringone, 50 μg/ml kanamycin, and 25 μg/ml rifampicin until an absorbance at 600 nm reached to 1.8 to 2.0. Subsequently, Agrobacterium were resuspended in infiltration buffer (10 mM MES, pH 5.6, 200 μM acetoxyringone, and 10 mM MgCl₂) to achieve a final absorbance of 1.0 at 600 nm and incubated at room temperature for 2 to 4 h. Before agroinfiltration, Agrobacterium containing pTRV1 and pTRV2 plasmids were mixed in equal ratio. For the silencing of ApPDS and ApUGT12, Agrobacterium harboring pTRV2 construct targeting 3’ or 5’ region of ApPDS/ApUGT12 cDNAs were mixed in equal ratio before adding (equal proportion) to Agrobacterium carrying pTRV1. Vacuum infiltration of Agrobacterium into kalmegh seedlings was done as described previously (53). After agroinfiltration, excess liquid from seedlings was soaked in blotting paper, and seedlings were planted into soilrite mix, covered with polyethylene bags, and maintained in dark for 12 to 24 h. Seedlings were subsequently grown at 24 to 25 °C with a 16:8 h light:dark cycle in a glass house. The seedlings at 30 to 35 days of agroinfiltration frequently showed ApPDS-silencing phenotype in leaves, but the new leaves emerged thereafter did not always have silencing phenotype (53). Therefore, the leaves were harvested after 30 to 35 days of agroinfiltration, flash frozen in liquid nitrogen, and stored at −80 °C until further use.

### UGT assay

To isolate proteins from kalmegh tissues and ApUGT12-transformed N. benthamiana leaves for in vitro UGT assays, frozen samples (1 g) were ground to fine powder in liquid nitrogen using mortar and pestle, and protein was extracted in 5 ml precooled buffer (100 mM Tris–Cl, pH 7.5, 5 mM DTT, 150 mM NaCl, 5% glycerol, 0.1 mM PMSF, and 1× protease inhibitor cocktail) at 4 °C for 2 h. Soluble protein fraction was collected by centrifugation (18,000g at 4 °C for 30 min) and quantified in Bradford assays using BSA standard. Until otherwise mentioned, in vitro UGT assay was carried out in 50 μl assay buffer (10 mM Tris–Cl, pH 7.5, and 1 mM DTT) containing 250 μM sugar acceptor (andrograpanin, andrographolide, 14-deoxy-11,12-didehydroandrographolide, or other phytochemicals), 2 mM UDP-glucose, and 30 to 50 μg of total protein extract or 2.5 μg of Diterpene scaffold selectivity of UGT86C11
purified protein, at 35 °C for 15 to 60 min with continuous shaking at 200 rpm. The reaction was stopped by adding 200 μl ethyl acetate, the organic phase was collected by centrifugation (15,000g for 5 min), air dried, and finally dissolved in methanol for TLC analysis using chloroform:methanol:water (66:26:8) solvent mixture as a mobile phase (40) or processed for HPLC and LC–QTOF–MS analysis as described later. To determine optimum pH for ApUGT12 activity, reactions were set in 100 mM sodium acetate (pH 4.5–5.0), 100 mM MES–HCl (pH 5.5–6.5), 100 mM Tris–Cl (pH 7.0–8.5), or 100 mM sodium carbonate–bicarbonate buffer (pH 9.0–10.0). To know optimum temperature for ApUGT12 activity, the assays were performed at temperatures ranging from 20 to 50 °C in assay buffer. For determining the kinetic parameters, in vitro assays were done using 15 to 1500 μM andrograpanin or 14-deoxy-11,12-didehydroandrographolide, 2.5 mM UDP-glucose, and 2.5 μg purified ApUGT12 in 50 μl assay buffer at 35 °C for 15 min. The kinetic constants for UDP-glucose were determined in assays containing 30 to 2500 μM UDP-glucose and 500 μM andrograpanin. The product formed in the assays was estimated with a comparison to standard curve by HPLC analysis, and kinetic parameters (V_{max}, K_m, and k_cat) were determined by nonlinear regression analysis and fitting into Michaelis–Menten model using GraphPad Prism 9.1.2 (GraphPad Software, Inc) for Windows (www.graphpad.com). Because of unavailability of pure 14-deoxy-11,12-didehydroandrographolide standard, the rate of reaction using 14-deoxy-11,12-didehydroandrographolide was calculated based on the amount of substrate consumed in reactions. ApUGT12 activity toward varied phytochemicals (steviol, kaempferol, gallic acid, oleanolic acid, maslinic acid, arjunic acid, 11-keto-β-boswellic acid) was analyzed in assay buffer as described previously. In planta, ApUGT12 activity in N. benthamiana leaves was also done as described previously with few modifications (72). Six days after agroinfiltration, N. benthamiana leaf discs (2 cm diameter) were prepared and dipped in infiltration buffer containing 250 μM andrograpanin, and vacuum was applied (600 mm Hg for 3 min). Subsequently, leaf discs were placed on wet blotting paper in Petri dish and kept in plant growth chamber. After 24 h of incubation, leaf discs were washed thoroughly in pure water, flash frozen in liquid nitrogen, lyophilized, and stored at −80 °C until further use.

### Metabolite extraction and HPLC analysis

Flash-frozen plant samples were ground to fine powder in liquid nitrogen using pestle and mortar. The ground tissue (250 mg) was extracted twice with 5 ml methanol. The organic phase was collected by centrifugation (18,000g for 15 min), evaporated to dryness, and finally reconstituted in methanol. Lyophilized leaf discs were ground to fine powder in liquid nitrogen, and ground tissue (100 mg) was extracted with 5 ml methanol. HPLC separation of plant metabolites and UGT assay products was carried out in an isocratic elution mode using a Waters Spherisorb ODS2 column (250 × 4.6 mm, 5 μm particle size) and a Waters Alliance e2695 separation module HPLC system consisting of a 2998 photodiode array detector, autosampler, vacuum degasser, and quaternary pump (Waters) as described previously (73). The mobile phase consisted of a mixture of solvents: acetoniitrile (solvent A: 15%) and 60:40 methanol–water (solvent B: 85%) with 0.6 ml/min constant flow rate and 25 °C column temperature. The stock solutions (1 mg/ml) of andrographolide, andrographiside, andrograpanin, neoandrographolide, and 14-deoxy-11,12-didehydroandrographolide (Sigma–Aldrich and Natural Remedies) were made in methanol for standard curve preparation. Metabolite peaks were monitored at 200 to 220 nm considering retention time and UV spectra, and quantification was performed with a comparison to the standard curve. Steviol, kaempferol, gallic acid, arjunic acid, oleanolic acid, corosolic acid, maslinic acid, and 11-keto-β-boswellic acid were analyzed as per HPLC methods described previously (74–76).

### LC–QTOF–MS analysis

The Agilent 1290 Infinity II UPLC system coupled with an Agilent 6545A QTOF mass spectrometer was used for LC–QTOF–MS analysis. The UPLC system consisted of a solvent reservoir, a degasser, a G7120A binary pump, a G7130A column oven, a G7129B vial sampler, and a G4212B diode array detector. The mass spectrometer was based on an Agilent multimode ion source. The liquid chromatographic separation was performed considering gradient elution on a Zorbax Eclipse Plus C18 Rapid Resolution HD (2.1 × 50 mm, 1.8 μm) column with a set flow rate at 0.2 ml/min and temperature at 30 °C (77). The mobile phase consisted of 0.1% formic acid aqueous solution (solution A) and acetonitrile (solution B), and the gradient elution was carried out as follows: 0 to 3 min, 20% B; 3 to 4.5 min, 30% B; 4.5 to 6 min, 50% B; 6 to 8.5 min, 70% B; 8.5 to 10 min, 90% B; 10 to 12 min, 50% B; and 12 to 15 min, 10% B. Before sample analysis, the column was saturated using the mobile phase for at least 30 min. Diode array detector spectrum was recorded in the wavelength range from 190 to 600 nm with a peak width >0.1 min (2 s response time) at 2.5 Hz. The QTOF–MS was run in both positive and negative modes. The chromatographic and spectra data (.d) were obtained using Agilent Mass Hunter Data Acquisition software (version B.06.01). The operation conditions of the mass spectrometer were as follows: drying gas (nitrogen) temperature, 325 °C; drying gas flow rate, 10 l/min; nebulizer gas (nitrogen) pressure, 35 psi; capillary voltage, 2500 V; fragmentor voltage, 180 V; skimmer voltage, 45 V; and octopole radiofrequency voltage, 750 V. The data were acquired by MS1 mode. The MS scan range was 100 to 1700 m/z at a scan rate of 1.5 spectra/s. The mass spectrometer was calibrated and tuned before analysis for accuracy in the mass.

### Data availability

The sequence data are deposited in the GenBank with accession numbers MW589262-MW589284 (ApUGT1-ApUGT23) and KU516822 (ApPDS).

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**Supporting information**—This article contains supporting information (41).

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Diterpene scaffold selectivity of UGT86C11

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Abbreviations—The abbreviations used are: BSA, bovine serum albumin; cDNA, complementary DNA; EYFP, enhanced yellow fluorescent protein; GT, glycosyltransferase; MelA, methyl jasmonate; mRFP, monomeric red fluorescent protein; Ni–NTA, nickel–nitrilotriacetic acid; qRT–PCR, quantitative RT–PCR; QTOF, quadrupole TOF; TRV, tobacco rattle virus; UGT, UDP-glycosyltransferase; VIGS, virus-induced gene silencing.

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