Saponin Biosynthesis in *Saponaria vaccaria*. cDNAs Encoding β-Amyrin Synthase and a Triterpene Carboxylic Acid Glucosyltransferase1[OA]

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*Saponaria vaccaria* (Caryophyllaceae), a soapwort, known in western Canada as cowcockle, contains bioactive oleanane-type saponins similar to those found in soapbark tree (*Quillaja saponaria*; Rosaceae). To improve our understanding of the biosynthesis of these saponins, a combined polymerase chain reaction and expressed sequence tag approach was taken to identify the genes involved. A cDNA encoding a β-amyrin synthase (*SvBS*) was isolated by reverse transcription-polymerase chain reaction and characterized by expression in yeast (*Saccharomyces cerevisiae*). The *SvBS* gene is predominantly expressed in leaves. A *S. vaccaria* developing seed expressed sequence tag collection was developed and used for the isolation of a full-length cDNA bearing sequence similarity to ester-forming glucosyltransferases. The gene product of the cDNA, classified as UGT74M1, was expressed in *Escherichia coli*, purified, and identified as a triterpene carboxylic acid glucosyltransferase. UGT74M1 is expressed in roots and leaves and appears to be involved in monodesmoside biosynthesis in *S. vaccaria*.

*Saponaria vaccaria* (synonyms *Vaccaria segetalis*, *Vaccaria hispanica*; Caryophyllaceae) is an annual herb widely distributed in Asia, Europe, and other parts of the world. The seeds of this plant are known in traditional Chinese medicine as Wang-Bu-Liu-Xing, which is prescribed for the treatment of amenorrhea, breast infections, and the stimulation of lactation.

While relatively little is known about the later stages of saponin biosynthesis in *S. vaccaria*, the likely route to both mono- and bisdesmosides is represented in Figure 1. This is based on biochemical and molecular genetic work from other species (Abe et al., 1993; Hostettmann and Marston, 1995; Mackenzie et al., 1997; Vogt and Jones, 2000; Haralampidis et al., 2002). The saponins of the Caryophyllaceae family, such as those of *S. vaccaria*, are almost completely based on β-amyrin (see Fig. 1). The most common aglycones found in this family are quillaic acid, gypsogenic acid, and gypsogenin, which have hydroxy and carboxylate groups at C-3 and C-28, respectively (Jia et al., 2002). In *S. vaccaria*, the saponins can be divided into two groups, the monodesmosides that contain one ester-linked oligosaccharide, typically at C-28 of gypsogenic acid (e.g., vaccaroside B; Fig. 1; Koike et al., 1998) and the bisdesmosides that contain acetal- and ester-linked oligosaccharides, typically at C-3 and C-28, respectively, of quillaic acid (e.g., vaccaroside E; Fig. 1; Jia et al., 2002).

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cyclases (OSC; Abe et al., 1993; Haralampidis et al., 2002). Judging from the structure of the saponins that accumulate in *S. vaccaria*, the next steps in the pathway presumably include: (1) oxidation of β-amyrin at positions 16, 23, and/or 28; (2) glycosylation at position 28 and, for the major bisdesmosides, position 3; and (3) the acylation of sugars with acetyl and 2-hydroxy-2-methylglutaryl moieties. Apart from the obvious structural prerequisites (C-28 must be oxidized prior to glycosylation), little is known about the order of the reactions involved. As an example, studies in *Calendula officinalis* (Asteraceae) suggest that the sapogenin oleanolic acid (having a carboxyl at C-28) is formed by stepwise oxidation prior to glycosylation at C3 (Wilkomirski and Kasprzyk, 1979). It is possible that this is a general feature of saponin biosynthesis.

The enzymes involved in oxidation of β-amyrin may include cytochrome P450s and other hydroxylases,

![Figure 1](image_url)

*Figure 1.* Structure and biosynthesis of *S. vaccaria* saponins. A, Sapogenins of *S. vaccaria* and related compounds. B, Proposed biosynthesis in *S. vaccaria* of examples of a monodesmoside (vaccaroside B) and a bisdesmoside (vaccaroside E).
and alcohol and aldehyde dehydrogenases. Monodesmosides are presumably formed by stepwise transfer of activated monosaccharide donors. For *S. vaccaria*, the first transfer would be expected to be Glc to the carboxyl at C-28. As in a wide variety of glycosyltransferase reactions in nature, this reaction is likely to be catalyzed by an activated monosaccharide such as UDP-Glc, in this case forming an ester linkage (Vogt and Jones, 2000; Li et al., 2001). Additional sugars would then be transferred and in some cases acylated. For example, vaccaroside B has three additional Glc moieties, one of which is esterified with 2-hydroxy-2-methylglutarate (Fig. 1). Similarly, more complex schemes may be postulated for bisdesmosides.

As part of a broader study of the biochemical genetics of saponin biosynthesis in *S. vaccaria*, we report progress in understanding two of the steps shown in Figure 1 involved in monodesmoside formation through the identification and characterization of cDNAs encoding BAS and an ester-forming triterpene glucosyltransferase.

**RESULTS**

*S. vaccaria* BAS

Our investigation of the molecular genetics of saponin biosynthesis in *S. vaccaria* included BAS, which catalyzes the first committed step in the pathway. Degenerate oligonucleotide primers based on known plant BAS genes were used in reverse transcription (RT)-PCR experiments with RNA from germinating seeds as the template. The RACE method was used to obtain a full-length cDNA, as pDM057, corresponding to the gene designated *SvBS*. The *SvBS* open reading frame (ORF) consists of 2,283 nucleotides encoding a 760-amino acid protein of 87.5 kD. The *SvBS* amino acid sequence revealed 81%, 80%, and 72% identity with the BAS of *Glycyrrhiza glabra* (GenBank accession no. AB037203), *Medicago truncatula* (GenBank accession no. AJ430607), and *Arabidopsis* (*Arabidopsis thaliana*; GenBank accession no. NM106544), respectively. *SvBS* possesses the amino acid motif DCTAE, thought to form part of the active site of OSC (Abe et al., 1993; Abe and Prestwich, 1995) and the four QW motifs characteristic of the OSC superfamily (Poralla et al., 1994). In addition, *SvBS* amino acid sequence contains the Trp residue in the MWCYCR motif that plays an important role in the formation of β-amyrin in *Panax ginseng* (Kushiro et al., 2000).

The identity of the enzyme encoded by *SvBS* was confirmed by expression in yeast (*Saccharomyces cerevisiae*). Figure 2 shows the results of gas chromatography-mass spectrometry (GC-MS) analysis of extracts of the yeast strain MKP-0/pDM067. When compared with the control strain, MKP-0/pSCW231, extracts showed a single additional compound whose retention time and mass spectrum (GC-MS [electron impact {EI}] 70 eV, mass-to-charge ratio [m/z]; relative intensity): 218 [100], 203 [52], 69 [25], 95 [25], 35 [24], 135 [18], 189 [17], 426 [M]+ [3], 257 [2], 411 [M-CH3]+ [1]) are indistinguishable from authentic β-amyrin (Fig. 2). Thus, the *SvBS* gene product appears to be a BAS that presumably acts on the endogenous to yeast. Expression analysis by RT-PCR indicates that the *SvBS* gene is highly expressed in leaf and to a lesser extent in roots and germinating seeds (Fig. 3). The sequence of the *SvBS* cDNA was deposited in GenBank as accession number DQ915167.

A Triterpene Glucosyltransferase from *S. vaccaria*

The latter stages of saponin biosynthesis involve glycosylation of specific sapogenins. In *S. vaccaria*, the pathway appears to be split into two major routes

*Figure 2.* GC-MS analysis of the underivatized product of *SvBS* expressed in *S. cerevisiae*. Total ion chromatograms are shown for extracts of the *S. cerevisiae* strain MKP-0/pDM067, expressing *SvBS* (A) and the control strain MKP-0/pSCW231 (B). See “Materials and Methods” for details. Chromatograms represent equal volumes of *S. cerevisiae* cultures. Arrow indicates peak identified as β-amyrin.

*Figure 3.* Expression of *SvBS* and UGT74M1 relative to 18S rRNA in tissues of *S. vaccaria*. Results of RT-PCR are shown for RNA isolated from leaves (L), flowers (F), roots (R), germinating seeds (G), and developing seeds (D). See “Materials and Methods” for details. The sizes in base pairs of the amplified products are shown on the right.
directed toward mono- and bisdesmosides (Fig. 1). These are differentiated by both the identity of the aglycone and of the sugar esterified at C-28. The latter is predominantly Glc in monodesmosides and Fuc in bisdesmosides. To investigate this part of the pathway, a search was made for cDNAs encoding glycosyltransferases that could play a role. The rationale behind this expressed sequence tag (EST)-based approach is that most of the ESTs corresponding to genes involved in saponin biosynthesis will have sequences that readily reflect their enzyme class (e.g. glycosyltransferases). In addition, for compounds such as saponins, which are abundant in the tissue of interest, the relevant genes would be expected to be expressed at moderate to high levels and therefore be represented in moderately sized EST collections. Similar approaches have been used to isolate cDNAs encoding other enzymes of secondary metabolism (van de Loo et al., 1995; Cahoon and Kinney, 2004).

Analysis of 7,200 ESTs from a S. vaccaria developing seed library indicated that 10 ESTs in four groups showed similarity to plant glycosyltransferases containing the plant secondary product glucosyltransferase domain (Vogt and Jones, 2000). Given its similarity to gene-encoding, ester-forming glucosyltransferases (see below), pSv33B05, a singleton full-length cDNA representing one of the four groups, was investigated as a candidate for involvement in C-28 glycosylation in saponin biosynthesis (the investigation of the additional glycosyltransferase cDNAs, as well as other cDNAs, as candidates involved in saponin biosynthesis is in progress). The S. vaccaria gene corresponding to pSv33B05 was classified by Peter Mackenzie (Mackenzie et al., 1997) and given the name UGT74M1 (and the specific allele UGT74M1-1; see below).

DNA sequence analysis of pSv33B05 revealed an ORF corresponding to 478 amino acids and a predicted molecular mass of 53.3 kD. Southern hybridization results indicate that a single copy of UGT74M1 gene is present in the genome of S. vaccaria (data not shown). To identify possible introns in this gene, genomic DNA of S. vaccaria was used as a template for PCR with gene-specific primers corresponding to the 5'- and 3'-untranslated regions of the UGT74M1 gene. A product larger than expected from the cDNA was cloned into the vector pCR2.1-TOPO-TA and sequenced. It was found that this clone contained one intron of 354 bp corresponding to positions 712 to 1,065 in the genomic DNA sequence obtained (deposited in Figure 4.

**Figure 4.** Phylogenetic analysis of UGT74M1 and related sequences. An unrooted distance tree shows the relationship of UGT74M1 to various other glycosyltransferases. The function, species, and GenBank accession numbers of the sequences used are indicated in Table I. Boxes indicate ester-forming UGTs.

<table>
<thead>
<tr>
<th>Table I. Glycosyltransferase amino acid sequences used in Figure 4</th>
<th>Function</th>
<th>Species</th>
<th>GenBank No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anthocyanin Vhy</td>
<td>Anthocyanin 5-O-glucosyltransferase</td>
<td>Verbena × hybrida</td>
<td>BAA36423</td>
</tr>
<tr>
<td>Anthocyanin Phy</td>
<td>Anthocyanin 5-O-glucosyltransferase</td>
<td>Petunia × hybrida</td>
<td>BAA89009</td>
</tr>
<tr>
<td>Anthocyanin1 Pfr</td>
<td>Anthocyanin 5-O-glucosyltransferase</td>
<td>Perilla frutescens</td>
<td>BAA36422</td>
</tr>
<tr>
<td>Anthocyanin2 Pfr</td>
<td>Anthocyanin 5-O-glucosyltransferase</td>
<td>P. frutescens</td>
<td>BAA36421</td>
</tr>
<tr>
<td>Crocetin Csa</td>
<td>Crocetin glucosyltransferase</td>
<td>Crocus sativa</td>
<td>AAP94878</td>
</tr>
<tr>
<td>Diterpene Sre</td>
<td>Steviol glucosyltransferase</td>
<td>S. rebaudiana</td>
<td>ARB09290</td>
</tr>
<tr>
<td>IAA Zma</td>
<td>Indole-3-acetate glucosyltransferase</td>
<td>Zea mays</td>
<td>Q41819</td>
</tr>
<tr>
<td>Jasmonate Ath</td>
<td>Jasmonic acid glucosyltransferase</td>
<td>Arabidopsis</td>
<td>ABA39729</td>
</tr>
<tr>
<td>Limonoid Cun</td>
<td>Limonoid glucosyltransferase</td>
<td>Citrus unshiu</td>
<td>Q9M873</td>
</tr>
<tr>
<td>Monoterpenoid1 Epe</td>
<td>Monoterpene glucosyltransferase</td>
<td>Eucalyptus periniana</td>
<td>BAD9035</td>
</tr>
<tr>
<td>Monoterpene2 Epe</td>
<td>Monoterpene glucosyltransferase</td>
<td>E. periniana</td>
<td>BAD9034</td>
</tr>
<tr>
<td>Phenol Les</td>
<td>UDP-Xyl:phenolic glucosyltransferase</td>
<td>Lycopersicon esculentum</td>
<td>CA62049</td>
</tr>
<tr>
<td>Phenylpropanoid Nta</td>
<td>Phenylpropanoid O-glucosyltransferase</td>
<td>N. tabacum</td>
<td>BABB8935</td>
</tr>
<tr>
<td>Resveratrol Vla</td>
<td>Resveratrol/hydroxycinnamic acid O-glucosyltransferase</td>
<td>Vitis labrusca</td>
<td>ABH03018</td>
</tr>
<tr>
<td>Salicylate Nta</td>
<td>Salicylic acid glucosyltransferase</td>
<td>N. tabacum</td>
<td>AA61647</td>
</tr>
<tr>
<td>Thiohydroximate Bna</td>
<td>Thiohydroximate S-glucosyltransferase</td>
<td>B. napus</td>
<td>AAL09350</td>
</tr>
<tr>
<td>Unknown Ath</td>
<td>Unknown</td>
<td>Arabidopsis</td>
<td>NP172059</td>
</tr>
</tbody>
</table>


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GenBank as accession no. DQ915168). The position and phase of this intron matches that of introns in Arabidopsis genes corresponding to a subset of plant glucosyltransferases that has been called cluster L of Arabidopsis genes (Li et al., 2001). The relationship of UGT74M1 to other plant glucosyltransferases was also assessed through phylogenetic analysis (Fig. 4; Table I) of deduced amino acid sequences. UGT74M1 was found to lie within a clade that includes members of family 1, cluster L. A number of UDP-glucosyltransferases (UGTs) in this cluster are known to form ester or sulfur linkages. The known enzymes that showed the highest amino acid sequence similarity include Nicotiana tabacum salicylic acid glucosyltransferase (GenBank accession no. AAF61647), Brassica napus thiohydroximate glucosyltransferase (GenBank accession no. AAO06920), S. rebaudiana glucosyltransferase from Nicotiana tabacum (GenBank accession no. AAD30627); UGT74G1, UDP-glucosyltransferase 74G1 from S. rebaudiana (GenBank accession no. AAR06920).

Figure 5. Partial amino acid sequence alignment of UGT74M1 and related glycosyltransferases in the region of the polyAsn tract (AA175-216). UGT74M1, from this study; SA-GT, UDP-Glc:salicylic acid glucosyltransferase from N. tabacum (GenBank accession no. AA61647); UGT74E2, indole-3-acetate β-glucosyltransferase from Arabidopsis (GenBank accession no. AAD30627); UGT74G1, UDP-glucosyltransferase 74G1 from S. rebaudiana (GenBank accession no. AAR06920). The substrate specificity of UGT74M1

Table II. The substrate specificity of UGT74M1

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Relative Activity</th>
<th>C-23</th>
<th>C-16</th>
</tr>
</thead>
<tbody>
<tr>
<td>16α-Hydroxygypsogenic acid</td>
<td>177</td>
<td>COOH</td>
<td>OH</td>
</tr>
<tr>
<td>Gypsogenic acid</td>
<td>100</td>
<td>COOH</td>
<td>H</td>
</tr>
<tr>
<td>Gypsogenin</td>
<td>22</td>
<td>CHO</td>
<td>H</td>
</tr>
<tr>
<td>Quilliac acid</td>
<td>19</td>
<td>CHO</td>
<td>OH</td>
</tr>
<tr>
<td>Echinocystic acid</td>
<td>9</td>
<td>CH3</td>
<td>OH</td>
</tr>
<tr>
<td>Hederagenin</td>
<td>4.5</td>
<td>CH3OH</td>
<td>H</td>
</tr>
<tr>
<td>Betulinic acid</td>
<td>2.5</td>
<td>CH3</td>
<td>H</td>
</tr>
</tbody>
</table>
respectively. Although the enzyme was routinely as-
sayed with 10 mM MgCl₂, this was not required for
activity (data not shown).

The kinetic constants shown in Table III indicate that
16-hydroxygypsogenic acid was the most efficiently
converted to glucoside by the UGT74M1 gene product.
Gypsogenin and quillaic acid had apparent
\( K_m \) values comparable to 16-hydroxygypsogenic acid but lower
\( k_{cat} \) values. The presumed major substrate in
S. vac-
caria, gypsogenic acid, had a \( k_{cat} \) value similar to
16-hydroxygypsogenic acid but a higher apparent
\( K_m \).

In general, glucosylation of the sapogenins can lead
to the formation of Glc esters or acetals. The two types
of reaction products can be distinguished by alkaline
hydrolysis. The products obtained from the enzyme
assay using a variety of S. vaccaria-derived sapogenin
substrates were found to be unstable in the presence of
1 M KOH at 80°C for 2 h (data not shown). This
indicates that the product of the enzyme is a Glc ester
(see Fig. 1). This was confirmed for the product of
gypsogenin glucosylation by NMR (see Table IV). The
measured \(^1\)H-NMR spectrum of gypsogenin has signals at chemical shifts of 3.30 and 5.48 ppm, which,
based on previous NMR studies of gypsogenin glyco-
sides (Delay et al., 1997; Bouget-Bonnet et al., 2002),
can be assigned to C-18 and C-12, respectively. In

DISCUSSION

The cloning of cDNAs encoding BAS and UGT74M1
provides some insights into saponin biosynthesis in
S. vaccaria. The expression of the two genes appears to
be tissue specific but not tightly coordinated. For
example, some expression of SvBS is observed in
germinating seeds for which no UGT74M1 expression
was detected. Based on the observed expression levels
for UGT74M1, it is not surprising that it is represented
only once in the developing seed EST collection. Thus,
the molecular cloning of UGT74M1 reported appar-
ently corresponds to the isolation of a rare cDNA from
a rare mRNA.

The characterization of the UGT74M1 product in-
dicates that it is a triterpene carboxylic acid gluco-
syltransferase. In vitro, the enzyme is capable of
 glucosylating a variety of oleanane triterpenes as
well as having low activity with the lupane triterpe-
noid, betulinic acid. NMR analysis of the glucosylation
product of gypsogenin indicates that it forms a Glc
ester at C-28.

It is noteworthy to consider the activity of UGT74M1
in relation to the saponin profile of S. vaccaria seeds.
The monodesmosides consist primarily of the vaccaro-
sides A to D, having gypsogenic acid as the aglycone
and a Glc linked to the carboxyl at C-28. Alternatively,
the bisdesmosides (vaccarosides E–H), in addition to
GlcUA at C-3, have a Fuc esterified to C-28. Based on
our experiments with GDP-Fuc and a variety of agly-
cones, UGT74M1 does not appear to be involved in
making the Fuc ester linkage found in bisdesmosides.
It is possible, however, that, in this regard, the correct
combination of donor and acceptor was not tested.

### Table III. Kinetic constants for UGT74M1 and various sapogenins substrates

<table>
<thead>
<tr>
<th>Substrate</th>
<th>( K_m ) (( \mu )M)</th>
<th>( k_{cat} ) (s⁻¹)</th>
<th>( (k_{cat}/K_m) ) (s⁻¹·M⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gypsogenic acid</td>
<td>170</td>
<td>1.13</td>
<td>( 6.5 \times 10^4 )</td>
</tr>
<tr>
<td>16-OH gypsogenic acid</td>
<td>51</td>
<td>1.29</td>
<td>( 2.5 \times 10^4 )</td>
</tr>
<tr>
<td>Gypsogenin</td>
<td>42</td>
<td>0.125</td>
<td>( 3.0 \times 10^4 )</td>
</tr>
<tr>
<td>Quillaic acid</td>
<td>37</td>
<td>0.111</td>
<td>( 3.0 \times 10^4 )</td>
</tr>
</tbody>
</table>

*Constants for gypsogenic acid are considered estimates, because solubility limited the maximum concentration used to 100 \( \mu \)M.*
Thus, the data presented suggest that UGT74M1 is specific for UDP-Glc and involved in the formation of monodesmosides. It is possible that glucosylation of triterpene carboxylic acids represents a branchpoint in the biosynthesis of mono- and bisdesmosides.

UGT74M1 is a member of cluster L of family 1 GTs. This group includes a number of other ester-forming GTs involved in plant secondary metabolism and hormone metabolism (Li et al., 2001; Lim, 2005). It is notable that the legume *M. truncatula* contains oleanane saponins that have C-28 Glc ester moieties. While UGT71G1 and UGT73K1 have been reported to have activity on hederagenin, it is not clear that they are capable of catalyzing ester formation at C-28 (Achnine et al., 2005). Therefore, it is interesting to speculate that a GT more similar to UGT74M1 and other family 1, cluster L enzymes may be involved in saponin biosynthesis in legumes as well as other members of the Caryophyllaceae.

Clearly, further work is required to elucidate the enzymes and genes involved in other steps in the biosynthesis of saponins in *S. vaccaria*. Our EST collection, in combination with heterologous expression and other experiments, should provide an effective basis with which to uncover the enzymes that catalyze the oxidation, glycosylation, and acyl transfer steps involved. The oxidation of β-aminin to various saponins will be of particular interest and relevance to a number of saponin-producing taxa.

### Materials and Methods

#### Chemicals

α-Amyrin, β-aminin, echinocystic acid, erythrodiol, lupeol, oleanolic acid, and hederagenin were purchased from the Indofine Chemical Company. Asiatic acid, betulinic acid, cholesterol, caffeic acid, diosgenin, quercetin, and salicylic acid were obtained from Sigma-Aldrich. Spinasterol and benzoic acid were obtained from Chromadex and Fisher Scientific, respectively. Cyanidin was obtained from APIN Chemicals. 

N,N,O-bis(trimethylsilyl)-acetamide (BSA) was obtained from Aldrich.

#### Preparation of Gypsogenin and Mixture of Gypsogenic Acid, 16-Hydroxygypsogenic Acid, and Seco-Gypsogenic Acid from *Saponaria vaccaria* Saponins

Two hundred grams of *Saponaria vaccaria* seed was milled in a coffee grinder. The solids remaining after diethyl ether extraction were air dried and extracted twice with 70% methanol (aq, 700 mL) for 4 h at 50°C. The combined extract was concentrated in vacuo to about 100 mL and applied to an Amberchrom CG-300C (500 g) open column and made up in 10% methanol. The column was eluted with a methanol gradient from 20% to 100% in 20% increments and fractions collected, monitored by thin-layer chromatography, and checked by liquid chromatography (LC)-MS-photodiode array detection (PDA) for composition. Fractions obtained with 60% methanol were enriched in gypsogenic acid saponins such as vaccaroside B (molecular weight [MW] 1,278; see Fig. 1), while fractions obtained with 100% methanol were enriched in the gypsogenin saponin, segetoside H (MW 1,448). The appropriate fractions were combined and evaporated to dryness, affording segetoside H-enriched and vaccaroside B-enriched materials.

### Table IV. Assignment of selected 1H-NMR chemical shifts (given in parts per million) in the 3 to 6.5 ppm range for the gypsogenin glucoside product of UGT74M1

<table>
<thead>
<tr>
<th>Atom No.</th>
<th>Gypsogenin 3-O-Glucuronide</th>
<th>Gypsogenin</th>
<th>Gypsogenin Glucoside (UGT74M1 Product)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12</td>
<td>5.45</td>
<td>5.48(t)</td>
<td>5.48(t)</td>
</tr>
<tr>
<td>18</td>
<td>3.29</td>
<td>3.30(dd)</td>
<td>3.20(dd)</td>
</tr>
<tr>
<td>1’</td>
<td>4.87</td>
<td>–</td>
<td>6.36(d)</td>
</tr>
</tbody>
</table>

*aFrom Bouget-Bonnet et al. (2002).*
Gypsogenin

The segetoside H-enriched material from above (approximately 200 mg) was dissolved in 1.5 M HCl (7 mL) and EtOAc (1 mL) and heated to 90°C for 27 h. The reaction was cooled to ambient temperature and diluted with brine (20 mL). The pH was adjusted to approximately 5 with 1 M NaOH and citric acid, and the mixture was extracted with EtOAc (3 x 25 mL). The combined organic extract was washed with brine (1 x), dried (Na2SO4), and concentrated in vacuo. The residue was chromatographed on silica gel using EtOAc as eluent to afford a mixture consisting predominantly of seco-gypsogenic acid (MW 486) and 16-hydroxygypsogenic acid (MW 502).

Sapogenin Diacid Mixture

Vaccaroside B-enriched fractions were combined and evaporated to dryness. The residue (approximately 200 mg) was dissolved in 1 M NaOH and heated at 60°C for 5 h under a nitrogen atmosphere. The reaction mixture was cooled to ambient temperature and the pH adjusted to about 5 with 1 M citric acid. The mixture was diluted with brine (20 mL) and extracted with EtOAc (3 x 25 mL). The combined organic extract was washed with brine (1 x), dried (Na2SO4), and concentrated in vacuo. The residue was chromatographed on silica gel using diethyl ether (EtOAc)/C176 CH3CN, 0.12% CH3COOH to 35% CH3CN, 0.12% CH3COOH at a flow rate of 45 mL/min, 5 mm, 10 mm, 209 nm, and a fraction collector. A Phenomenex Gemini C18 column (250 x 10 mm, 5 μm particle size) was used with an elution gradient from 22.5% CH3CN, 0.12% CH3COOH to 35% CH3CN, 0.12% CH3COOH at a flow rate of 3 mL min⁻¹ for 30 min. One-minute fractions were collected and the fractions containing detected peaks were analyzed by GC-MS for presence of the sapogenins of interest. The fractions that contained individual sapogenins were combined, and solvent was removed by nitrogen stream and the sapogenins were extracted into ethyl acetate. Trimethylsilyl derivatives of the isolated sapogenins were prepared by addition to a mixture of 50% pyridine, 50% BSA before analysis. The purity of gypsogenic acid (as TMS derivative) GC-MS(EI) 70 eV, m/z (relative intensity): 73 (100), 203 (95), 202 (95), 143 (47), 342 (30), 294 (30), 585 [M-COOTMS]⁺ (15), 292 (14), 687 [M-CH3⁺]⁺ (11), 495 (9), 467 (8), 381 (3), 377 (3), 612 [M-TMSOH]⁺ (2), 702 [M⁺]⁺ (1); 16α-hydroxygypsogenic acid (as TMS derivative) GC-MS(EI) 70 eV, m/z (relative intensity): 73 (100), 147 (8), 201 (7), 129 (5), 725 (12), 719 (12), 790 [M⁺]⁺ (1), 775 [M-CH3⁺]⁺ (1), 700 [M-TMSOH]⁺ (1), 672 [M-HCOOTMS]⁺ (1), 610 (1), 583 (1), 493 (1), 393 (1); quillaic acid (as TMS derivative) GC-MS(EI) 70 eV, m/z (relative intensity): 73 (100), 187 (25), 143 (24), 129 (17), 275 (12), 585 [M-COOTMS]⁺ (10), 702 [M⁺]⁺ (8), 305 (12) [M-H]⁻ (1), 612 [M-TMSOH]⁺ (1), 393 (3), 495 (2), 687 [M-CH3⁺]⁺ (1); and gypsogenin (as TMS derivative) GC-MS(EI) 70eV, m/z (relative intensity): 73 (100), 203 (90), 202 (90), 143 (45), 189 (42), 119 (25), 496 [M-COOTMS]⁺ (17), 320 (17), 307 (6), 614 [M⁺]⁺ (1), 599 [M-CH3⁺]⁺ (1), 407 (1) was 95%, 94%, 82%, and 99%, respectively, by GC-MS.

Final Purification of Sapogenins

Final purification of each sapogenin (gypsogenic acid, 16β-hydroxygypsogenic acid, and quillaic acid from the diacid mixture) was accomplished by HPLC fractionation using an Agilent 1100 series HPLC with a quaternary pump, diode array detector monitoring at 209 nm, and a fraction collector. A Phenomenex Gemini C18 column (250 x 10 mm, 5 μm particle size) was used with an elution gradient from 22.5% CH3CN, 0.12% CH3COOH to 35% CH3CN, 0.12% CH3COOH at a flow rate of 3 mL min⁻¹ for 30 min. One-minute fractions were collected and the fractions containing detected peaks were analyzed by GC-MS for presence of the sapogenins of interest. The fractions that contained individual sapogenins were combined, and solvent was removed by nitrogen stream and the sapogenins were extracted into ethyl acetate. Trimethylsilyl derivatives of the isolated sapogenins were prepared by addition to a mixture of 50% pyridine, 50% BSA before analysis. The purity of gypsogenic acid (as TMS derivative) GC-MS(EI) 70 eV, m/z (relative intensity): 73 (100), 203 (95), 202 (95), 143 (47), 342 (30), 294 (30), 585 [M-COOTMS]⁺ (15), 292 (14), 687 [M-CH3⁺]⁺ (11), 495 (9), 467 (8), 381 (3), 377 (3), 612 [M-TMSOH]⁺ (2), 702 [M⁺]⁺ (1); 16α-hydroxygypsogenic acid (as TMS derivative) GC-MS(EI) 70 eV, m/z (relative intensity): 73 (100), 147 (8), 201 (7), 129 (5), 725 (12), 719 (12), 790 [M⁺]⁺ (1), 775 [M-CH3⁺]⁺ (1), 700 [M-TMSOH]⁺ (1), 672 [M-HCOOTMS]⁺ (1), 610 (1), 583 (1), 493 (1), 393 (1); quillaic acid (as TMS derivative) GC-MS(EI) 70 eV, m/z (relative intensity): 73 (100), 187 (25), 143 (24), 129 (17), 275 (12), 585 [M-COOTMS]⁺ (10), 702 [M⁺]⁺ (8), 305 (12) [M-H]⁻ (1), 612 [M-TMSOH]⁺ (1), 393 (3), 495 (2), 687 [M-CH3⁺]⁺ (1); and gypsogenin (as TMS derivative) GC-MS(EI) 70eV, m/z (relative intensity): 73 (100), 203 (90), 202 (90), 143 (45), 189 (42), 119 (25), 496 [M-COOTMS]⁺ (17), 320 (17), 307 (6), 614 [M⁺]⁺ (1), 599 [M-CH3⁺]⁺ (1), 407 (1) was 95%, 94%, 82%, and 99%, respectively, by GC-MS.

S. vaccaria cv. Pink Beauty seeds were obtained from CN Seeds. Plants were grown under 16 h light (150 μm s⁻²) at 22°C and 8 h dark at 16°C. 1°T.

Table V. Oligonucleotide primers used in this study

<table>
<thead>
<tr>
<th>Primer Name</th>
<th>Oligonucleotide sequence (5' to 3')</th>
</tr>
</thead>
<tbody>
<tr>
<td>BS-forward</td>
<td>GATTTTTGTAACACGTACGTAACAGCG</td>
</tr>
<tr>
<td>BS-reverse</td>
<td>CCTCGATCTCGGGTACGTTAGTTAGT</td>
</tr>
<tr>
<td>BS-forward1</td>
<td>GAATTCGACGAGATTTAGGCCGATTTAGT</td>
</tr>
<tr>
<td>BS-forward2</td>
<td>GCCGTCAGACCCGGCCTCGGATCAGGC</td>
</tr>
<tr>
<td>BS-reverse2</td>
<td>CGGGTCTCCAACACAGCGCCCAACATAC</td>
</tr>
<tr>
<td>GT33-F1</td>
<td>CCGTCTCGAGATGTGTCATAATGAAAAACACATGC</td>
</tr>
<tr>
<td>GT33-R1</td>
<td>CCGTCTGAGTTAAGAAGCCAGAGGCA</td>
</tr>
<tr>
<td>GT33-SEQ3</td>
<td>CAGCAGCGTCGGGCTCGTGTGGTTGTTGTA</td>
</tr>
<tr>
<td>GT33-SEQ4</td>
<td>GCACGAGCATGCCGGTTCGCGAGATTTC</td>
</tr>
<tr>
<td>GT33-SEQ5</td>
<td>ATGTGAATATTTTGTCACCATATGC</td>
</tr>
<tr>
<td>GT33-SEQ6</td>
<td>GCATATGGTCAATATTATCATATGAGGCTCTATTTAG</td>
</tr>
<tr>
<td>SQ4</td>
<td>A/G(A/C)/G(T/C)/G(C/G)/A/T/CT/TC/CC/CCCC/A/C/C/C/CC</td>
</tr>
</tbody>
</table>
Cloning of Putative BAS cDNA from *S. vaccaria*

Based on the highly conserved amino acid regions of known OSCs, four degenerate oligonucleotide primers were synthesized. The nucleotide sequences of these primers (SQ4, SQ5, SQ6, and SQ9) are shown in Table V. First, PCR with SQ5 and SQ4 primers corresponding to amino acid DGGWGLH1 and LSYSEGWWG, respectively, was carried out for 30 cycles (95°C, 1 min, 50°C, 1 min, 72°C, 1.5 min, and final extension at 72°C, 10 min) using cDNA from 7-to-10-d germinating seeds of *S. vaccaria* as a template. The product of the first PCR was applied to a QIAquick spin column (Qiagen) to remove the primers. Next PCR was carried out with SQ6 and SQ9 primers corresponding to amino acid SFLMMHPAK and EQAGAPEWA, respectively, with first purified PCR product (5 μL) as a template under the same conditions as the first PCR except the extension time at 72°C for 1 min. The expected size (800-900 bp) fragments were separated by electrophoresis (1.2% agarose gel) and purified using a QIAquick gel extraction kit (Qiagen). This DNA fragment was ligated into a pCR2.1-TOPO vector (Invitrogen). To obtain the entire sequences of **Sv**BS, the 5' and 3' regions were amplified separately using the Marathon cDNA Amplification kit (BD Biosciences CLONTECH) according to the manufacturer's instructions. The primer SQ10 was used to amplify the 3' region. The complete sequences were then amplified using specific primers BS-Forward and BS-Reverse and Vent polymerase (New England Biolabs). The resulting bands were gel purified, cloned into a pCR2.1-TOPO vector (Invitrogen) to give plasmid pDM057, and sequenced. The gene corresponding to pDM057 was designated **Sv**BS.

Functional Characterization of **Sv**BS

**SvBS** was characterized by expression in yeast (*Saccharomyces cerevisiae*). Two oligonucleotide primers (BS-Forward1 and BS-Reverse1), including EcoR1 sites to facilitate subsequent manipulation, were used to amplify the **Sv**BS coding region of pDM057 using Vent polymerase. After treatment with Taq polymerase and dATP, the amplified PCR product was directly ligated into the pCR2.1-TOPO vector (Invitrogen). The plasmid was then digested with EcoR1 and ligated into pSCW231 yeast expression vector to generate the plasmid pDM067. The DNA sequence of the insert was confirmed to be identical to that of pDM057 and in the sense orientation relative to the ADH1 promoter.

The yeast strain MKP-0 (MALL cant-100 ade2-1 his3-1 leu3-3 111 his3-Δ200 trp-Δ301) (covello and reed, 1996) was separately transformed with pSCW231 and pDM067 by the lithium acetate method (giertz et al., 1992) and selected on minimal agar plates lacking Trp. The resulting yeast strains were designated MKP-0/pSCW231 and MKP-0/pDM067. For assessment of enzyme activity, recombinant yeast cells were grown until stationary phase in 50 mL at 28°C on minimal medium (synthetic dropout) lacking Trp. Mkp-0 yeast containing the empty plasmid pSCW231 was used as a negative control.

For analysis of **Sv**BS products in yeast, the cells of 50 mL of saturated cultures were collected and saponified with 2 mL 10% KOH/methanol at 80°C for 1 h. After extraction with the same volume of hexane and water, the extract was dried and the residue was dissolved in 100 μL of BSA (Aldrich)/pyridine (1:1). GC-MS analysis was carried out using DB-5MS column (J&W Scientific), as described previously (li et al., 2006).

**DNA Extraction and Southern-Blot Analysis**

Genomic DNA was isolated from leaves of *S. vaccaria*, essentially as described previously (bekesiowa et al., 1999). Southern-blot analyses were carried out using standard methods. Ten micrograms of *S. vaccaria* genomic DNA were digested with EcoR1, EcoRv, or HindIII (Biolab), resolved on 1% agarose gel, and then transferred to HybondN membrane (amersham biosciences). This was followed by hybridization at 65°C for 2 h with a 715-bp NcoI cDNA probe that had been radiolabeled with c-13° PdCTP using a Random Primers DNA Labeling kit (Invitrogen).

This fragment containing a partial **Sv**GT1 **G**DNA fragment was obtained from the digestion of pDM060. The filter was washed once in 2× sodium chloride/sodium phosphate/EDTA (SSPE, 0.1% SDS for 10 min then 1× SSPE, 0.1% SDS for 15 and 0.1× SSPE, 0.1% SDS for 10 min at the hybridization temperature. The blot was exposed to Super Rx Fujii Medical x-ray film for 3 d.

**RNA Isolation**

Total RNA was isolated from field-grown *S. vaccaria*. The RNeasy Plant Mini kit (Qiagen) was used for the total RNA isolation from leaves, flowers, roots, and germinating seeds. For developing seeds, RNA was first isolated by the method of Wang and Vodkin (1994) prior to use of the RNeasy Plant Mini kit. Genomic DNA contamination was eliminated by on-column DNase digestion step with RNase-free DNase set (Qiagen).

**Relative Expression of **UGT74M1** and **Sv**BS by RT-PCR**

To investigate gene expression by RT-PCR, first-strand cDNA was synthesized from 5 μg of total RNA using Thermoscript RT-PCR system in a 20-μL reaction with random primers (Invitrogen) according to the manufacturer's instructions. Two microliters of the first-strand reaction was then used as a template for PCR amplification using Platinum Taq DNA Polymerase (Invitrogen). The specific primers BS-Forward2 and BS-Reverse2 were used for the amplification of **Sv**BS, and GT33-SEQ3 and GT33-SEQ4 were used for **UGT74M1**. The 18S PCR primer pair (universal 18s internal standards, Amboim) was used for the amplification as an internal control. The PCR was carried out with 3 min at 95°C, followed by 26 cycles of 30 s at 95°C, 30 s at 55°C, and 40 s at 72°C for **Sv**BS and 35 cycles for **UGT74M1**. The PCR products were then analyzed on a 1.5% agarose gel.

**Phylogenetic Analysis**

Using BLASTP to search public databases maintained at the National Center for Biotechnology Information, amino acid sequences with known function and similarity to **UGT74M1** were identified. With software hosted at the European Bioinformatics Institute (chenna et al., 2003), amino acid sequences encoding glycosyltransferases were aligned using ClustalW using default parameters including the Gonnet scoring matrix, a gap penalty of 10, and a gap extension penalty of 0.2. The resulting alignment was used to generate an unrooted phylogenetic tree using the neighbor joining method. The tree was visualized using TREEVIEW (PAG, 1996).

**UGT74M1 PolyAsn Variants**

To test for genetic variation in the polyAsn tract of **UGT74M1**, full-length cDNAs were cloned by RT-PCR using total RNA from 7-to-10-d germinating seeds of *S. vaccaria* as a template. The primers GT33-F1 and GT33-R1 (Table V) were used to PCR amplify a DNA fragment of the **UGT74M1** ORF under the following conditions: 3 min at 95°C, followed by 30 cycles of 30 s at 95°C, 60 s at 57°C, and 90 s at 72°C and one cycle of 10 min at 72°C using BD Sprint Advantage PCR kit (clontech). The PCR products were then cloned into pCR2.1-TOPO vector. Twelve individual plasmids were chosen for sequencing. The plasmid that contained the allele with 12 contiguous Asn codons in **UGT74M1** was named **pDM060**.

To construct a **UGT74M1** clone in which the polyAsn tract is deleted, the Gene Splicing by Overlay Extension method (horton et al., 1990) was used. The primers used for the upper fragment were GT33-F1 and GT33-SEQ5 and for the lower fragment were GT33-R1 and GT33-SEQ6 using pDM060 as the template. The pDM060 was originally from the clone **pSv**338605, obtained from a developing seed library of *S. vaccaria*. To obtain the corresponding full-length cDNA, PCR was carried out with the primers GT33-F1 and GT33-R1 and the upper and lower fragments as templates. The resulting PCR product was cloned into pCR2.1-TOPO vector, digested with Xhol, and ligated into the Escherichia coli expression vector pET14b to generate the plasmid pDM066. To obtain the full-length cDNAs of **Asn**<sub>14</sub>-containing **UGT74M1** alleles, PCR was carried out with the primers GT33-F1 and GT33-R1 using pDM060 and pDM065, respectively. The resulting PCR products were cloned into pCR2.1-TOPO and digested with Xhol. Both fragments were cloned into pET14b to generate the plasmid pDM064 and pDM065 for **Asn**<sub>14</sub>-containing **UGT74M1** alleles, respectively. The DNA sequences of the insert for pDM066, pDM064, and pDM065 were confirmed to be identical to that of the original plasmids and in the sense orientation relative to the T7 promoter.

**Expression of and Purification Recombinant **UGT74M1****

Single colonies of the *E. coli* strain Rosetta2Pls**Sv**pDM064 were used to inoculate 10 mL. Luria-Bertani medium containing ampicillin (100 μg/mL) and chloramphenicol (34 μg/mL) at 37°C overnight. The fresh culture was used to inoculate 50 volumes of Luria-Bertani medium containing the same antibiotics. Bacteria were grown at 37°C to OD<sub>600</sub> = 0.5 to 1.0. Induction was achieved by addition of 0.4 mM isopropyl-β-d-thiogalactopyranoside. The
cells were maintained overnight at 30°C, harvested by centrifugation (5,000 rpm, 15 min), and stored at –80°C.

Cell pellets were suspended in buffer A (20 mM sodium phosphate, pH 7.4, 10% [v/v] glycerol, 0.5 mM Nicotinamide, 10% bovine serum albumin) containing 20 µM EDTA, 50% glycerol, 0.1 M dithiothreitol, and disrupted using a French press. The cell lysate was centrifuged and the supernatant used to reduce the solubilizing recombining enzyme was passed through a 1 mL HisTrap FF Crude column (Amersham Biosciences) packed with precharged Ni Sepharose 6 FastFlow. After washing with 20 mM volumes, the bound enzyme was eluted with buffer A containing 300 mM imidazole. One-millilitre fractions were collected, and those containing UGT74M1 activity were pooled and concentrated by ultrafiltration (Amicon Ultra-4, 50 K NMWL; Millipore). The protein solution was subsequently applied to Bio-Sil TSK 250 HPLC gel filtration column (300 mm × 7.5 mm; Bio-Rad) at a flow rate of 1 mL min⁻¹ using 20 mM HEPES, pH 7.5, and 10% glycerol as eluent. Five hundred-microlitre fractions were collected, and those corresponding to OD_{280} peaks were assayed for UGT74M1 activity. The fraction showing the highest activity was frozen in aliquots (50 µL) and stored at –80°C. Protein concentration was determined according to Bradford (1976) using bovine serum albumin as standard.

Enzyme Reactions

Unless otherwise stated, glucosyltransferase enzyme reactions were carried out in 100 µL containing 0.6 to 0.8 mg/mL of purified recombinant protein, 250 µM UDP-Glc, 1 mM acceptor substrate, 100 mM HEPES, pH 7.5, 15% glycerol, 1 mM dithiothreitol, and 10% MgCl₂. The reaction mixture was incubated at 30°C for 10 min and stopped by the addition of 1 mL ethyl acetate. For radiochemical assays, the reaction mixture included UDP-[U-¹⁴C]Glc (Amersham; final specific activity, 37 kBq/nmol), UDP-[U-¹⁴C]GlcUA (Amersham; specific activity, 15.47 GBq/nmol), or GDP-[U-¹⁴C]Fuc (Perkin Elmer; specific activity, 37 kBq/nmol). The radioactivities were measured using a Rackbeta liquid scintillation counter.

The pH optimum of UGT74M1 was evaluated using the radiochemical assay and one mM UDP-Glc from pH 5.0 to 10.0 using five buffer systems: 100 mM MES-NaOH, pH 5.0 to 7.0; 100 mM MOPS-NaOH, pH 6.0 to 8.0; 100 mM HEPES-NaOH, pH 6.5 to 8.0; 100 mM Tris-HCl, pH 7.0 to 9.0; and 100 mM 2-(cyclohexylamino)ethanesulfonic acid (CHES)-NaOH, pH 8.5 to 10.0. Similarly, the optimal temperature was evaluated from 20°C to 60°C in 10°C intervals at pH 7.5.

For kinetic studies, the radiochemical assay was used, and substrate concentrations and reaction times were varied for UDP-Glc (10–100 µM, 10 min), UDP-GlcUA (10–100 µM, 30 min), 16-hydroxy-UDP-Glc (10–500 µM, 5 min), and quilliaic acid (10–500 µM, 30 min). The kinetic constants were estimated from Lineweaver-Burke plots using the average of triplicate measurements. The K_m values were calculated using the predicted molecular mass of 53,352 g mol⁻¹.

In some cases, unlabeled UDP-Glc was used, and the extracted products were concentrated and subjected to LC-MS (see below). For product analysis by NMR, a 20-ml reaction mixture containing 1 mM gypsogenin was incubated overnight and extracted twice with 50 mL ethyl acetate. After evaporation of the solvent, the reaction mixture was dissolved in pyridine-CH₃CN, 0.12% CH₃COOH over 30 min at a flow rate of 0.2 mL min⁻¹. An elution fraction corresponding to a peak with a retention time of 23 min was evaporated to complete dryness, and after dissolving in pyridine-CH₃CN, the proton NMR spectrum was recorded on a Bruker Avance DRX 500 MHz spectrometer equipped with a CryoProbe.

LC-MS

A 2695 Alliance chromatography system, coupled to a ZQ mass detector and a 2996 photodiode array detector (Waters) was used for LC-MS-PDA analysis. A Waters SunFire 3.5-µm RP C₁₇, 150 × 2.1 mm at 35°C with a flow rate of 0.2 mL/min was used. The binary solvent system consisted of 90:10 (v/v) water/acetonitrile containing 0.12% acetic acid (solvent A) and acetonitrile containing 0.12% acetic acid (solvent B). The gradient program used was: 0 to 3 min, 75:25 A:B; 3 to 28 min, 75:25 to 50:50 A:B; 28 to 31 min, 50:50 to 100:0 A:B; and 31 to 39 min, 100:0 A:B. Mass analysis was performed using negative electrospray ionization under the following conditions: capillary, 2.70 kV; cone ramped, from –20 to –60 V; extractor, –3.50 V; RF lens, –0.7 V.

GC-MS

GC-MS analysis was performed using an Agilent 6890 GC equipped with an autosampler split 30:1 onto a DB-5MS column (30 m × 0.25 mm i.d., J&W Scientific), which was temperature programmed from 125°C to 300°C at 5°C min⁻¹. The column was connected to a mass selective detector (Agilent 5973) operating under standard EI conditions.

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession numbers DQ915167 and DQ915168.

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