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53 Abstract

Catabolism of sulfoquinovose (SQ, 6-deoxy-6-sulfoglucose), the ubiquitous sulfosugar produced by photosynthetic organisms, is an important component of the biogeochemical carbon and sulfur cycles. Here, we describe a new pathway for SQ degradation that involves oxidative desulfurization to release sulfite and enable utilization of the entire carbon skeleton of the sugar to support the growth of the plant pathogen Agrobacterium tumefaciens. SQ or its glycoside sulfoquinovosyl glycerol (SQGro) are imported into the cell by an ABC transporter system with an associated SQ binding protein. A sulfoquinovosidase hydrolyses the SQ glycoside and the liberated SQ is acted on by a flavin mononucleotide-dependent sulfoquinovose monooxygenase, in concert with an NADHdependent flavin reductase, to release sulfite and 6-oxo-glucose. An NADPH-dependent oxidoreductase reduces the 6-oxo-glucose to glucose, enabling entry into primary metabolic pathways. Structural and biochemical studies provide detailed insights into the recognition of key metabolites by proteins in this pathway. Bioinformatic analyses reveal that the sulfoquinovose monooxygenase (smo) pathway is distributed across Alpha- and Betaproteobacteria and is especially prevalent within the Rhizobiales order. This strategy for SQ catabolism is distinct from previously described pathways as it enables the complete utilization of all carbons within SQ by a single organism with concomitant production of inorganic sulfite.

Significance Statement

Sulfoquinovose, a sulfosugar derivative of glucose, is produced by most photosynthetic organisms and contains up to half of all sulfur in the biosphere. Several pathways for its breakdown are known, though they provide access to only half of the carbon in sulfoquinovose and none of its sulfur. Here, we describe a fundamentally different pathway within the plant pathogen *Agrobacterium tumefaciens* that features oxidative desulfurization of sulfoquinovose to access all carbon and sulfur within the molecule. Biochemical and structural analyses of the pathway's key proteins provided insights how the sulfosugar is recognized and degraded. Genes encoding this sulfoquinovose monooxygenase pathway are present in many plant pathogens and symbionts, alluding to a possible role for sulfoquinovose in plant host–bacteria interactions.

Introduction

Sulfoquinovose (SQ; 6-deoxy-6-sulfoglucose) is an anionic sulfosugar found in plant and cyanobacterial sulfolipids, and in S-layer proteins in archaea (1). It is estimated that SQ holds around half of all sulfur in the biosphere, with 10 billion tonnes produced each year in Nature, and so its cycling is a significant component of the biogeochemical sulfur cycle (2). SQ is primarily found as the headgroup of the plant sulfolipid sulfoquinovosyl diacylglycerol, and its close association with photosynthetic membranes and proteins supports roles in their structure and function (3). Microbial communities play a dominant role in SQ cycling and usually more than one organism is required to completely assimilate this source of carbon and sulfur. Organisms with a tier 1 pathway, termed sulfoglycolysis, perform scission of the C3-C4 bond of SQ to give two threecarbon fragments; carbons 1-3 enter central metabolism, while carbons 4-6 bearing the sulfonate are excreted as dihydroxypropanesulfonate (DHPS) or sulfolactate (SL). Organisms with a tier 2 pathway are those that process DHPS and SL to access the remaining three carbon fragment and release inorganic sulfur. To date, three tier 1 pathways have been described: the sulfoglycolytic Embden-Meyerhof-Parnas (sulfo-EMP) (4), Entner-Doudoroff (sulfo-ED) (5, 6) and sulfofructose transaldolase (sulfo-SFT) pathways (7, 8). Tier 2 metabolism has been described for various specialized bacteria that utilize SL or DHPS and perform 'biomineralization' to release inorganic sulfite, which under aerobic conditions is readily oxidized to sulfate (1). While many of the steps in the three tier 1 sulfoglycolysis pathways differ, all three pathways share the presence of a specialized glycoside hydrolase, a sulfoquinovosidase (SQase), which catalyzes the hydrolysis of SQ glycosides, such as SQGro, to release SQ (9, 10).

While the tier 1 and 2 pathways described to date require two or more organisms to complete the 'biomineralization' of SQ, there is some evidence that this can also be accomplished by a single organism. Roy and co-workers have reported that an *Agrobacterium* strain from soil can completely consume SQ, with release of sulfate, although the genetic and biochemical details behind this process were not investigated (11). We previously reported that *A. tumefaciens* C58 contains a functional SQase, with the ability to hydrolyze SQGro (9). However, analysis of its genome did not reveal any genes homologous to those expected for known tier 1 sulfoglycolysis pathways.

Here, we investigate the 'biomineralization' of SQ by *Agrobacterium tumefaciens* (*Agrobacterium fabrum*) strain C58 and show that this organism effects the oxidoreductive desulfurization of SQ to release inorganic sulfite and glucose, which can feed into primary metabolism. We show that this pathway involves: a novel SQ/SQGro solute binding protein and associated ATP-binding cassette (ABC) transporter; an SQase to release SQ from its glycosides; a flavin-dependent SQ

monooxygenase with paired flavin-reductase to effect oxidative desulfurization of SQ to sulfite and 6-oxo-glucose; and a NADPH-dependent oxidoreductase to reduce 6-oxo-glucose to glucose. X-ray structures determined for each of these proteins in complex with relevant metabolites reveal the molecular basis of substrate binding and catalysis. We show through bioinformatics analyses that this pathway – the first to enable the complete assimilation of SQ – is distributed across Alpha- and Betaproteobacteria and is particularly well-represented within the Rhizobiales order.

Results

Differential expression of a gene cluster in the presence sulfoquinovose

To determine if A. tumefaciens C58 can utilize SQ as a carbon source, we attempted to grow this organism in M9 minimal media containing SQ as the sole carbon source. A. tumefaciens C58 exhibited robust growth in this media and analysis of spent culture supernatant failed to detect DHPS or SL. Instead, the culture supernatant accumulated sulfate, but with a lag between consumption of SQ and sulfate release (Fig. 1a, Fig. S1), as was previously reported by Roy and co-workers for Agrobacterium sp. strain ABR2 (11). Noting that sulfite is generally released from organosulfonate degradation pathways (1, 12), we analyzed the supernatant for sulfite (SO₃²⁻), and observed that SQ consumption is coincident with production of sulfite, which slowly undergoes autooxidation to sulfate. To investigate the metabolism of the carbon skeleton of SQ, we cultured A. tumefaciens on ¹³C₆-SQ (13) and analyzed the culture supernatant using ¹³C NMR spectroscopy (Fig. S2). The only significant ¹³C-labelled product we could detect was ¹³C-bicarbonate, which formed transiently during exponential phase growth, and the ¹³C-labelled bicarbonate signal disappeared at stationary phase, presumably through exchange with atmospheric CO₂. A. tumefaciens grew on other sulfoquinovosides, including SQGro and methyl α-sulfoquinovoside (MeSQ), but did not grow on other alkylsulfonates including DHPS, SL, sulfoacetic acid, taurine, pentanesulfonate, MES, MOPS, HEPES, PIPES, cysteic acid or methanesulfonic acid (Fig. S3). Collectively, this data demonstrates that A. tumefaciens effects the complete metabolism of the carbon backbone of SQ with concomitant release of sulfite.

We performed comparative proteomic experiments to identify changes associated with the growth of *A. tumefaciens* on SQ compared to glucose at mid-log phase (**Fig. 1b**). The largest and most significant change we observed was an increase in the abundance of proteins encoded by a single cluster of genes (*Atu3277-Atu3285*) for cells grown on SQ. Proteins encoded by *Atu3283* and *Atu3284* were not observed; however, they are predicted to be integral membrane proteins that can be difficult to detect using conventional proteomic workflows (14). Thus, the gene cluster *Atu3277-Atu3285*, which was subsequently renamed *smoA-smoI*, appeared to be important for growth on SQ (**Fig. 1c**). While the protein encoded by *Atu3285* was previously identified as an SQase (9), the proteins encoded by other genes in the cluster were not annotated with functions that were consistent with any tier 1 pathway, suggesting that *A. tumefaciens* uses a different approach for the catabolism of SQ. The automated annotations ascribed to the respective gene products in the cluster, which included a putative ABC transporter system, sulfonate monooxygenase, SDR oxidoreductase, flavin reductase and exporters, enabled development of a hypothetical biochemical pathway that could explain the complete assimilation of SQ by *A. tumefaciens* (**Fig. 1d**). We

proceeded to biochemically validate this hypothesis and gain structural insights into the proteins involved.

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Atu3282 (smoF) encodes an ABC transporter solute-binding protein that binds SQGro

- Within the gene cluster identified through proteomics, Atu3281 (smoE), Atu3283 (smoG), and Atu3284 (smoH) were annotated as an ABC transporter system, with Atu3282 (smoF) encoding an associated periplasmic solute-binding protein. The substrate preferences of solute binding proteins are useful for assigning functions to their associated ABC transporters (15). Accordingly, we produced recombinant SmoF (**Fig. S4**) and demonstrated that it binds SQGro with $K_d = 0.29\pm0.17$ μ M ($\Delta H = -11\pm0.4$ kcal mol⁻¹, $\Delta S = -7\pm2$ cal mol⁻¹ deg⁻¹) (**Fig. 2a, Fig. S5, Table S3**). No binding
- was observed for the stereochemically-related monosaccharides D-glucose and D-glucuronic acid.

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To delineate how SmoF recognizes its ligand, we used X-ray diffraction methods to obtain a high-172 173 resolution 3D structure of SmoF in its ligand-free apo state and in complex with SQGro (Fig. 2b, 174 Table S4). Like most ABC transporter solute-binding proteins, SmoF possesses two globular 175 domains with a similar α/β fold forming a deep cleft lined with aromatic and polar residues to capture the ligand. Comparisons of the structures for ligand-free SmoF and the SQGro complex 176 177 revealed a large conformational change in the protein resulting from inter-domain rotation upon SQGro binding. The relative movement of domains was assessed using the DynDom server, which 178 179 indicated a hinge rotation of 31° about four linker regions connecting the two domains (Fig. S6). SOGro is buried deep within the inter-domain cleft and residues from both domains accommodate 180 181 this ligand through a network of hydrogen-bonding interactions (Fig. 2c,d). The sulfonate of SQGro, which is the defining feature of this sulfosugar, is accommodated by hydrogen-bonds to the 182 183 side-chain of Thr220 (2.6 Å), backbone amides of Gly166 (3 Å) and Ser43 (2.8 Å), and an ordered water molecule that in turn hydrogen-bonds to the sidechain of His13 (3 Å) and Gln46 (3.2 Å) (Fig. 184 185 2c,d). These and the other interactions in the SQGro-bound 'closed' state stabilized SmoF substantially, as evidenced by a 15 °C increase in the protein melting temperature (Fig. S7). 186

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The structural basis of SQGro recognition by the SQase Atu3285 (SmoI)

We previously reported that *Atu3285 (smoI)* encodes an SQase that preferentially hydrolyses 2'*R*SQGro, the natural stereoisomer of this glycoside (9). To understand the molecular basis of the
preference SmoI has for this stereoisomer, we determined the 3D structure of a pseudo-Michaelis
complex: the inactive acid/base mutant SmoI-D455N in complex with 2'*R*-SQGro (**Fig. 2e,f**). SmoID455N•SQGro crystallized with four protomers in the asymmetric unit, each showing unambiguous
density of the substrate bound at the active site. As described previously, the overall fold is an

195 (α/β)₈ barrel appended with small β sheet domain and the sulfonate group is recognized by 196 Arg283/Trp286/Tyr491 triad⁸. Arg438 and Glu135 make hydrogen-bonding interactions with the 197 glyceryl aglycone of 2'*R*-SQGro. Only Arg438 interacts with the C2-hydroxyl group of the glyceryl 198 aglycone and thus this residue appears to drive selectivity for the 2'*R*-SQGro stereoisomer.

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Atu3277 (smoA) encodes a flavin mononucleotide (FMN) reductase

201 SmoA, annotated as a flavin reductase, was recombinantly expressed in E. coli and maintained a 202 yellow color throughout purification, suggesting that it had co-purified with a flavin co-factor. A 203 sample of this protein was heat-denatured to release the co-factor and the supernatant analyzed by 204 LC-MS to reveal that FMN was the sole detectable flavin (Fig. S8). Michaelis-Menten kinetics 205 were conducted for SmoA with saturating FMN and NADH or NADPH to determine which of these reductants was preferred by the enzyme. With NADH the kinetic parameters were $K_{\rm M} = 35\pm5~\mu{\rm M}$, 206 $k_{\rm cat} = 14.5 \pm 0.5 \text{ s}^{-1}$ and $k_{\rm cat}/K_{\rm M} = 4.1 \times 10^5 \text{ M}^{-1} \text{ s}^{-1}$; while for NADPH saturation was not observed and 207 $k_{\text{cat}}/K_{\text{M}} = 6.8 \times 10^2 \text{ M}^{-1} \text{ s}^{-1}$, indicating that NADH is the preferred cofactor for SmoA (**Fig. 3a, Fig.** 208 209 S9, Fig. S10). Owing to difficulties in obtaining structural data for this enzyme, we also studied a 210 close homologue from Rhizobium oryzae (RoSmoA, UniProt accession number: A0A1X7D6Q3), 211 which possesses a syntenic gene cluster to Atu3277-Atu3285. Recombinant RoSmoA also copurified with FMN (Fig. S8) and utilized the NADH cofactor with $K_{\rm M} = 16\pm5~\mu{\rm M}, k_{\rm cat} = 33\pm2~{\rm s}^{-1}$ 212 and $k_{\text{cat}}/K_{\text{M}} = 2.1 \times 10^6 \,\text{M}^{-1} \,\text{s}^{-1}$ (**Fig. S9**). 213

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Atu3279 (smoC) encodes an SQ monooxygenase that desulfurizes SQ

216 SmoC is annotated as an alkanesulfonate monooxygenase, though it possesses only 30% sequence 217 identity with the well-characterized alkanesulfonate monooxygenase SsuD, from E. coli. SsuD catalyzes the FMNH₂- and O₂-dependent oxidation of alkanesulfonates to produce the 218 219 corresponding aldehyde and sulfite, with a preference for pentanesulfonate (16). The mechanism of 220 this and related enzymes have been intensively studied yet remain enigmatic. The transformation is 221 thought to involve initial formation of a C4a-peroxy or N5-peroxy flavin species on-enzyme. One 222 mechanism posits that the terminal peroxide oxygen attacks the sulfonate sulfur of the substrate 223 before undergoing a rearrangement to effect C-S bond fissure and release of the aldehyde and 224 sulfite products (Fig. S12a) (17). An alternative mechanism suggests the peroxide deprotonates C6, 225 which is then oxidized to an α -hydroxysulfonate that undergoes elimination to produce sulfite and 226 the aldehyde (Fig. 12b) (18). To demonstrate activity for recombinant SmoC (Fig. S4), we adapted assays developed for SsuD that use Ellman's reagent to detect sulfite released by the enzyme (19). 227 228 Direct detection of the putative sugar product, 6-oxo-glucose (6-OG), is not trivial as this molecule 229 exists as a complex equilibrium of (hemi)acetals and hydrates that have poor stability. Thus, SmoC

was incubated with SQ in the presence of SmoA, FMN and NADH, which generate FMNH₂ in situ, and the concentration of sulfite determined periodically using Ellman's reagent (**Fig. 3b**). Maximal substrate conversion was approximately 200 μ M (**Fig. S12c**), which is commensurate with the solubility of molecular oxygen in water under standard conditions, with peak activity observed at pH 8.5 (**Fig. S12d**). No activity was observed when SQ was replaced with other sulfonates, including SQGro (the precursor to SQ) or HEPES (an unrelated sulfonate) demonstrating that, unlike the promiscuous SsuD, SmoC has high specificity for SQ (**Fig. S12c**). As such, the hydrolysis of SQGro by SmoI necessarily precedes oxidative desulfurization by SmoC. This observation is further supported by ITC, where SQ was found to bind SmoC with $K_d = 3 \mu$ M in the absence of any flavin-based cofactors, whereas no binding was detected for SQGro (**Fig. 3c, Fig. S13, Table S3**). The unique SQ monooxygenase activity of SmoC defines this pathway: it is the enzyme that effects fissure of the C–S bond in SQ, and so it was chosen as the namesake for this gene cluster and Atu3277-Atu3285 were renamed the <u>SQ MonoOxygenase cluster</u> (smoA-I).

While we could readily crystallize SmoC, these crystals only diffracted to a maximum resolution of 3.4 Å. The corresponding low-resolution map suggested that SmoC exists as a dimer, which was confirmed in solution by SEC-MALS (Fig. S14). To obtain structural information for an SQ monooxygenase, we turned to the homolog from R. oryzae (RoSmoC). Recombinant RoSmoC exhibited similar activity and substrate selectivity for SQ to SmoC (Fig. S12e) and provided crystals that diffracted to 1.9 Å. Importantly, the low-resolution structure of A. tumefaciens SmoC superimposed with the high-resolution RoSmoC structures with a peptide backbone rmsd of 0.4 Å across the entire structure, providing confidence that both enzymes shared a common structure and function (Fig. S15). Both SQ monooxygenases consist of a core $(\alpha/\beta)_8$ TIM barrel with three additional insertion regions, analogous to monooxygenases from the bacterial luciferase family. The protomers exist as a homodimer that buries 4697 Å² of surface area, amounting to 18% of total accessible surface area for each protomer (Fig. 3d). Pairwise structural analysis using the DALI server identified close relationships to a putative luciferase-like monooxygenase (3RAO.pdb) with an rmsd of 2.4 over 314 residues and a Z score of 34.3, the FMNH₂-dependent methanesulfonate monooxygenase MsuD (7K14.pdb, rmsd 2.0/322 residues, Z-score of 41.0), and the FMNH₂dependent alkanesulfonate monooxygenase SsuD (1M41.pdb, rmsd 1.8/317 residues, Z-score of 41.2).

Comparisons of the *Ro*SmoC structure with MsuD (7K14.pdb) in complex with FMN enabled identification of the FMN binding for site *Ro*SmoC: a deep hydrophobic pocket that accommodates the isoalloxazine ring system and extends out to the protein-solvent interface, which is gated by

conserved phosphate-binding residues Tyr136 and Ser189 (**Fig. 3e**) (18). The close structural and functional relationship of *Ro*SmoC to MsuD is evident from the conservation of a putative sulfonate binding site comprised of the side-chains Trp206, Arg236, His238, Tyr341 and His343 (18). Aside from conferring these enzymes with an ability to bind sulfonates, these conserved active-site residues have been suggested to contribute to the stabilization of a peroxyflavin intermediate in MsuD and SsuD (18, 19). Efforts to obtain crystals of a *Ro*SmoC–SQ complex were unsuccessful, limiting further insights into the origin of enzyme specificity towards SQ over other sulfonates.

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Atu3278 (smoB) encodes an oxidoreductase that converts 6-oxo-glucose to glucose

SmoB is annotated as a short-chain dehydrogenase/reductase (SDR) and we had hypothesized that it was responsible for reduction of 6-OG to glucose (Fig. 1d). Since 6-OG is difficult to study directly, we tested our hypothesis by looking for SmoB-mediated isotope incorporation into glucose at equilibrium (Fig. 4a). Assuming our hypothesis to be true, and as a consequence of microscopic reversibility, incubation of SmoB with a nicotinamide co-factor and glucose in H₂¹⁸O should result in transient formation of 6-OG, rapid and reversible hydration/dehydration with H₂¹⁸O to competeout ¹⁶O at C6 for ¹⁸O, and reduction to give 6-¹⁸O-glucose. In parallel to this process, ¹⁸O incorporation will occur at C1 of glucose through a similar series of hydration/dehydration reactions. Before proceeding with these experiments, we used ITC to establish which nicotamide cofactor was suitable for SmoB: NADPH bound to SmoB with $K_d \sim 2 \mu M$, while no binding was observed for NADH (Fig. S16, Table S3). Thus, glucose pre-equilibrated in $H_2^{18}O$ was incubated with SmoB and NADP⁺ then analyzed by mass spectrometry to reveal the formation of a product 4 Da greater in mass than glucose, presumably due to the incorporation of two ¹⁸O atoms into glucose. The crude reaction mixture was subjected to peracetylation (Ac₂O/pyridine) then LC-MS analysis to confirm that the +4 Da product co-eluted with authentic D-glucose-pentaacetate (Fig. S17). To determine that the ¹⁸O label was being incorporated at C6 of glucose, we used electronimpact GC-MS, which required conversion of the reaction product to the acyclic pentapropionate aldonitrile (Fig. S18) (20). This approach provided diagnostic C1-C5 and C5-C6 fragment ions. The ¹⁸O-labelled product gave a C5-C6 fragment that was 2 mass units higher (m/z 173 versus 175), whereas the C1-C5 fragment was the same as unlabelled glucose reference (m/z 370), demonstrating that the ¹⁸O is incorporated at C6. Only enzymatic reactions conducted in the presence of NADP⁺ produced product labelled with ¹⁸O at C6: NAD⁺ failed to produce any product. supporting our observations by ITC and defining the cofactor specificity of SmoB.

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We determined the 3D structure of SmoB using X-ray diffraction methods. This initial structure revealed that SmoB exists as a compact trimer, however the C-terminal His₆-tag in this construct

occupied the putative active site of adjoining subunits, making co-crystallization with cofactors difficult (Fig. S19). To overcome this issue, SmoB was subcloned into a different vector and expressed with a cleavable N-terminal purification tag. This protein maintained the same catalytic activity and SEC-MALS confirmed it remained a trimer in solution (Fig. S20). This SmoB construct was co-crystallized with NADPH and a ternary product complex obtained by soaking crystals with D-glucose (Fig. 4b). These crystals diffracted to a resolution of 1.5 Å and the resulting model revealed that SmoB is an $(\alpha/\beta)_8$ TIM barrel fold with a C-terminal cofactor binding site. The overall fold has high structural conservation with members of the aldo-keto reductase (AKR) superfamily. SmoB binds NADPH with the 2'-phosphate oxygens hydrogen-bonded to Thr284, Arg289 and backbone amide of Asn285 and the adenine ring stacked between Arg289 and Phe241 at the C-terminus (Fig. 4c). NADPH binds in an extended anti-conformation and the nicotinamide ring is located at the base of the substrate binding pocket. Trp232 makes a π - π stacking interaction with the nicotinamide ring that positions the reactive center (C4) at a distance of 3 Å from C-6 of glucose, appropriate for hydride transfer (Fig. 4d). Within the SmoB•NADP⁺•glucose complex, glucose interacts with Arg152 (2.9 Å) and Lys120 (3 Å), as well as His151 (2.8 Å) and Tyr76 (2.7 Å) within the conserved catalytic tetrad His/Tyr/Lys/Asp that is common to the AKR superfamily (**Fig. 4e**) (21).

SMO pathways occur in the Alphaproteobacteria and Betaproteobacteria

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319 To ascertain how widespread this pathway for SQ utilization might be, a Multigene BLAST search 320 was conducted of the non-redundant protein set of the NCBI for gene clusters that contain 321 homologous SQases and SQ monoxygenases. This identified many putative *smo* gene clusters 322 across the Agrobacterium and Rhizobium genus within the Rhizobiales order and evidence of some 323 broader expansion into the Alphaproteobacteria and Betaproteobacteria classes (Fig. 5). Amongst 324 these putative *smo* gene clusters, some were syntenic while others were substantially rearranged 325 (non-syntenic) or modified to make use of other (non-ABC) transporter systems. The use of diverse 326 transport systems is not surprising: a similar phenomenon has been observed for the tier-1 sulfo-ED 327 pathway (5, 6). Indeed, sulfo-ED gene clusters have been identified in several Rhizobiales (5, 6), suggesting that there has been ample opportunity for genetic exchanges between these pathways 328 329 during their evolution.

Discussion

While existing pathways for the breakdown of SQ require two different organisms and involve scission of the carbon chain into two 3-carbon fragments, we describe here a fundamentally different approach that features complete utilization of the SQ carbon skeleton. The SMO pathway features several proteins with hitherto undescribed activities, including: an SQGro-binding protein; an FMNH₂- and O₂-dependent SQ monooxygenase that defines this 'SMO' pathway by catalyzing scission of the C-S bond in SQ; and an oxidoreductase dedicated to the NADPH-dependent reduction of 6-OG to glucose. Like all other sulfoglycolytic pathways studied to date, the SMO pathway also possesses a conserved SQase, which is essential for liberating SQ from its precursor glycoside SQGro (9, 10).

The SMO pathway is reminiscent of other sugar-metabolizing pathways in bacteria. For example, the SmoI (SQase), SmoF (SQGro binding protein) and SmoE/G/H (ABC transporter) proteins encoded by the *smo* cluster are analogous to MalP (maltodextrin phosphorylase), MalE (maltose binding protein) and MalF/G/K (ABC transporter) encoded by the *mal* operon of *E. coli* that imports and degrades maltose (22). Additionally, the SmoC (SQ monooxygenase) and SmoA (flavin reductase) proteins of the SMO pathway are reminiscent of the SsuD (FMNH₂-dependent alkylsulfonate monooxygenase) and SsuE (NADPH-dependent FMN reductase) pair encoded by the *ssu* operon of *E. coli* that degrades alkanesulfonates (16). Indeed, it is likely that the SMO pathway arose through the recombination and neofunctionalization of analogous sugar- and sulfonate-metabolising pathways.

Through structural analysis we identified key residues involved in sulfosugar recognition and processing, in order to provide greater confidence to bioinformatic analyses of putative *smo* gene clusters: an approach that has proven valuable for the identification of tier 1 sulfoglycolytic pathways (9, 23, 24). This includes the Thr220-Gly166-Ser43-H₂O(His13-Gln46) cluster of SmoF for the recognition of SQGro, the Arg283-Tryp286-H₂O(Tyr491) triad of SmoI for the recognition of SQGro; and the Trp206-Arg236-His238-Tyr341-His343 constellation of SmoC for the recognition of SQ. Given the importance of the SQ monooxygenase SmoC to the SMO pathway, further empirical and computational work is warranted to understand what interactions drive its selectivity for SQ, which lies in contrast with the promiscuity exhibited by alkanesulfonate monoxygenases like SsuD.

The prevalence of the SMO pathway in Alphaproteobacteria of the *Rhizobiales* order is intriguing, since many bacteria of this order are plant symbionts or pathogens. Indeed, those bacteria that do

not possess an SMO pathway often possess a complementary tier 1 sulfo-ED pathway (5). Accordingly, it appears that plant sulfolipid catabolism is important for rhizobiales, whether they be plant pathogens/symbionts or free-living organisms adopting an oligotrophic saprophytic lifestyle in substrate replete with decaying plant tissues. Symbiotic bacteria of the Rhizobiales order reside within the root nodules of their plant host, where they harness four-carbon substrates from the host for energy and central metabolism (25). Sugawara and co-workers showed that sulfonate utilization gene clusters were expressed by the plant symbiont Bradyrhizobium diazoefficiens USDA 110 within these nodules and that this may be important for utilizing diverse sulfur sources to support symbiotic and possibly free-living lifestyles (26). With sulfolipid representing a large and accessible pool of sulfur in plants, one possible purpose of the SMO pathway may be to salvage sulfur for these bacteria. This is an important distinction between the SMO pathway and the tier 1 sulfoglycolytic pathways: the latter supports two-member microbial communities containing a second member with a tier 2 pathway to provide access to the sulfur of SO (27). In this sense, use of the SMO pathway, which enables the complete utilization of the carbon skeleton and access to the sulfur of the monosaccharide can be considered a 'selfish' metabolic strategy, and could provide an advantage in the highly competitive soil environment or in the absence of other bacterial species within colonized plant tissues. Combined with the pathway's requirement for molecular oxygen to effect C-S bond fissure, this may explain why the SMO pathway occurs within those bacteria that are commonly associated with plants. Understanding how the SMO and tier 1 pathways impact fitness within different environmental niches remains an important question, with answers that have significant implications for understanding plant diseases and symbioses, as well as soil chemistry.

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387 **Methods**

- 388 Specialist reagents
- 389 SQ and methyl α-sulfoquinovoside were purchased from MCAT GmbH (Donaueschingen,
- 390 Germany), (13C₆)SQ, glycer-1-yl α-sulfoquinovoside (SQGro), and dicyclohexylammonium
- 391 sulfolactate, cyclohexylammonium dihydroxypropanesulfonate were synthesized as described (13,
- 392 28). All other sulfonates were purchased from Sigma-Aldrich.

393

- 394 *Growth studies*
- 395 Cultures of A. tumefaciens C58 were grown in a phosphate-buffered mineral salts media (M9, pH
- 396 7.2), with glucose or SQ (10 mM) as the sole carbon source. Cultures were incubated at 30 °C (250
- 397 rpm), with adaptation and robust growth observed within 2–3 days. These were sub-cultured (1%
- 398 inoculum) into the same media (10 mL) and grown at 30 °C (250 rpm). Bacterial growth was
- 399 quantitated using a Varian Cary50 UV/visible spectrophotometer to measure OD₆₀₀. Growth
- 400 experiments were replicated twice.

401

- 402 Reducing sugar assay for culture supernatant
- The reducing sugar assay was performed according to the procedure of Blakeney and Mutton (29).
- 404 This assay uses pre-prepared alkaline diluent and 4-hydroxybenzoic acid hydrazide (PAHBAH)
- working solution. Alkaline diluent was prepared by the addition of sodium hydroxide (20 g, 0.50
- 406 mol) to a solution of 0.10 M trisodium citrate (50 mmol, 500 mL) and 0.02 M calcium chloride (13
- 407 mmol, 500 mL). PAHBAH working solution was prepared by dissolving 4-hydroxybenzhydrazide
- 408 (PAHBAH) (0.25 g, 1.6 mmol) in alkaline diluent (50 mL). The PAHBAH working solution should
- be made fresh shortly before use. To determine reducing sugar concentration, 0.90 mL of PAHBAH
- 410 working solution was added to 0.10 mL of sample. The mixture was heated at 98 °C for 4 min then
- 411 0.5 mL of the mixture was diluted into 1.0 mL of deionized water and the absorbance read at 415
- nm using a Varian Cary50 UV/visible spectrophotometer. Concentrations of SQ were determined
- with reference to a standard curve constructed using SQ.

- 415 Turbidometric sulfate assay for culture supernatant
- The sulfate assay was performed according to the procedure of Sörbo (30). This assay uses a Ba-
- PEG reagent, which contains PEG to stabilize BaSO₄ crystals and a small amount of pre-formed
- BaSO₄ seed crystals to improve the reproducibility and linearity of the assay. The Ba-PEG reagent
- should be prepared fresh before use. Ba-PEG reagent was prepared by dissolving BaCl₂ (42 mg,
- 420 0.20 mmol) and polyethylene glycol 6000 (0.75 g) in deionized water (5.0 mL). A small amount of
- Na₂SO₄ (10 μL, 50 mM) was added to this solution, with efficient magnetic stirring to generate

422 preformed BaSO₄ seed crystals. Individual sulfate assays were conducted as follows. An aliquot of

423 culture supernatant obtained after pelleting of cells for 5 min at 5000 g (typically 100 μL,

- 424 containing a maximum of 2.5 μmol of Na₂SO₄) was diluted to 0.1 mL with deionized water before
- 425 the addition of 0.5 M HCl (0.1 mL) followed by Ba-PEG reagent (0.1 mL). The mixture was mixed
- 426 vigorously and the absorbance of the sample at 400 nm determined using a Varian Cary50
- 427 UV/visible spectrophotometer. Concentrations of sulfate were determined by reference to a
- standard curve constructed using Na₂SO₄. This curve was linear up to 2.5 µmol of Na₂SO₄.

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- 430 Colorimetric fuchsin sulfite assay for culture supernatant
- The fuchsin sulfite assay was performed according to the procedures of Brychkova et al. (31) and
- Kurmanbayeva et al. (32). This procedure requires three pre-prepared solutions, Reagents A, B and
- 433 C. Reagent A was prepared by dissolution of basic fuchsin (4.0 mg, 12 μmol) in deionized water
- 434 (8.25 mL) at 0 °C, prior to the addition of 98% H₂SO₄ (1.25 mL). Reagent B was prepared by
- diluting formaldehyde (36% in H₂O, 0.32 mL) in deionized water (9.68 mL) at 0°C. Reagent C was
- prepared by dilution of Reagent A (1 mL) in deionized water (7 mL), prior to the addition of
- solution reagent B (1 mL). Individual sulfite assays were performed by addition of Reagent C (516
- 438 μL) to a mixture of sample (72 μL) and 0.5 mM Na₂SO₃ (12 μL), with the latter providing a stable
- background signal for reference. The sample was incubated at 20-22°C for 10 min and the
- 440 absorbance of the sample at 570 nm determined using a Varian Cary50 UV/visible
- spectrophotometer. Concentrations of sulfite were determined by reference to a standard curve
- constructed using Na₂SO₃.

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- NMR analysis of metabolites produced from ($^{13}C_6$)SQ
- M9 minimal media (5 mL) containing 10 mM glucose was inoculated with A. tumefaciens C58 and
- grown to stationary phase at 30 °C (250 rpm). A 50 µL aliquot of this culture was used to inoculate
- 2 mL of M9 minimal media containing 10 mM (¹³C₆)SQ and the culture incubated at 30 °C (250
- rpm). At OD₆₀₀ 0.27 and OD₆₀₀ 0.49, 950 μL samples of culture supernatant were diluted with 100
- 449 µL of D₂O and ¹³C-NMR spectra acquired using a 400 MHz spectrophotomer (100 MHz for ¹³C).

- 451 Growth of A. tumefaciens C58 on diverse alkanesulfonates
- M9 minimal media (5 mL) containing 10 mM glucose was inoculated with A. tumefaciens C58 and
- grown to stationary phase at 30 °C (250 rpm). A 50 µL aliquot of this starter culture was used to
- 454 inoculate 2 mL of M9 minimal media containing 10 mM of the alternative alkanesulfonate
- 455 substrate: SQ (positive control), methyl α-sulfoquinovoside (MeSQ), glycer-1-yl α-
- 456 sulfoquinovoside (SQGro), dicyclohexylammonium sulfolactate, cyclohexylammonium

457 dihydroxypropanesulfonate, sulfoacetic acid, taurine, sodium pentanesulfonate, cysteic acid, 458 MOPS, HEPES, PIPES, MES and methanesulfonic acid. Cultures were incubated for 30 days at 30 459 °C (250 rpm) with daily observations of optical density at 600 nm. Each experiment was performed

460 in duplicate. Growth was observed on SQ (positive control), MeSQ, and SQGro, but not on any 461

other sulfonate. Control experiments established that A. tumefaciens grows on glucose in the

presence and absence of cyclohexylamine or dicyclohexylamine, and does not grow on

cyclohexylamine or dicyclohexylamine alone.

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- 465 Digestion of samples for quantitative proteomics
- 466 Freeze dried A. tumefaciens whole-cell pellets were resuspend in 500 µL lysis buffer (4% SDS, 50
- 467 mM Tris pH 8.5, 10 mM DTT) and boiled at 95 °C for 10 min with shaking at 2000 rpm to shear
- 468 DNA and inactivate protease activity. Lysates were cooled to room temperature and protein
- 469 concentration determined using a BCA assay. Each sample (200 µg of protein) was acetone
- 470 precipitated by mixing 4 volumes of ice-cold acetone with one volume of sample. Samples were
- 471 precipitated overnight at -20 °C and then centrifuged at 4000 × g for 10 min at 4 °C. The
- 472 precipitated protein pellets were resuspended with 80% ice-cold acetone and precipitated for an
- 473 additional 4 h at -20 °C. Samples were centrifuged at 17000 × g for 10 min at 4 °C to collect
- precipitated protein, the supernatant was discarded and excess acetone driven off at 65 °C for 5 min. 474
- 475 Dried protein pellets were resuspended in 6 M urea, 2 M thiourea, 40 mM NH₄HCO₃ and
- reduced/alkylated prior to digestion with Lys-C (1/200 w/w) then trypsin (1/50 w/w) overnight as 476
- 477 previously described (33). Digested samples were acidified to a final concentration of 0.5% formic
- 478 acid and desalted using C18 stage tips (34) before analysis by LC-MS.

- 480 Quantitative proteomics using reversed phase LC-MS
- 481 Purified peptides were resuspended in Buffer A* (2% MeCN, 0.1% TFA) and separated using a
- 482 Proflow-equipped Dionex Ultimate 3000 Ultra-Performance Liquid Chromatography system
- 483 (Thermo Fisher Scientific) with a two-column chromatography set up composed of a PepMap100
- 484 C18 20 mm × 75 µm trap and a PepMap C18 500 mm × 75 µm analytical column (Thermo Fisher
- Scientific). Samples were concentrated onto the trap column at 5 µL min⁻¹ with Buffer A (2%) 485
- MeCN, 0.1% FA) for 6 min and then infused into an Orbitrap Q-Exactive HF Mass Spectrometer 486
- (Thermo Fisher Scientific) at 250 nl min⁻¹. Peptides were separated using 124-min gradients 487
- 488 altering the buffer composition from 2% Buffer B (80% MeCN, 0.1% FA) to 8% B over 14 min,
- 489 then from 8% B to 30% B over 80 min, 30% B to 45% B over 10 min, 45% B to 95% B over 2 min,
- 490 holding at 95% B for 10, then dropped to 2% B over 1 min and holding at 2% B for the remaining 7
- 491 min. The Q-Exactive HFTM Mass Spectrometer was operated in a data-dependent mode

492 automatically switching between the acquisition of a single Orbitrap MS scan (120,000 resolution) 493

and a maximum of 20 MS-MS scans (HCD NCE 28, maximum fill time 40 ms, AGC 2×10⁵ with a

resolution of 15,000).

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- Mass spectrometry data analysis
- 497 Proteomics datasets were searched using MaxQuant (v1.5.3.3) (35) against the A. tumefaciens C58
- 498 proteome (Uniprot proteome id UP000000813, downloaded 27/01/2018, 5344 entries). Searches
- 499 were performed with carbamidomethylation of cysteine set as a fixed modification and oxidation of
- 500 methionine as well as acetylation of protein N-termini allowed as variable modifications. The
- 501 protease specificity was set to trypsin allowing 2 miscleavage events with a maximum false
- 502 discovery rate (FDR) of 1.0% set for protein and peptide identifications. To enhance the
- 503 identification of peptides between samples the Match Between Runs option was enabled with a
- 504 precursor match window set to 2 min and an alignment window of 10 min. For label-free
- 505 quantitation, the MaxLFQ option within Maxquant(36) was enabled in addition to the re-
- 506 quantification module. The resulting protein group output was processed within the Perseus
- 507 (v1.4.0.6) (37) analysis environment to remove reverse matches and common protein contaminates
- 508 prior. For LFQ comparisons missing values were imputed using Perseus and Pearson correlations
- 509 visualized using R. The mass spectrometry proteomics data have been deposited to the
- 510 ProteomeXchange Consortium via the PRIDE (38) partner repository with the dataset identifier
- 511 PXD014115.

512

- 513 Cloning
- 514 Oligonucleotides encoding Atu3277 (SmoA), Atu3278 (SmoB), Atu3279 (SmoC) and Atu3282
- 515 (SmoF) were amplified by PCR using Phusion polymerase HF master mix (NEB), the appropriate
- 516 primers listed in Table S1 and A. tumefaciens C58 gDNA as template. Oligonucleotides encoding
- 517 RoSmoA and RoSmoC were synthesized (IDT) to provide the sequences listed in **Table S1**. These
- 518 were cloned into the pET29b(+) vector at the NdeI and XhoI sites and sequence-verified by Sanger
- 519 sequencing to give expression vectors for SmoA, SmoB, SmoC, SmoF, RoSmoA and RoSmoC.
- 520 Due to interference from the SmoB C-terminal His6-tag during structural studies, the smoB
- (Atu3278) gene was sub-cloned into the pET-YSBLIC3C vector (39) by PCR amplification with the 521
- relevant primers in Table S1 and In-Fusion® cloning (Clontech Laboratories, Inc.) into linearized 522
- 523 YSBLIC3C vector according to the manufacturer's protocol. The expression plasmid was sequence-
- verified by Sanger sequencing. 524

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Protein expression and purification

527 All vectors were transformed into 'T7 Express' E. coli (NEB), except for the vector encoding SmoF 528 (Atu3282), which was transformed into 'Shuffle® T7' E. coli (NEB), and all were plated onto LBagar (50 µg mL⁻¹ kanamycin) and incubated at 37 °C for 16 h. A single colony was used to 529 inoculate 10 mL of LB media containing 50 µg mL⁻¹ kanamycin and the cultures incubated at 37 °C 530 531 for 16 h. These starter cultures were used to inoculate 1000 mL of S-broth (35 g tryptone, 20 g yeast extract, 5 g NaCl, pH 7.4) containing 50 µg mL⁻¹ kanamycin, which was incubated with shaking 532 (250 rpm) at 37 °C until it reached an OD₆₀₀ of 0.8. Each culture was cooled to room temperature, 533 534 isopropyl thiogalactoside (IPTG) added to a final concentration of 400 µM, and incubation with 535 shaking (200 rpm) continued at 18 °C for 19 h. Cells were harvested by centrifugation at 8,000 g for 20 min at 4 °C then resuspended in 40 mL binding buffer (50 mM NaP_i, 300 mM NaCl, 5 mM 536 537 imidazole, pH 7.5) containing protease inhibitor (Roche cOmplete EDTA-free protease inhibitor cocktail) and lysozyme (0.1 mg mL⁻¹) by nutating at 4 °C for 30 min. Benzonase (1 µL, 250 U) was 538 539 added to the mixture then lysis was effected by sonication [10× (15 s on / 45 s off) at 45% 540 amplitude]. The lysate was centrifuged at 18,000 g for 20 min at 4 °C and the supernatant collected. 541 The supernatants were filtered (0.45 µm) and loaded onto a 1 mL HiTrap TALON IMAC column 542 (GE). The column was washed with 3×10 mL of binding buffer, then the protein was eluted using 543 elution buffer (50 mM NaP_i, 300 mM NaCl, 400 mM imidazole, pH 7.5). Fractions containing product, as judged by SDS-PAGE, were further purified by size exclusion chromatography on a 544 545 HiPrep 16/60 Sephacryl S-200 HR column (GE) using 50 mM NaP_i, 150 mM NaCl, pH 7.5 (Atu3277 SmoA; Atu3278, SmoB; Atu3279, SmoC) or 50 mM sodium citrate, 150 mM NaCl, pH 546 547 5.5 (Atu3282, SmoF) as buffer (Fig. S2). SmoI (Atu3285 or AtSQase) was prepared as previously described (9). 548

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SEC-MALS analyses

551 Experiments were conducted on a system comprising a Wyatt HELEOS-II multi-angle light 552 scattering detector and a Wyatt rEX refractive index detector linked to a Shimadzu LC system 553 (SPD-20A UV detector, LC20-AD isocratic pump system, DGU-20A3 degasser and SIL-20A 554 autosampler). Experiments were conducted at room temperature (20 ± 2 °C). Solvents were filtered through a 0.2 µm filter prior to use and a 0.1 µm filter was present in the flow path. The column 555 was equilibrated with > 2 CV of buffer (50 mM NaPi, 300 mM NaCl pH 7.4) before use and buffer 556 was infused at the working flow rate until baselines for UV, light scattering and refractive index 557 detectors were all stable. The sample injection volume was 100 µL of protein at 6 mg mL⁻¹ in 50 558 mM NaPi buffer, 300 mM NaCl pH 7.4. Shimadzu LC Solutions software was used to control the 559 560 LC and Astra V software for the HELEOS-II and rEX detectors. The Astra data collection was 1 561 min shorter than the LC solutions run to maintain synchronization. Blank buffer injections were

- used as appropriate to check for carry-over between sample runs. Data were analyzed using the
- Astra V software. Molecular weights were estimated using the Zimm fit method with degree 1. A
- value of 0.158 was used for protein refractive index increment (dn/dc).

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- 566 Isothermal Titration Calorimetry
- 567 ITC experiments were performed using a MicroCal iTC200 (GE Healthcare) at 25 °C, with a 750
- r.p.m. stirring speed and a reference power of 10 μCal.s⁻¹. Proteins and substrates were equilibrated
- into degassed and filter-sterilized buffer (50 mM NaPi, 200 mM NaCl, pH 7.4 for SmoC/F and 25
- 570 mM NaPi, pH 7.5 for Smo B). Protein concentration was determined by BCA assay (Thermo
- 571 Fisher) before intiating experiments. For SmoC–SQ binding, 600 μM SQ was titrated into the ITC
- 572 cell containing 40 μ M SmoC as a series of 10 \times 3.94 μ L injections with a pre-injection of 1 \times 0.4
- 573 μL. For SmoF–SQGro binding, 200 μM SQGro was titrated into the ITC cell containing 20 μM
- SmoF as a series of $15 \times 2.94 \mu L$ injections with a pre-injection of $1 \times 0.4 \mu L$. The delay between
- 575 injections was set at 120 s, with an initial injection delay of 60 s. For SmoB-NAD(P)H binding, 1
- 576 mM NADH was titrated into the ITC cell containing 40 μ M SmoB as a series of 19 \times 3 μ L
- injections with a pre-injection of $1 \times 4 \mu L$. The delay between injections was set at 150 s, with an
- 578 initial injection delay of 180 s. All data analysis was performed in MicroCal ITC Origin Analysis
- 579 software (Malvern).

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- Nano Differential Scanning Fluorescence analysis of SmoF
- Thermal stability analysis for SmoF in the presence and absence of SQGro ligand was performed on
- a Prometheus NT.48 (NanoTemper) at 15% excitation, scanning from 20 °C to 65 °C at 0.5 °C min
- ¹. All protein samples were at a concentration of 1 mg mL⁻¹ in 50 mM citrate, 150 mM NaCl at pH
- 585 5.5, with a 10 µL capillary load per sample. Data acquisition and analysis was performed with
- 586 PR.ThermControl (NanoTemper) software.

- 588 Identification of the flavin co-factor that co-purified with SmoA
- 589 100 μL of recombinant flavin reductase (SmoA or *Ro*SmoA) at a concentration of 20 mg mL⁻¹ in 50
- 590 mM Tris, 150 mM NaCl, pH 8.5 was heated at 90 °C for 10 min. The sample was clarified by
- 591 centrifugation (16,000 ×g, 10 min, 4 °C) and the supernatant filtered (0.2 μm). Samples were
- analyzed by LCMS on an Agilent LCMS system (G6125B mass detector, 1290 Infinity G7120A
- 593 high speed pump, 1290 Infinity G7129B autosampler, and 1290 Infinity G7117B diode array
- detector). Conditions for LC were as follows: column: Phenomenex 00B-4752-AN Luna Omega 1.6
- μ m PS C₁₈ 100Å (50 × 2.1 mm); injection volume: 1 μL; gradient: 3 to 100% B over 20 min

596 (solvent A: water + 0.1% FA; solvent B: MeCN + 0.1% FA); flow rate: 0.6 mL min⁻¹; DAD – 254

597 and 214 nm.

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- 599 Michaelis-Menten kinetic analyses of SmoA and RoSmoA
- Reactions were conducted at 25 °C in 96-well plate format and involved the addition of SmoA or
- 601 RoSmoA (final concentration of 20 nM for NADH and 500 nM for NADPH) to 20-800 μM
- NAD(P)H in 50 mM NaPi, 150 mM NaCl, 30 µM FMN, 0.01% BSA, pH 7.4 at a total volume of
- 603 100 μL. The progress of the enzyme-catalyzed conversion of NAD(P)H to NAD(P)⁺ was monitored
- by measuring loss of absorbance at 340 nM over time using an Envision Multimodal Plate Reader
- 605 (Perkin Elmer). Initial rates for each reaction were calculated after first subtracting the rate of
- spontaneous NAD(P)H oxidation (determined using an enzyme-free control) and an empirically
- determined extinction coefficient for NAD(P)H under these conditions. Each initial rate was
- determined in triplicate and fit to a Michaelis-Menten equation using Prism 8 (GraphPad).

609

- 610 Sulfoquinovose monooxygenase assay
- This SQ monooxygenase activity assay is based on a previously described alkanesulfonate
- 612 monooxygenase activity assays (19) and uses Ellman's reagent to quantify sulfite released by these
- enzymes. A 2 mL reaction containing 1 mM SQ, 1 mM NADH, 3 µM FMN, 0.01% (w/v) BSA, 100
- nM SmoA or RoSmoA and 300 nM SQ monooxygenase (SmoC or RoSmoC) in buffer (25 mM Tris
- pH 9.1, 25 mM NaCl) was incubated at 30 °C, along with controls lacking reaction components or
- using alternate sulfonate substrates. Reactions were initiated by the addition of SmoA or RoSmoA
- 617 to the mixture. Sulfite concentration in the samples was determined at discrete time points by
- 618 quenching 40 μL of the reaction in 160 μL of Ellman's reagent (0.125 mg mL⁻¹ in 25 mM NaPi pH
- 7.0, prepared fresh) within a 96-well plate. After 60 s, the absorbance of the sample at 405 nm was
- determined using an Envision Multimodal Plate Reader (Perkin Elmer). The sulfite concentration
- was interpolated using a calibration curve generated under these conditions: a linear relationship
- between sulfite concentration and absorbance at 405 nm was observed for 5–1000 μM Na₂SO₃. The
- activity of SQ monooxygenases at different pH was determined by modifying the buffer in the
- above reactions (MES: pH 6.0, 6.5 and Tris: pH 7.0, 7.5, 8.0, 8.5, 9.1) using an endpoint of t = 30
- 625 min.

- 627 Equilibrium isotope labelling using SmoB
- In order to pre-label the anomeric position, glucose was incubated in 98% H₂¹⁸O with heating at 80
- °C for 2 days, then evaporated to dryness to give C1-18O-labelled glucose. Labelling was determined
- to be 95% by mass spectrometry based on intensities of the M and M+2 peaks. Using ${\rm H_2}^{18}{\rm O}$ buffer

(100 mM potassium phosphate, pH 7.0), NAD+ and NADP+ were each added at 0.05 molar 631 632 equivalent to C1-18O-glucose and SmoB. Four control experiments were conducted: one without enzyme, one without NAD⁺ and NADP⁺, one in H₂¹⁶O, and one in H₂¹⁶O with unlabeled glucose. 633 Reactions were monitored by mass spectrometry. Only in the experimental sample containing 634 enzyme, H₂¹⁸O and NAD⁺/NADP⁺ was an M+4 signal observed and this reached a maximum 635 intensity after 72 h. Two additional reactions were performed using SmoB, glucose and either 636 NADP⁺ or NAD⁺ in H₂¹⁸O and only the reaction containing NADP⁺ generated the M+4 species. To 637 confirm that the M+4 species was glucose with two ¹⁸O labels, we studied the product by HPLC. 638 However, under aqueous HPLC conditions the ¹⁸O-label at C1 is lost through chemical exchange 639 with solvent. Therefore, we acetylated the product to form the pentaacetate to ensure no exchange at 640 641 the anomeric position during HPLC analysis. The reaction mixture from above was evaporated 642 under reduced pressure. The crude residue was treated with acetic anhydride in pyridine (1:2, 1 mL) 643 overnight. The product was extracted with EtOAc and washed with sat. CuSO₄ to remove pyridine. The organic solution containing peracetylated glucose was analyzed by LCMS on an Agilent LCMS 644 system (G6125B mass detector, 1290 Infinity G7120A high speed pump, 1290 Infinity G7129B 645 autosampler, and 1290 Infinity G7117B diode array detector). Conditions for LC were as follows: 646 647 column: Phenomenex 00B-4752-AN Luna Omega 1.6 μm PS C₁₈ 100Å (50 × 2.1 mm); injection volume: 1 μL; gradient: 0 to 65% B over 20 min (solvent A: water + 0.1% FA; solvent B: MeCN + 648 0.1% FA); flow rate: 0.6 mL min⁻¹. Peaks with m/z 413 [M+Na]⁺, m/z 415 [M+2+Na]⁺, and m/z 417 649 [M+4+Na]⁺ had the same retention time as an authentic glucose pentaacetate standard. 650

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652 GC-MS analysis of isotopically-labelled carbohydrates

653 A 0.1 μ L aliquot of SmoB-glucose reaction mixture (containing \approx 2.5 nmol glucose) was transferred to a GC vial insert (deactivated) together with 1 nmol scyllo-inositol as an internal standard. 654 655 Samples were derivatized as described in Antoniewicz et al. (20), with minor modifications. Briefly, samples were dried (in vacuo, 35 °C with a 40 µL methanol wash), followed by addition of 656 hydroxylamine hydrochloride (20 mg mL⁻¹ in 25 µL pyridine) and incubation at 90 °C for 1 h. Vials 657 were cooled briefly at 20-22°C before the addition of propionic anhydride (50 µL) and incubation at 658 659 60 °C for 30 min. Samples were evaporated to dryness under a stream of nitrogen at 60 °C and resuspended in EtOAc (40 µL). Control samples of U-12C-glucose, U-13C-glucose, 1,2-13C₂-glucose 660 and 6.6-2H₂-glucose were also prepared at a 2.5 nmol scale in the assay buffer mixture. Samples 661 662 were blinded for analysis. The derivatized labelled glucose samples (Fig. S13 and Table S7) were analyzed by GC-MS using a DB5 capillary column (J&W Scientific, 30 m, 250 µm inner diameter, 663 664 0.25 µm film thickness) with a 10 m inert duraguard. The injector insert and GC-MS transfer line temperatures were 270 °C and 250 °C, respectively. The oven temperature gradient was 665

- of 666 programmed as follows: 70 °C (1 min); 70 °C to 295 °C at 12.5 °C min⁻¹; 295 °C to 320 °C at 25 °C
- 667 min⁻¹; 320 °C for 2 min. Glucose and *scyllo*-inositol were identified by reference to authentic
- standards. A calibration curve was generated using glucose standard in assay buffer (starting
- concentration 50 nmol, 2-fold dilution series). Fig. S12 shows the fraction of labelled fragments,
- 670 corrected for isotope natural abundance by DExSI analysis (40).
- 671
- 672 Protein crystallization
- 673 Initial crystallization screening was performed using commercially available INDEX (Hampton
- Research), PACT premier and CSSI/II (Molecular Dimensions) screens in 96-well sitting drop
- 675 trays. Further optimization was carried out in a 48-well sitting drop or 24-well hanging-drop format
- 676 to obtain optimal crystals for X-ray diffraction. Unless otherwise stated, all crystals were grown at
- 677 20 °C.
- 678
- 679 Crystals of apo-SmoF were obtained by mixing 0.15 μL of protein stock (50 mg mL⁻¹ protein in 50
- 680 mM citrate, 150 mM NaCl, pH 5.5) with 0.15 μL mother liquor (0.3 M ammonium acetate, 0.1 M
- Bis-Tris, 25% w/v PEG 3350, pH 5.5) housed in a Rigaku Xtaltrak plate hotel to enable consistent
- growth and monitoring at 6 °C. Crystals were harvested with nylon CryoLoopsTM (Hampton
- Research) and cryopreserved in liquid nitrogen without additional cryoprotectants.
- 684
- Crystals of SmoF were initially obtained by mixing 0.15 µL of protein stock (3.5 mg mL⁻¹ protein
- with 2'R-SQGro at a 1:10 molar ratio in 50 mM citrate, 150 mM NaCl, pH 5.5) with 0.15 μL
- mother liquor (30% (w/v) polyethylene glycol 4000, 0.2 M sodium acetate, 0.1 M tris chloride, pH
- 8.5). The resulting crystals were used to prepare a seed stock by mixing the crystallization drop
- with 100 µL mother liquor and vortexing for 60 s with one teflon bead. An optimisation plate was
- setup with drops comprised of 0.1 µl of various mother liquors (28-36% (w/v) polyethylene glycol
- 691 4000, 0.2 M sodium acetate, 0.1 M tris chloride, pH 7.1-9.1), 50 nl seed stock solution, and 0.15 μL
- protein stock (4 mg mL⁻¹ protein with 2'R-SQGro at a 1:10 molar ratio in 50 mM citrate, 150 mM
- NaCl, pH 5.5). A single crystal grown at 31.8% (w/v) polyethylene glycol 4000, 0.2 M sodium
- acetate, 0.1 M tris chloride, pH 8.95, was harvested with a nylon CryoLoopTM (Hampton Research)
- and cryopreserved in liquid nitrogen with 25% (v/v) ethylene glycol as cryoprotectant.
- 696
- 697 Crystals of SmoI-D455N-E370A-E371A were obtained by mixing 0.4 µL of protein stock (35 mg
- 698 mL⁻¹ protein in 50 mM NaPi, 300 mM NaCl, pH 7.4) with 0.5 μL mother liquor (26% PEG 3350
- 699 w/v, 0.2 M KSCN, 0.1 M Bis-Tris propane, pH 6.5). Crystals were soaked with solid SQGro in

mother liquor for 2 min prior to harvesting with nylon CryoLoopsTM (Hampton Research) and cryopreserved without additional cryoprotectants.

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Crystals of apo-SmoC were obtained by mixing 0.6 μL of protein stock (60 mg mL⁻¹ protein in 50 mM Tris, 300 mM NaCl, pH 7.5) with 0.5 μL mother liquor (0.2 M NaCl, 0.1 M MES pH 6, 26% PEG 6000 w/v and 10 mM SQ-glucitol). Crystals of apo-*Ro*SmoC were obtained by mixing 0.1 μL of protein stock (11.7 mg mL⁻¹ protein in 50 mM Tris, 300 mM NaCl, pH 7.5) with 0.2 μL mother liquor (0.2M NaNO₃, 20% PEG 3350 w/v and 10 mM SQ). Crystals were harvested with nylon CryoLoopsTM (Hampton Research) and cryopreserved in liquid nitrogen without additional

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cryoprotectants.

- 711 Crystals of SmoB-apo (YSBLIC3C construct) were obtained by mixing 0.15 μ L of protein stock 712 (20 mg mL⁻¹ protein in 50 mM NaPi, 150 mM NaCl, pH 7.4) with 0.15 μ L mother liquor (0.2 M
- sodium malonate dibasic monohydrate, 0.1 M Bis-Tris propane pH 8.5, 20% w/v PEG 3350). For
- 714 the SmoB•NADPH complex, crystals were obtained by mixing 0.15 μL of protein stock (20 mg
- 715 mL^{-1} protein in 50 mM NaPi, 150 mM NaCl, 2 mM NADPH, pH 7.4) with 0.15 μ L mother liquor
- 716 (0.1 M succinic acid, sodium dihydrogen phosphate, glycine buffer (SPG buffer, Qiagen), 25% w/v
- 717 PEG 1500 at pH 6.0). For the SmoB•NADPH•Glc complex, crystals were obtained in a hanging
- drop by mixing 1 μL of protein stock (13 mg mL⁻¹ protein in 50 mM NaPi 150 mM NaCl, pH 7.4)
- vith 1 μL of mother liquor (2 mM NADPH, 0.1 M SPG (Qiagen), 25% w/v PEG 1500 at pH 6).
- 720 Crystals were soaked with solid glucose in mother liquor for 1 min prior to harvesting with nylon
- 721 CryoLoopsTM (Hampton Research) and cryopreserved without additional cryoprotectants.

- 723 *X-ray data collection, processing and refinement*
- The data were processed and integrated using XDS (41) and scaled using SCALA (42) included in
- the Xia2 processing system (43). Data reduction was performed with AIMLESS, and resolution was
- cut until CC1/2 = 0.5. The structure of the SmoI•SQGro complex was determined using molecular
- 727 replacement using 50HS (9) as the initial model. For SmoF, the structure was solved by molecular
- 728 replacement using PHASER (44) with a search model created from PDB ID: 6DTQ (45). The
- structure of RoSmoC was solved by molecular replacement using the ensemble based on PDB ID:
- 730 1M41 (19) as an initial search model. The structure of SmoB was determined using molecular
- 731 replacement with the monomer of an aldo-keto reductase from S. enterica (PDB ID: 4R9O) as the
- 732 initial model. The apo-SmoF structure was solved using a dissected C-terminal domain of the
- 733 SmoF•SQGro structure. Structures were built and refined by iterative cycles using Coot (46) and
- 734 REFMAC (47) or Phenix (48), the latter employing local NCS restraints. Following building and

- refinement of the protein and water molecules, clear residual density was observed in the omit maps for co-complex structures, respective ligands were modelled into these. The coordinate and refinement library files were prepared using ACEDRG (49). The final structures gave R_{cryst} and R_{free}
- values along with data and refinement statistics that are presented in **Table S4-6.** Data were
- 739 collected at Diamond light source, Didcot, Oxfordshire, U.K., on beamlines I24 (SmoI-
- 740 D455N•SQGro, to 2.15 Å; SmoF-apo, to 1.88 Å), I04 (RoSmoC to 1.75 Å) and I04-1 (SmoC-apo,
- 741 to 3.2 Å; SmoB-apo_YSBLIC3C, to 1.5 Å; SmoB-apo; pET29a; SmoB•NADPH and
- 742 SmoB•NADPH•Glc) and at the Australian Synchrotron using the MX2 beamline (At3282•SQGro
- complex, to 1.7 Å). The coordinate files and structure factors have been deposited in the Protein
- 744 DataBank (PDB) with the coordinate accession numbers 70FX (SmoI-D455N•SQGro), 7NBZ
- 745 (SmoF-apo), 7OFY (SmoF•SQGro), 7OH2 (RoSmoC), 7OLF (SmoC-apo), 7BBY (SmoB-apo;
- 746 pET29a), 7BBZ (SmoB-apo; YSBLIC3C), 7BC0 (SmoB•NADPH) and 7BC1
- 747 (SmoB•NADPH•Glc).

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- 749 Structure-based analyses
- 750 Crystal packing interactions were analyzed using the protein interactions, surfaces, and assemblies
- 751 (PISA) server (50). Structural comparisons and structure-based sequence alignments were
- conducted using PDB25 search on DALI server against a representative subset of the Protein Data
- Bank (51). All structure figures were generated using ccp4mg (52).
- 755 Bioinformatic analysis SMO pathway prevalence
- Each gene within the A. tumefaciens C58 SMO gene cluster (Atu3277-Atu3285) was submitted as a
- query to the NCBI BLASTp algorithm to search a database comprised of non-redundant protein
- sequences with A. tumefaciens (taxid: 358) sequences excluded. Standard algorithm parameters
- were used, except the maximum target sequences was set to 10,000. Results were filtered to only
- 760 retain protein sequences with E-value $\leq 1.19 \times 10^{-51}$. The corresponding nucleotide accession
- numbers for each protein from all nine searches were extracted, combined and duplicates removed
- to provide a list of candidate genome sequences. This was converted into a reference library for
- 763 MultiGeneBLAST (53) and queried using the A. tumefaciens C58 SMO gene cluster. Clusters
- identified by this workflow with both an SQ monoxygenase and SQase homolog were regarded as
- putative SMO gene clusters. Clusters representative of the observed diversity were visualized using
- 766 Clinker (54). A phylogenetic tree of species possessing a putative SMO gene cluster was generated
- by pruning the All-Species Living Tree Project's 16s rRNA release 132 (55) using iTOL (56).

Data Availability Statement

Structure coordinates have been deposited in the Protein Data Bank (https://www.rcsb.org/) under accession codes 70FX, 70FY, 7NBZ, 70H2, 70LF, 7BBZ, 7BC0, 7BC1 and 7BBY. Proteomics data are available via ProteomeXchange (57) (http://www.proteomexchange.org/) with the identifier PXD014115. Scripts used to screen for the related gene clusters listed in Figure 5 is available on GitHub (https://github.com/jmui-unimelb/Gene-Cluster-Search-Pipeline).

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Additional information

- 934 Supplementary information
- Correspondence should be addressed to S.J.W, G.J.D or E.D.G.-B.

Figure 1. A. tumefaciens utilizes SQ and its glycosides as a carbon source. (a) Optical density of A. tumefaciens C58 culture (blue) and [SQ] (red), change in [sulfite] (green) and change in [sulfate] (yellow), with respect to time. This data is representative of two independent experiments (see Fig. S1), error bars denote observational error (derived by propagation of estimated random errors). (b) Manhattan plot of comparative proteomics data for A. tumefaciens C58 grown on SQ vs glucose, demonstrating that the most heavily upregulated proteins belong to a single gene cluster. (c) A cartoon of the upregulated cluster with automated annotations for each of the gene products. These would later be renamed smoABCDEFGHI, to reflect the importance of the sulfoquinovose monooxygenase enzyme activity to this new biochemical pathway. (d) A cartoon illustrating the hypothetical roles played by the gene products of this pathway to complete the catabolism of SQGro.

Figure 2. Biochemical and structural analyses of the SQGro-binding protein SmoF (Atu3282) and SQase SmoI (Atu3285). (a) Isothermal titration calorimogram for SmoF titrated against its cognate ligand 2'*R*-SQGro. The data is representative of two independent experiments (see **Fig. S5**). (b) Ribbon diagrams (with transparent surface) for the open and closed (liganded) conformations of SmoF. 2'*R*-SQGro is bound tightly in the inter-domain cleft and is inaccessible to the bulk solvent in the closed conformation. (c) Interactions between protein and ligand within the SmoF•2'*R*-SQGro complex: SmoF is in grey, 2'*R*-SQGro is in green, and the 2Fo – Fc map at 1.5σ is in blue. (d) A cartoon highlighting key interactions from c. (e) Interactions between protein and ligand within the complex pf SmoI-D455N SQase and 2'*R*-SQGro: SmoI is in gold, 2'*R*-SQGro is in green, and the 2Fo – Fc map at 1.5σ is in blue. (f) A cartoon highlighting key interactions from e: red spheres represent ordered water molecules; dotted lines represent proposed hydrogen bonds.

Figure 3. Biochemical and structural analyses of the flavin reductase SmoA and SQ monooxygenase SmoC. (a) Michaelis-Menten kinetics for SmoA-catalysed reduction of FMN by NADH. The data is representative of two independent replicates (see **Fig. S10**), error bars denote observational errors (derived by propagation of estimated random error). (b) SmoC activity assessed using sulfite release assay with Ellman's reagent in the presence of FMN, flavin reductase, NADH and SQ. The data is representative of two independent experiments (see **Fig. S11**), error bars denote observational error (derived by propagation of estimated random errors). (c) Isothermal titration calorimogram of interaction of SmoC with SQ as determined by ITC. The data is representative of two independent experiments (see **Fig. S13**). (d) Transparent molecular surface and ribbon diagram of *Ro*SmoC homodimer showing cofactor binding pocket and active site (dotted circle). (e) Alternative orientation of *Ro*SmoC monomer (in gold) overlaid with the MsuD·FMN·CH₃SO₃⁻ complex (7K14.pdb in ice blue) showing FMN from the latter. Expansion shows view of proposed substrate-binding pocket and conserved residues lining the active site of *Ro*SmoC.

Figure 4. Biochemical and structural analyses of 6-oxo-glucose reductase SmoB. (a) Top: Equilibrium oxygen exchange at C-6 of Glc via 6-OG facilitated by SmoB when incubated with NADP⁺ in H₂¹⁸O. Bottom: Derivatization and MS fragmentation allows localization of ¹⁸O to C6 of Glc. (b) Transparent molecular surface and ribbon diagram of SmoB in complex with NADPH and Glc. (c) Closeup view of SmoB•NADPH•Glc ternary complex. Backbone and carbon atoms of SmoB are shown in ice blue and NADPH and glucose are shown in cylinder format. Electron density for NADPH corresponds to the 2Fo – Fc map in blue at levels of 1σ. (d) Substrate binding pocket of SmoB depicting hydrogen bonding interactions of glucose with the active site residues including the conserved catalytic residues Asp71, Lys 104, His151 and Tyr76. Electron density corresponds to the 2Fo – Fc map (in blue) at levels of 1σ. The geometry of the SmoB-Glc complex indicates the likely trajectory of hydride addition to 6-OG. (e) Proposed mechanism of SmoB catalyzed reduction of 6-OG by NADPH showing hydride transfer from C4 of nicotinamide ring of NADPH to C6 carbonyl and Y76 (within the catalytic tetrad) as the proton donor. The red sphere is a bound water molecule; dotted lines are proposed hydrogen bonds.

Figure 5. Prevalence of the SMO pathway. (a) Architecture of the SMO gene cluster in *A. tumefaciens* and homologous gene clusters in other organisms. Colored links indicate ≥ 30% protein sequence similarity. Only those clusters encoding putative SQ monoxygenases and SQases were annotated as putative SMO gene clusters. (b) A phylogenetic tree demonstrating the diversity of organisms possessing putative SMO gene clusters. The tree was constructed by pruning of the All-Species Living Tree Project's 16s rRNA-based LTP release 132 (https://www.arb-silva.de/projects/living-tree/).









