UNIVERSITY of York

This is a repository copy of A sulfoglycolytic Entner-Doudoroff pathway in Rhizobium leguminosarum bv. trifolii SRDI565.

White Rose Research Online URL for this paper: <u>https://eprints.whiterose.ac.uk/161920/</u>

Version: Accepted Version

Article:

Li, Jinling, Epa, Ruwan, Scott, Nichollas E et al. (8 more authors) (2020) A sulfoglycolytic Entner-Doudoroff pathway in Rhizobium leguminosarum bv. trifolii SRDI565. Applied and Environmental Microbiology. ISSN 0099-2240

https://doi.org/10.1128/AEM.00750-20

Reuse

Items deposited in White Rose Research Online are protected by copyright, with all rights reserved unless indicated otherwise. They may be downloaded and/or printed for private study, or other acts as permitted by national copyright laws. The publisher or other rights holders may allow further reproduction and re-use of the full text version. This is indicated by the licence information on the White Rose Research Online record for the item.

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



A sulfoglycolytic Entner-Doudoroff pathway in *Rhizobium leguminosarum* bv. *trifolii* SRDI565

Jinling Li,¹ Ruwan Epa,¹ Nichollas E. Scott,³ Dominic Skoneczny,⁴ Mahima Sharma,⁵

Alexander J.D. Snow,⁵ James P. Lingford,² Ethan D. Goddard-Borger,² Gideon J. Davies,⁵

3

4

5

Malcolm J. McConville,⁴ Spencer J. Williams¹* 6 7 8 ¹School of Chemistry and Bio21 Molecular Science and Biotechnology Institute and 9 University of Melbourne, Parkville, Victoria 3010, Australia ²ACRF Chemical Biology Division, The Walter and Eliza Hall Institute of Medical Research, 10 11 Parkville, Victoria 3010, Australia and Department of Medical Biology, University of Melbourne, Parkville, Victoria 3010, Australia 12 ³ Department of Microbiology and Immunology, University of Melbourne at the Peter 13 14 Doherty Institute for Infection and Immunity, Parkville, VIC, 3010, Australia 15 ⁴Department of Biochemistry and Molecular Biology, Bio21 Molecular Science and 16 Biotechnology Institute, University of Melbourne, Parkville, Victoria 3010, Australia ⁵York Structural Biology Laboratory, Department of Chemistry, University of York, 17 18 Heslington YO10 5DD, United Kingdom 19 20 Keywords 21 Sulfoglycolysis; metabolomics; sulfur cycle; rhizobia; carbohydrates; metabolism

- 22
- 23 E-mail: sjwill@unimelb.edu.au
- 24
- 25

26 Abstract

27 Rhizobia are nitrogen fixing bacteria that engage in symbiotic relationships with plant hosts 28 but can also persist as free-living bacteria with the soil and rhizosphere. Here we show that 29 free living Rhizobium leguminosarum SRDI565 can grow on the sulfosugar sulfoquinovose 30 (SQ), or the related glycoside SQ-glycerol, using a sulfoglycolytic Entner-Doudoroff (sulfo-31 ED) pathway resulting in production of sulfolactate (SL) as the major metabolic end-product. 32 Comparative proteomics supports the involvement of a sulfo-ED operon encoding an ABC 33 transporter cassette, sulfo-ED enzymes and an SL exporter. Consistent with an oligotrophic 34 lifestyle, proteomics data revealed little change in expression of the sulfo-ED proteins during 35 growth on SQ versus mannitol, a result confirmed through biochemical assay of 36 sulfoquinovosidase activity in cell lysates. Metabolomics analysis showed that growth on SQ 37 involves gluconeogenesis to satisfy metabolic requirements for glucose-6-phosphate and 38 fructose-6-phosphate. Metabolomics analysis also revealed the unexpected production of 39 small amounts of sulfofructose and 2,3-dihydroxypropanesulfonate, which are proposed to 40 arise from promiscuous activities of the glycolytic enzyme phosphoglucose isomerase and a 41 non-specific aldehyde reductase, respectively. The discovery of a rhizobium isolate with the 42 ability to degrade SQ builds our knowledge of how these important symbiotic bacteria persist 43 within soil.

44 Importance

Sulfonate sulfur is a major form of organic sulfur in soils but requires biomineralization before it can be utilized by plants. Very little is known about the biochemical processes used to mobilize sulfonate sulfur. We show that a rhizobial isolate from soil, *Rhizobium leguminosarum* SRDI565, possesses the capability to degrade the abundant phototrophderived carbohydrate sulfonate SQ through a sulfoglycolytic Entner-Doudoroff pathway. Proteomics/metabolomics demonstrated the utilization of this pathway during growth on SQ

2

and provided evidence for gluconeogenesis. Unexpectedly, off-cycle sulfoglycolytic species were also detected pointing to the complexity of metabolic processes within cells under conditions of sulfoglycolysis. Thus rhizobial metabolism of the abundant sulfosugar SQ may contribute to persistence of the bacteria in the soil and to mobilization of sulfur in the pedosphere.

56 Introduction

57 Sulfur is essential for plant growth and is the fourth most important macronutrient after nitrogen, phosphorus, and potassium. Up to 10 kg/ha/y of sulfur is deposited in rain, 58 59 especially near industrialized areas (1). However, sulfur dioxide emissions from industrial 60 sources have decreased in recent decades as a result of pollution mitigation and the move to 61 low sulfur fuels and renewable energy sources, and quantities received from atmospheric 62 sources is now at levels below that required by most crops (2). Sulfur deficiency in soils is 63 primarily combated by application of sulfur-containing fertilizers such as superphosphate, 64 ammonium sulfate and gypsum (3), which are applied across all major cropping and pasture 65 areas worldwide (4). Soils contain significant amount of sulfur, yet plants can only use sulfur 66 in the form of sulfate and it has been shown that 95-98% of sulfur in soils is in the form of 67 unavailable biological sulfur (4). Thus, effective microbial cycling of sulfur from biological 68 to inorganic forms within the soil is important (5) and has the potential to enhance crop yields 69 and reduce reliance on fertilizers.

70 X-ray absorption near-edge spectroscopy measurements have led to estimates that 71 approx. 40% of sulfur within various sediments and humic substances exist as sulfonate (6). 72 Chemical methods of analysis applied to a range of forest soils revealed sulfonate sulfur was 73 at 40% of total organic sulfur pool in the majority of cases (7). Little detail is known on the 74 speciation of organic sulfonates in soils but one important input is phototroph-derived litter. 75 It is estimated that around 10 billion tonnes per annum of the sulfosugar sulfoquinovose (SQ) is produced annually by photosynthetic organisms, including plants, cyanobacteria and algae 76 77 (8). SQ is primarily found as the glycerolipid sulfoquinovosyl diacylglycerol (SQDG), and 78 land plants can contain as much as 10% SQDG in their thylakoid membrane glycerolipids 79 (9). Very little is known about how SQ is metabolized within soils, although it has been 80 shown to undergo very rapid mineralization to inorganic sulfate (10).

81 Bacteria are likely to be primarily responsible for the biomineralization of SQ, 82 possibly by using SQ as a carbon source and catabolizing it via a modified version of glycolysis, termed sulfoglycolysis (11). Two sulfoglycolytic processes have been described: 83 84 the sulfoglycolytic Embden-Meyerhof-Parnas (sulfo-EMP) pathway (12), and the 85 sulfoglycolytic Entner-Doudoroff (sulfo-ED) pathway (Fig. 1) (13). The sulfo-ED pathway 86 was first reported in *Pseudomonas putida* strain SQ1, a bacterium isolated from freshwater sediment, catabolised of SQ with excretion of equimolar amounts of sulfolactate (SL) (13). 87 88 The sulfo-ED operon of P. putida SQ1 contains 10 genes including a transcriptional 89 regulator, an SQ importer and SL exporter, a sulfoquinovosidase, SQ mutarotase, SQ 90 dehydrogenase, SL lactonase, SG dehydratase, KDSG aldolase and SLA dehydrogenase 91 enzymes. Based on genome-wide annotation studies, the sulfo-ED pathway is predicted to 92 occur in a range of alpha-, beta- and gamma-proteobacteria (13). However, no direct 93 evidence for this pathway has been reported for any organism other than P. putida SQ1. 94 Other members the microbial community can catabolize of SL and 2.3-95 dihydroxypropanesulfonate (DHPS; the product of the sulfo-EMP pathway) to inorganic 96 sulfur (14), completing the biomineralization of SQ.

97 Rhizobium leguminosarum bv. trifolii SRDI565 (syn. N8-J), hereafter Rl-SRDI565, 98 was isolated from a soil sample collected in western New South Wales but has the capacity to 99 colonize Trifolium subterraneum subsp. subterraneum (subterranean clover) and other 100 Trifolium spp. (15). Colonization of trifolium species with Rl-SRDI565 results in suboptimal 101 nodulation and nitrogen fixation in some species and ineffective nitrogen fixation in others, leading to reduced shoot nitrogen content relative to other commercial strains (16). 102 103 Interestingly, the genome of Rl-SRDI565 encodes all the genes needed for a functional sulfo-104 ED pathway (17), although there is no evidence to show that this is operative and/or that Rl-105 SRDI565 can use SQ as a major carbon source.

106 Rhizobia participate in sophisticated symbiotic relationships with leguminous host 107 plants that allow them to fix atmospheric dinitrogen to provide a growth advantage to the host 108 (18). Symbiosis is triggered by molecular communication between the bacterium and the host 109 resulting in nodule formation on the root and colonization by the bacterium. Within nodule 110 bacteroids the energy intensive fixation of nitrogen is supported by C₄-dicarboxylates 111 (primarily malate, fumarate, and succinate) sourced from glycolysis of sucrose photosynthate 112 within the plant host (18). Owing to the importance of biological nitrogen fixation for input 113 of nitrogen into the biosphere, the symbiosis of rhizobia and leguminous hosts has been well 114 studied. However, rhizobia can also exist as free-living bacteria within the soil and 115 rhizosphere (19). Here, like other soil bacteria, they adopt a saprophytic and oligotrophic 116 lifestyle where they utilize a variety of alternative carbon sources, including a wide range of 117 carbohydrates (20). Most likely, the ability of various rhizobia to persist in the pedosphere 118 depends upon their ability to utilize diverse carbohydrate and non-carbohydrate substrates 119 and establish an appropriate niche. SQ or its glycosides are likely to be a common soil 120 constituent and nutrient given its ubiquitous production by plants. Possibly, the sulfo-ED 121 pathway in *Rl*-SRDI565 might provide it with the capacity to survive on plant derived SQ or 122 SQDG in the rhizosphere and in the soil.

123 Here we investigated whether the sulfo-ED pathway is active in *Rl*-SRDI565 and its 124 potential role in utilizing plant-derived SQ or SQDG in the rhizosphere and in the soil. We 125 show that *Rl*-SRDI565 can grow on SQ and sulfoquinovosyl glycerol (SQGro) as sole carbon 126 source. Growth on SQ leads to excretion of SL into the growth media indicating active 127 sulfoglycolysis. This was supported by proteomic analyses, which showed that several 128 proteins in the sulfo-ED operon show increased expression when bacteria are grown on SQ, 129 while metabolomic analyses confirm the presence of characteristic intermediates of the sulfo-130 ED pathway, as well as the unexpected production of intracellular DHPS. Overall, we show

- 131 that *Rl*-SRDI565 has an active pathway for SQ utilization which may support growth of this
- 132 bacterium in the environment, and in turn provides a new model organism for the study of the
- 133 sulfo-ED pathway.
- 134
- 135

136 **Results**

137 Analysis of the genome of *Rl*-SRDI565 revealed a sulfo-ED operon that had the same genes, but no synteny with the P. putida SQ1 operon (Fig. 1). Genes with high sequence identity to 138 139 the P. putida proteins included a putative SQase, SQ dehydrogenase, SL lactonase, SG 140 dehydratase, KDSG aldolase and SLA dehydrogenase, and an SL exporter (see Fig. S1-S6). 141 The Rl-SRDI565 operon contains some important differences compared to that of P. putida 142 SQ1. In particular, it lacks a putative SQ mutarotase (21), and appears to use an ABC 143 transporter cassette to import SQ/SQGro in place of an SQ/SQGro importer/permease. The 144 putative sulfo-ED pathway in *Rl*-SRDI565 is consistent with the proposed protein functions 145 outlined in Fig. 1b, with a comparison to the classical ED pathway in Fig. 1c.

146 Initial attempts were made to grow *Rl*-SRDI565 in completely defined medium, such as M9 minimal media containing 125 μ g mL⁻¹ biotin (22), to allow assessment of different 147 148 carbon sources on bacterial growth. However, optimal growth could only be achieved using a 149 yeast extract-based medium (16). In particular robust growth was achieved using a 5% dilution of 1 g L^{-1} yeast extract (Y_{5%} media) containing 5 mmol mannitol (Y_{5%}M), while no 150 151 detectable bacterial growth was observed on Y_{5%} media alone. Significantly, Rl-SRDI565 also grew robustly on $Y_{5\%}$ media containing 5 mM SQ ($Y_{5\%}$ SQ) and reached the same final 152 OD₆₀₀ value as in Y_{5%}M (Fig. 2a). *Rl*-SRDI565 also grew on Y_{5%} media containing glucose, 153 although to a lower final OD₆₀₀ than in $Y_{5\%}M$ or $Y_{5\%}SQ$. ¹³C NMR spectroscopic analysis of 154 the culture media of stationary phase *Rl*-SRDI565 grown in Y_{5%}SQ revealed the presence of 155 156 three major signals corresponding to SL (Fig. 2b). A fourth signal was also observed but not 157 assigned and was also present in stationary phase media of cells grown on Y_{5%}M, suggesting it is derived from other carbon sources in the yeast extract. *Rl*-SRDI565 also grew on $Y_{5\%}$ 158 159 containing SQGro, but less robustly than on SQ.

160 We next examined changes in the proteome of Rl-SRDI565 cultivated on mannitol 161 versus SQ. Label-free based quantitative proteomic analysis of five experimental replicates 162 of *Rl*-SRDI565 cultivated on each carbon source, identified 2954 proteins, with 1943 proteins 163 quantified in at least 3 experimental replicates under each growth condition (Supplementary 164 Table 1). Expression levels of 17 proteins potentially associated with SQ metabolism were 165 significantly elevated $(-\log_{10}(p)>2)$ and a fold change greater than $2 \log_2 (p) \log_2 (p)$ in bacteria 166 cultivated in Y_{5%}SQ (Fig. 2e and 2f). In particular, a suspected KDSG aldolase (annotated as 167 alpha-dehydro-beta-deoxy-D-glucarate aldolase, WP_017967308.1), a member of the 168 proposed sulfo-ED pathway, was significantly increased $(-\log_{10}(p) = 4.74429)$ and a fold 169 change of 2.38 log₂). Consistent with the involvement of this pathway we also observed a 170 significant yet less dramatic increase in the proposed SQase (annotated alpha-glucosidase, 171 WP_017967311.1) ($-\log_{10}(p) = 1.43643$ and a fold change of $1.02 \log_2$). Additional members 172 of the predicted pathway expressed at higher levels in SQ-fed bacteria included the suspected 173 SQ dehydrogenase (annotated as SDR family oxidoreductase, WP_017967310.1) identified 174 by MS/MS events in 4 out of 6 SQ experiments compared to 1 mannitol experiment and the suspected SG dehydratase (annotated as dihydroxy-acid dehydratase, WP 017967307.1) 175 176 identified by MS/MS events in 3 out of 6 SQ experiments compared to 0 mannitol 177 experiments; however, owing to their low abundance they could not be accurately quantified 178 (Fig. S7).

179 Other proteins that were significantly increased in SQ-fed bacteria included a 180 NAD(P)-dependent oxidoreductase (WP 017965793.1), NADH-quinone oxidoreductase NAD-dependent 181 subunit NuoH (WP_017963854.1), succinate-semialdehyde а (WP_017967313.1) and a citrate synthase/methylcitrate 182 dehydrogenase synthase (WP_017964386.1) supporting an alteration with the TCA cycle and oxidative 183 184 phosphorylation under conditions of growth on SQ (Fig. 2f).

9

185 To demonstrate activity for a representative sulfo-ED enzyme from *Rl*-SRDI565, we 186 cloned and expressed the gene encoding the putative SQase. To support future structural studies, we expressed the N-terminal hexahistidine tagged K375A/K376A variant, termed 187 188 RlSQase*, a mutant enzyme whose design was guided by the Surface Entropy Reduction 189 prediction (SERp) server (Fig. S8) (23). Size exclusion chromatography-multiple angle light 190 scattering (SEC-MALS) analysis of *Rl*SQase* revealed that the protein exists as a dimer in 191 solution (Fig. S8). Enzyme kinetics were performed using the chromogenic SQase substrate 4-nitrophenyl α-sulfoquinovoside (PNPSQ). *Rl*SQase* exhibited a bell-shaped pH profile 192 193 with optimum at pH 7-8 and consistent with titration of catalytically important residues of 194 $pKa1 = 6.5 \pm 0.4$ and $pKa2 = 8.6 \pm 0.3$. The enzyme displayed saturation kinetics with Michaelis-Menten parameters $k_{cat} = 1.08 \pm 0.17 \text{ s}^{-1}$, $K_{M} = 0.68 \pm 0.25 \text{ mM}$, and $k_{cat}/K_{M} =$ 195 $(1.59 \pm 0.83) \times 10^3$ M⁻¹ s⁻¹ (Fig. 3a and 3b). For comparison, the kinetic parameters for A. 196 tumefaciens SQase is $k_{cat} = 22.3 \pm 0.6 \text{ s}^{-1}$, $K_{M} = 0.21 \pm 0.03 \text{ mM}$, $k_{cat}/K_{M} = (1.1 \pm 0.1) \times 10^{5}$ 197 M^{-1} s⁻¹ and *E. coli* SQase YihQ is $k_{cat} = 32.7 \pm 0.6$ s⁻¹, $K_M = 0.15 \pm 0.01$ mM, $k_{cat}/K_M = (2.2)$ 198 ± 0.2) × 10⁵ M⁻¹ s⁻¹ (24). 199

200 Direct evidence for enzymatic activity associated with the sulfo-ED operon in Rl-201 SRDI565 was obtained by measuring SQase enzyme activity in cell lysates. The chromogenic 202 substrate 4-nitrophenyl α -sulfoquinovoside (PNPSQ), which was designed as an analogue of 203 the natural substrate SQGro, results in release of the chromophore 4-nitrophenolate, which 204 can be detected using UV-visible spectrophotometry with high sensitivity at 400 nm or at the isosbestic point, 348 nm (24, 25). Rl-SRDI565 was grown to mid-logarithmic phase in Y_{5%}M 205 206 and Y_{5%}SQ media, and the harvested cells used to prepare a cell-free lysate containing 207 soluble proteins. Incubation of Y_{5%}M and Y_{5%}SQ-derived lysates with PNPSQ both resulted 208 in production of 4-nitrophenolate at similar rates. The activity in the YSQ-derived lysate was inhibited by the addition of IFG-SQ, an azasugar inhibitor of SQases that makes key 209

interactions in the active site that mimic those required for substrate recognition (Fig. 3c)
(24). The similar levels of activity of SQase in both mannitol and SQ grown *Rl*-SRDI565 is
consistent with the abundance of the putative SQase WP_017967311.1 detected by proteomic
analysis.

214 To further confirm that a sulfo-ED pathway was operative in cells, a targeted 215 metabolomics approach was used to detect expected intermediates in bacteria grown on 216 Y_{5%}SQ media. Detected intermediates were identified based on their LC-MS/MS retention 217 time and mass spectra with authentic reference standards of the sulfo-EMP and sulfo-ED 218 pathway that were synthesized in-house. Sulfogluconate (SG) was synthesized by oxidation 219 of SQ with iodine (26) (Fig. S9), while SQ, SF, SFP, DHPS, SLA and SL were prepared as 220 previously reported (27). Rl-SRDI565 was grown to mid-log phase in Y_{5%}M or Y_{5%}SQ, 221 metabolically quenched and extracted polar metabolites analyzed by LC/MS-MS. SQ-grown 222 bacteria contained SQ, SF, SG, SL and DHPS, while SFP and SLA could not be detected 223 (Fig. 4a-e). The detection of SG is characteristic for a sulfo-ED pathway, and presumably 224 arises from the action of the putative SQ dehydrogenase and SGL lactonase. The 225 identification of DHPS and SF was unexpected, as these intermediates/products of the sulfo-226 EMP pathway (12). BLAST analysis of the genome of *Rl*-SRDI565 did not identify putative 227 genes for the sulfo-EMP pathway. SF may therefore be formed by the action of 228 phosphoglucose isomerase (PGI), while DHPS could be the product of a promiscuous 229 aldehyde reductase. Rl-SRDI565 was unable to utilize DHPS or SL as sole carbon source in 230 $Y_{5\%}$ medium, supporting the absence of an alternative pathway of sulfoglycolysis that utilizes 231 these intermediates. Unexpectedly, cytosolic levels of DHPS were 20-fold higher than SL, 232 suggesting that cells may lack a membrane transporter to export accumulated DHPS, in 233 contrast to the SL transporter.

234	NMR and LC-MS/MS analysis of the culture supernatant of both unlabeled and
235	$(^{13}C_6)$ -labelled SQ-cultivated <i>Rl</i> -SRDI565 confirmed that the substrate is almost completely
236	consumed by the time bacteria reach stationary growth (final concentration of 0.006±0.001
237	mM compared to 5.0±0.5 mM SQ in starting medium) (Fig. S10). Using a highly sensitive
238	cryoprobe ¹³ C NMR spectroscopic analysis revealed that both DHPS and SG were present in
239	culture supernatant of ¹³ C ₆ -SQ-cultivated <i>Rl</i> -SRDI565. Quantitative LC-MS/MS analysis
240	showed that consumption of SQ was associated with production of SL (5.70±0.12 mM), and
241	low levels of DHPS (0.081±0.010 mM), (SG 0.172±0.006 mM) and SF (0.002±0.0001 mM)
242	(Table 1). This experiment was repeated to assess the effect of growth of Rl-SRDI565 but
243	using SQGro as carbon source. As noted previously, Rl-SRDI565 grows inconsistently on
244	SQGro and complete consumption of SQGro could not be achieved. However, the results of
245	partial consumption broadly agreed with the results for growth on SQ, namely that SL is the
246	major terminal metabolite detected in the culture media, with much lower amounts of SF, SG
247	and DHPS (Table 1).

249 **Discussion**

250 We demonstrate here that *Rl*-SRDI565 has a functional sulfo-ED pathway that allows these 251 bacteria to utilize SQ as their major carbon source. Catabolism of SQ is primarily or 252 exclusively mediated by a sulfo-ED pathway with production of SL as the major end-product, 253 similar to the situation in *P. putida* SQ1, the only other experimentally described exemplar of 254 this pathway (13). In contrast to P. putida SQ1, Rl-SRDI565 also produces trace amounts of DHPS which could reflect the presence of enzymes which exhibit promiscuous activities 255 256 similar to those in the conventional sulfo-EMP pathway. This observation is reminiscent of 257 Klebsiella sp. strain ABR11 isolated from soil (28) that is also able to grow on SQ with 258 production of both SL and DHPS. *Klebsiella* sp. strain ABR11 possesses an NAD⁺-specific 259 sulfoquinovose-dehydrogenase activity (29), suggesting it has an operative sulfo-ED 260 pathway.

Various bacteria that can metabolize SQ have been isolated from soil including *Agrobacterium* sp., (29) *Klebsiella* sp., (29) and *Flavobacterium* sp., (30) as well as *P. putida* SQ1 (13), which was isolated from a freshwater littoral sediment. These bacteria may work cooperatively with species such as *Paracoccus pantotrophus* NKNCYSA that can convert SL to mineral sulfur, leading to stoichiometric recovery of sulfite/sulfate (14). Together these bacterial communities achieve the complete mineralization of SQ to sulfate, which is available for use by plants.

Proteomic and biochemical evidence suggests that the sulfo-ED pathway is constitutively expressed in *Rl*-SRDI565, with only relatively small increases in protein expression as shown by statistically significant increases in only KDSG aldolase and SQase. in the presence of SQ. As *Rl*-SRDI565 in the soil is likely to be oligotrophic, constitutive expression of the sulfo-ED pathway may allow simultaneous usage of multiple nonglycolytic substrates without requirement for significant transcriptional changes. Consistent 274 with this view, the proteomic abundance of the putative LacI-type regulator 275 WP_017967302.1 was unchanged between mannitol and SQ grown Rl-SRDI565. The sulfo-276 ED operon in *Rl*-SRDI565 differs from that described for *P. putida* SQ1 through the absence 277 of a putative SQ mutarotase. SQ undergoes mutarotation with a half-life of approximately 6 278 h, which is much slower than for the glycolytic intermediate Glc-6-P, which has a half-life of 279 just seconds (21). Aldose mutarotases are often relatively non-specific and possibly a 280 constitutive mutarotase not in the sulfo-ED operon expressed by the cell provides this 281 catalytic capacity. Alternatively, the SQ dehydrogenase may not be stereospecific, with the 282 ability to act on both anomers of SQ, or even that it acts on α -SQ (the product released from 283 SQGro by an SQase) at a high rate such that mutarotation to β -SQ is of insignificant 284 importance. A second difference in the sulfo-ED operon lies in the presence of an ABC 285 transporter cassette. ABC transporter cassettes are the most common solute transporters, and 286 can translocate their substrates in either a forward or reverse direction (31). While we 287 propose that the ABC transporter cassette operates in the forward direction, based on the presence of a signal sequence in the putative solute binding domain targeting it to the 288 289 periplasm, and consistent with a wide range of sugar import systems, the directionality of 290 transport and thus the choice of substrate (SQ/SQGro versus SL) may depend on the relative 291 abundance of these metabolites intra and extracellularly.

Sulfoglycolysis in *Rl*-SRDI565 leads to production of pyruvate and the excretion of the C3-organosulfonate SL (Fig. 5). In order to satisfy the demands of the pentose phosphate pathway and cell wall biogenesis, sulfoglycolytic cells must synthesize glucose-based metabolites such as glucose-6-phosphate and glucose-1-phosphate. Gluconeogenesis has been studied in *Rhizobium leguminosarum* strain MNF3841, and operates through a classical pathway involving fructose bisphosphate aldolase (32). Action of phosphoglucose isomerase on SQ might lead to production of SF, thereby explaining the observation of this metabolite 299 in *Rl*-SRDI565. This is not likely to be consequential, as the reversibility of this reaction will 300 ultimately allow complete consumption of any SF through isomerization back to SQ. The 301 formation of DHPS may result from a promiscuous aldehyde reductase. Analysis of spent culture media reveals that the production of DHPS is minor in terms of total carbon balance. 302 303 However, within the cytosol, DHPS accumulates to levels much higher than SL, presumably 304 because of the absence of a dedicated exporter for the former. Possibly, reduction of SLA to 305 DHPS is reversible and enables conversion of this metabolite to SL and subsequent excretion 306 from the cell. The observation of SG, SF and DHPS in the spent culture media at low levels 307 is suggestive of low levels of leakage of these metabolites from the cell, either through cell 308 lysis or leaky export systems.

309 Given that SQ contains a significant portion of organic sulfur within plants, the 310 pathways of SQ catabolism leading to release of its sulfur may be important to enable 311 recycling of this important macronutrient. Plants can only use sulfate, which is poorly 312 retained by most soils. Biomineralization of organic sulfur to sulfate is important to allow 313 plants to access this element. As one of just two known pathways for the catabolism of SQ, the sulfo-ED pathway is likely to be an important part of environmental breakdown of SQ 314 315 and may contribute to the persistence of symbiotic rhizobia within the pedosphere. The 316 present work lays the groundwork for a more detailed investigation of sulfoglycolysis in a 317 well-characterized bacterium with an established capability for symbiosis of a leguminous 318 plant host.

319

320 Materials and Methods

321 **Reagents**

322 SQ, $({}^{13}C_6)$ -SQ, SF, SFP, SLA, SL, and DHPS were chemically and chemoenzymatically 323 synthesized as described previously (27). IFG-SQ was chemically synthesized as described 324 (24).

325 Bacteria and culture conditions

326 *Rhizobium leguminosarum* bv. *trifolii* SRDI565 was a gift from Dr Ross Ballard (South 327 Australian Research and Development Institute, Adelaide, South Australia). Minimal salts 328 media consists of $0.5 \text{ g} \cdot \text{L}^{-1} \text{ K}_2 \text{HPO}_4$, $0.2 \text{ g} \cdot \text{L}^{-1} \text{ MgSO}_4$, $0.1 \text{ g} \cdot \text{L}^{-1} \text{ NaCl}$, 1 M CaCl₂ 3 mL·L⁻¹, 329 adjusted to pH 7.0. YM media consists of minimal salts media plus 1 g·L⁻¹ yeast extract, 10 330 g·L⁻¹ mannitol. Y_{5%}M consists of minimal salts media plus 50 mg·L⁻¹ yeast extract, 5 mM 331 mannitol. Y_{5%}SQ consists of minimal salts media plus 50 mg·L⁻¹ yeast extract, 5 mM SQ.

332

Growth curves were determined in a MicrobeMeter built in-house according to published plans (33) and blueprints available at <u>https://humanetechnologies.co.uk/</u> The MicrobeMeter was calibrated by performing serial 2-fold dilutions across the detection range of the MicrobeMeter (0-1023 units), starting with an OD_{600} approx. 1 culture of *Rl*-SRDI565. OD_{600} measurements were made with a UV/Vis spectrophotometer and plotted against the reading of the MicrobeMeter. The data was fit to a polynomial to obtain a calibration curve.

339

340 **Proteomic sample preparation:** Cells were washed 3 times in PBS and collected by 341 centrifugation at 10,000 x g at 4°C then snap frozen. Frozen whole cell samples were 342 resuspended in 4% SDS, 100 mM Tris pH 8.0, 20 mM DTT and boiled at 95°C with shaking 343 at 2000 rpm for 10 min. Samples were then clarified by centrifugation at 17,000 × g for 10 344 min, the supernatant collected, and protein concentration determined by bicinchoninic acid 345 assay (Thermo Scientific Pierce). 100 µg of protein from each sample was cleaned up using 346 SP3 based purification according to previous protocols (34). Briefly, reduced samples were cooled and then alkylated with 40 mM 2-chloroacetamide (CAA) for 1 hour at RT in the 347 348 dark. The alkylation reactions were then guenched with 40 mM DTT for 10 min and then 349 samples precipitated onto SeraMag Speed Beads (GE Healthcare, USA) with ethanol (final 350 concentration 50% v/v). Samples were shaken for 10 min to allow complete precipitation 351 onto beads and then washed three times with 80% ethanol. The precipitated protein-covered 352 beads were resuspended in 100 mM ammonium bicarbonate containing 2 µg trypsin (1/50 353 w/w) and allowed to digest overnight at 37 °C. Upon completion of the digests, samples were 354 centrifuged at 14000 g for 5 min to pellet the beads and the supernatant collected and 355 desalted using homemade C18 stage tips (35). The eluted material was dried and stored, until 356 analysed by LC-MS.

357

358 Proteomics analysis using reversed phase LC-MS: Purified peptides prepared were re-359 suspended in Buffer A* (2% ACN, 0.1% CF₃CO₂H) and separated using a two-column 360 chromatography set-up composed of a PepMap100 C18 20 mm × 75 µm trap and a PepMap 361 C18 500 mm \times 75 μ m analytical column (Thermo Fisher Scientific). Samples were 362 concentrated onto the trap column at 5 µL/min for 5 min and infused into an Orbitrap Elite™ 363 (Thermo Fisher Scientific). 120 min gradients were run altering the buffer composition from 364 1% buffer B (80% ACN, 0.1% formic acid) to 28% B over 90 min, then from 28% B to 40% 365 B over 10 min, then from 40% B to 100% B over 2 min, the composition was held at 100% B for 3 min, and then dropped to 3% B over 5 min and held at 3% B for another 10 min. The 366 367 Elite Orbitrap Mass Spectrometers was operated in a data-dependent mode automatically switching between the acquisition of a single Orbitrap MS scan (120,000 resolution) and a 368 maximum of 20 MS-MS scans (CID NCE 35, maximum fill time 100 ms, AGC $1*10^4$). 369

370

371 Mass spectrometry data analysis. Proteomic comparison of growth with and without 372 sulfoquinovose was accomplished using MaxQuant (v1.5.5.1) (36). Searches were performed 373 against Rhizobium leguminosarum bv. trifolii SRDI565 (NCBI Taxonomy ID: 935549, 374 downloaded 01-08-2019, 6404 entries) with carbamidomethylation of cysteine set as a fixed 375 modification. Searches were performed with Trypsin cleavage allowing 2 miscleavage events and the variable modifications of oxidation of methionine and acetylation of protein N-376 377 termini. The precursor mass tolerance was set to 20 parts-per-million (ppm) for the first 378 search and 10 ppm for the main search, with a maximum false discovery rate (FDR) of 1.0% 379 set for protein and peptide identifications. To enhance the identification of peptides between 380 samples the Match Between Runs option was enabled with a precursor match window set to 2 381 min and an alignment window of 10 min. For label-free quantitation, the MaxLFQ option 382 within MaxQuant (37) was enabled in addition to the re-quantification module. The resulting 383 peptide outputs were processed within the Perseus (v1.4.0.6)(38) analysis environment to 384 remove reverse matches and common protein contaminates with missing values imputed. The 385 mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium 386 via the PRIDE partner repository with the dataset identifier PXD015822.

387

388 Enzyme kinetics of *RISQase*

389 **Michaelis Menten plot.** Kinetic analysis of *RISQase** was performed using PNPSQ as 390 substrate, using a UV/visible spectrophotometer to measure the release of the 4-391 nitrophenolate ($\lambda = 348$ nm). Assays were carried out in 50 mM sodium phosphate, 150 mM 392 NaCl, pH 7.2 at 30 °C using 212 nM *RISQase** at substrate concentrations ranging from 0.05 393 µM to 4 mM. Using the extinction coefficient for 4-nitrophenolate of 5.125 mM⁻¹ cm⁻¹, 394 kinetic parameters were calculated using Prism. **pH profile.** For the determination of pH profile, specific activities of *RI*SQase* were monitored by measuring absorbance changes at $\lambda = 348$ nm in the presence of the following buffers: sodium acetate buffer (pH 5.6, sodium phosphate buffer (pH 6.0–8.5), and glycine NaOH buffer (pH 8.8–9.2). The assays were performed at 30 °C in duplicates and specific activities determined using extinction coefficient of PNP 5.125 mM⁻¹ cm⁻¹ at the isosbestic point (348 nM). One unit of SQase activity is defined as the amount of protein that releases 1 µmol PNP per min.

402

403 Cloning, expression and kinetic analysis of *Rl*-SRDI565 sulfoquinovosidase (*Rl*SQase*)

404 The gene sequence coding for RlSQase* SERp mutant was synthesised with codon 405 optimisation for expression in E. coli and was cloned within a pET-28a(+) vector with C-406 terminal His-tag through GenScript. The plasmid His₆-*Rl*SQase*-pET-28a(+) containing the 407 gene for target RlSQase* was transformed into E. coli BL21(DE3) cells for protein 408 expression. Pre-cultures were grown in LB-medium (5 mL) containing 30 µg/mL for 18 h at 409 37 °C, 200 rpm. Cultures (1 L LB-medium supplemented with kanamycin 30 µg/mL) were 410 inoculated with the pre-culture (5 mL) and incubated at 37 °C, 200 rpm until an OD₆₀₀ of 0.6-411 0.8 was achieved. Protein expression was induced by addition of IPTG (1 mM) and shaking 412 was continued overnight (20-22 h) at 18 °C, 200 rpm. The cells were harvested by centrifugation (5000 rpm, 4 °C, 20 min), resuspended in 50 mM Tris, 300 mM NaCl pH 7.5 413 414 buffer and were subjected to further cell lysis. Cells were disrupted using French press under 415 20 k Psi pressure and the lysate was centrifuged at 50,000 g for 30 min.

The N-terminal His₆-tagged protein was purified by immobilized metal ion affinity chromatography, followed by size exclusion chromatography (SEC) (Fig. S8). The lysate was loaded onto a pre-equilibrated Ni-NTA column, followed by washing with load buffer (50 mM Tris-HCl, 300 mM NaCl, 30 mM imidazole pH 7.5). The bound protein was eluted using 420 a linear gradient with buffer containing 500 mM imidazole. Protein containing fractions were 421 pooled, concentrated and loaded onto a HiLoad 16/600 Superdex 200 gel filtration column 422 pre-equilibrated with 50 mM Tris-HCl, 300 mM NaCl pH 7.5 buffer. The protein was 423 concentrated to a final concentration of 60 mg mL⁻¹ using a Vivaspin® 6 with a 300 kDa 424 MW cut-off membrane for characterization and enzyme assays.

425

426 SEC-MALS analysis

427 Experiments were conducted on a system comprising a Wyatt HELEOS-II multi-angle light 428 scattering detector and a Wyatt rEX refractive index detector linked to a Shimadzu HPLC 429 system (SPD-20A UV detector, LC20-AD isocratic pump system, DGU-20A3 degasser and 430 SIL-20A autosampler). Experiments were conducted at room temperature ($20 \pm 2^{\circ}$ C). 431 Solvents were filtered through a 0.2 µm filter prior to use and a 0.1 µm filter was present in 432 the flow path. The column was equilibrated with at least 2 column volumes of buffer (50 mM 433 Tris, 300 mM NaCl pH 7.5) before use and buffer was infused at the working flow rate until 434 baselines for UV, light scattering and refractive index detectors were all stable. The sample injection volume was 100 µL R/SQase* at 6 mg/mL in 50 mM Tris buffer, 300 mM NaCl pH 435 436 7.5. Shimadzu LC Solutions software was used to control the HPLC and Astra V software for the HELEOS-II and rEX detectors (Fig. S8). The Astra data collection was 1 min shorter than 437 438 the LC solutions run to maintain synchronisation. Blank buffer injections were used as 439 appropriate to check for carry-over between sample runs. Data were analysed using the Astra V software. Molar masses were estimated using the Zimm fit method with degree 1. A value 440 441 of 0.158 was used for protein refractive index increment (dn/dc).

442

443 Detection of SQase activity in cell lysates

444 *Rl*-SRDI565 was grown in 50 mL Y_{5%}M and Y_{5%}SQ media at 30 °C to mid log phase, 445 approximately $OD_{600} = 0.2$, measured using a Varian Cary50 UV/visible spectrophotometer. Cells were harvested by adding 3x volume of ice-cold PBS to metabolically quench the 446 447 samples then centrifuged at 2000 g, 4 °C for 10 min. The supernatant was discarded and the cells were washed 3 times with ice-cold PBS, with each wash involving resuspension and 448 centrifugation at 2000 g, 4 °C for 10 min. The cells were collected once more by 449 centrifugation at 10,000 g, 4 °C, for 1 min then snap frozen in liquid nitrogen and stored at -450 80 °C. 451

452 Cells were lysed by addition of 1000 µL pre-chilled PBS, 1 µL RNaseA, 1 µL DNase, 1 μ L 100 mg·mL⁻¹ hen egg white lysozyme (Sigma), and a 1× final concentration of 453 454 cOmplete EDTA-free protease inhibitor cocktail (Roche) to the cell pellet. The cells were 455 gently resuspended and mixed at 4 °C for 10 min. The suspension was placed on ice and 456 irradiated with a Sonoplus HD3200 MS 73 sonicator probe (Bandelin) at a frequency of 20 457 kHz, 20% amplitude, pulse 2s on 8s off, repeated for a total time of the sonication to 150 s, 458 then incubated on ice for 5 min. The suspension was clarified by centrifuging at 14,000 g, 4 °C for 1 min and the supernatant was filtered through a Nanosep mini centrifugal spin 459 column with a 0.2 µm filter (Pall) into a 1.5mL Eppendorf tubes and stored at 4 °C. Protein 460 461 concentration was determined using a BCA assay.

462 SQase activity was measured in triplicate using PNPSQ and an Agilent Cary UV 463 Workstation (G5191-64000) at 30°C. Reactions contained buffer consisted of 50 mM NaPi 464 and 150 mM NaCl, pH=7.4, and 2.5 mM PNPSQ. Reactions were initiated by addition of 465 SQ- or mannitol-derived lysate to a final concentration of 43.7 μ g·mL⁻¹ protein, and 466 absorbance was monitored at 400 nm for 3 h. After 3 h, IFGSQ was added to a final 467 concentration of 6.25 mM to the SQ-lysate sample, and absorption monitored for 3 h.

468

469 Metabolite analysis of *Rhizobium leguminosarum* cell extracts

470 Metabolic quenching and extraction. *Rl*-SRDI565 was grown on Y_{5%}SQ or Y_{5%} containing 471 35 mM glucose to mid-logarithmic phase (approx. 0.15), as calculated based on the OD_{600} 472 measured by Cary 50 UV/visible spectrophotometer, and were rapidly quenched in a 473 prechilled 15 mL Falcon tube containing phosphate buffered saline (PBS) at 4 °C. Ice-cold 474 PBS (11 mL) was infused into cell culture media (4 mL). The Falcon tubes were mixed by inversion and incubated in ice/water slurry for 5 min then were centrifuged at $2000 \times g$ at 1 475 476 °C for 10 min. The supernatant was removed by aspiration and cell pellets were washed twice 477 with 1 mL of ice-cold PBS (with resuspension each time) and transferred into 1.5 mL 478 Eppendorf tubes. Cells were pelleted by centrifugation at 14000 rpm and residual solvent was 479 carefully removed. Cell pellets were stored at -80°C until extraction. Cells were extracted in 480 200 μ L of extraction solution (methanol/water, 3:1 v/v) containing an internal standard, 5 μ M 481 $^{13}C_4$ -aspartate (Cambridge Isotopes), and subjected to 10 freeze-thaw cycles to facilitate cell 482 lysis (30 s in liquid nitrogen, followed by 30 s in dry ice/ethanol bath). Debris was pelleted 483 by centrifugation at 14000 rpm, 5 min, 1°C and cell lysate was transferred into a HPLC vial 484 insert for LC/MS analysis.

485 LC/MS analysis and identification of sulfonate metabolites. Separation and detection of 486 polar metabolites was performed using an Agilent Technologies 1200 series high 487 performance liquid chromatography (HPLC) coupled to a quadrupole time-of-flight mass spectrometer (6545 QTOF, Agilent Technologies) using a method modified from 488 489 Masukagami et al. (39). Metabolite extracts were transferred into 2 mL auto sampler vials 490 with glass inserts and placed in the auto sampler kept at 4 °C prior to analysis. Metabolite 491 separation was performed by injecting 7 µL of the extract into a SeQuant® ZIC-pHILIC 492 PEEK coated column (150 mm × 4.6 mm, 5 µm polymer, Merck Millipore) maintained at 493 25°C, with a gradient of solvent A (20 mM ammonium carbonate, pH 9.0, Sigma-Aldrich) 494 and solvent B (100% acetonitrile, Hypergrade for LCMS LiChrosolv, Merck) at a flow rate 495 of 0.3 mL/min. A 33.0 min gradient was setup with time (t) = 0 min, 80% B; t = 0.5 min, 496 80% B; t = 15.5 min, 50% B; t = 17.5 min, 30% B; t = 18.5 min, 5% B; t = 21.0 min, 5% B; t497 = 23.0 min, 80% B.

498 The LC flow was directed into an electrospray ionization (ESI) source with a capillary 499 voltage of 2500 V operating in negative ionization mode. Drying nitrogen gas flow was set to 500 10 L/min, sheath gas temperature and nebulizer pressure were set to 300 °C and 20 psi, 501 respectively. The voltages of fragmentor and skimmer were set at 125 V and 45 V, 502 respectively. Data was acquired in MS and MS/MS mode, with a scan range of 60 to 1700 503 m/z and 100 to 1700 m/z respectively, at a rate of 1.5 spectra/sec. MS/MS acquisition was 504 performed with four collision energies (0, 10, 20 and 40 V). The mass spectrometer was 505 calibrated in negative mode prior to data acquisition and mass accuracy during runs was 506 ensured by a continuous infusion of reference mass solution at a flow rate of 0.06 mL/min 507 (API-TOF Reference Mass Solution Kit, Agilent Technologies). Data quality was ensured by 508 multiple injections of standards (with 1.5 µM concentration each) and pooled biological 509 sample (a composite of cell extracts) used to monitor the instrument performance. Samples 510 were randomized prior to metabolite extraction and LC/MS analysis.

Standard preparation. Standards of selected metabolites (Supplementary Table 1) were prepared at 10 μ M in 80% acetonitrile (Hypergrade for LCMS LiChrosolv, Merck) and injected separately into a column connected to mass spectrometer interface. Retention time and detected molecular ion were used to create a targeted MS/MS acquisition method. The spectra, mass to charge (*m*/*z*) and retention time (RT) were imported into a personal compound database and library (PCDL Manager, version B.07.00, Agilent Technologies) used in data processing workflow. **Data analysis.** Data were analysed using MassHunter Qualitative and Quantitative Analysis software (version B.07.00, Agilent Technologies). Identification of metabolites was performed in accordance with metabolite identification (Metabolomics Standard Initiative, MSI) level 1 based on the retention time and molecular masses matching to authentic standards included in the personal database and library. Peak integration was performed in MassHunter quantitative software (version B.07.00, Agilent Technologies) on the spectra from identified metabolites.

525

526 Chemical synthesis of 6-deoxy-6-sulfo-D-gluconate (SG)

527 NaOH in methanol (4% w/v, 4 mL) was added dropwise to a stirred solution of 528 sulfoquinovose (100 mg, 0.410 mmol) and iodine (209 mg, 1.65 mmol) in water (1 mL) and 529 methanol (4 mL) held at 40 °C. As the sodium hydroxide was added the color of iodine dissipated. The solvent was evaporated under reduced pressure and the crude residue was 530 531 subjected to flash chromatography (EtOAc/MeOH/H₂O, 4:2:1 to 2:2:1, then water) to give 532 the 6-deoxy-6-sulfogluconate sodium salt (89.2 mg). An aqueous solution of the sodium salt 533 was eluted through a column of Amberlite IR120 (H⁺ form) resin. The acidic eluate was collected and concentrated under reduced pressure give SG (71.3 mg, 67%). ¹H NMR (400 534 535 MHz, D_2O): δ 4.23–4.15 (1 H, m, H2), 4.13 (1 H, d, J = 3.3 Hz, H3), 4.05 (1 H, t, J = 3.2 Hz, H5), 3.74 (1 H, dd, J = 6.5, 3.4 Hz, H4), 3.35 (1 H, d, J = 14.6 Hz, H6a), 3.05 (1 H, dd, J = 536 14.6, 9.7 Hz, H6b); ¹³C{¹H} NMR (100 MHz, D₂O) δ 178.7 (C1), 74.2 (C4), 73.8 (C2), 70.8 537 538 (C3), 67.8 (C5), 53.4 (C6); HRMS (ESI) calcd for C₆H₁₁O₉S [M⁻] 259.0129, found 259.0131. 539

540

541 Quantitation of metabolite levels in spent culture media

542 The metabolites (DHPS, SF, SQ, SL and SG) present in spent culture media were quantified 543 against standard solutions of pure metabolites by HPLC-ESI-MS/MS. Quantification was done with the aid of calibration curves generated by dissolving the pure standards in spent 544 545 media from *Rl*-SRDI565 grown on Y_{5%}M. Spiked spent media was diluted 100-fold with 546 water and then analysed by LC-MS/MS with α -MeSQ as internal standard. For experimental 547 determination of metabolites, spent culture media from *Rl*-SRDI565 grown in Y_{5%}SQ or $Y_{5\%}$ SQGro were diluted 100-fold with water and analysed by LC-MS/MS with α -MeSQ as 548 549 internal standard.

550 HPLC-ESI-MS/MS analysis was performed using a TSQ Altis triple quadrupole mass 551 spectrometer (Thermo Fisher Scientific) coupled with a Vanquish Horizon UHPLC system 552 (Thermo Fisher Scientific). The column was a ZIC-HILIC column (5 μ m, 50 \times 2.1 mm; 553 Merck). The HPLC conditions were: from 90% B to 40% B over 15 min; then 40% B for 5 min; back to 90% B over 1 min (solvent A: 20 mM NH₄OAc in 1% acetonitrile; solvent B: 554 acetonitrile); flow rate, 0.30 ml min⁻¹; injection volume, 1 µl. The mass spectrometer was 555 556 operated in negative ionization mode. Quantitation was done using the MS/MS selected 557 reaction monitoring (SRM) mode using Thermo Scientific XCalibur software and normalized 558 with respect to the internal standard, α -MeSQ. Prior to analysis, for each analyte, the 559 sensitivity for each SRM-MS/MS transition was optimized.

560 DHPS: ESI-MS/MS m/z of [M-H]⁻ 155, product ions 137, 95; retention time: 4.91 min

- 561 α -MeSQ (internal standard): ESI–MS/MS *m*/*z* of [M-H]⁻ 257, product ions 166, 81;
- 562 retention time: 6.31 min
- 563 SF: ESI-MS/MS m/z of [M-H]⁻ 243, product ions 207, 153; retention time: 6.81 min
- 564 SQ: ESI–MS/MS m/z of [M-H]⁻ 243, product ions 183, 123; retention time: 7.58 and 7.89 565 min for α / β
- 566 SL: ESI-MS/MS m/z of [M-H]⁻ 169, product ions 107, 71; retention time: 9.26 min

567 SG: ESI–MS/MS m/z of [M-H]⁻ 259, product ions 241, 161; retention time: 9.66 min 568 SQGro: ESI–MS/MS m/z of [M-H]⁻ 317, product ions 225, 165; retention time: 7.15 min 569

570 Acknowledgements

571 This work was supported by grants from the Australian Research Council (DP180101957), 572 the National Health and Medical Research Council of Australia (APP1100164, GNT1139549) and the Leverhulme Trust; support from The Walter and Eliza Hall Institute of 573 574 Medical Research, the Australian Cancer Research Fund, and a Victorian State Government 575 Operational Infrastructure support grant. MJM is an NHMRC Principal Research Fellow, 576 G.J.D. is a Royal Society Ken Murray Research Fellow. JL is supported by a PhD scholarship 577 from the Chinese Scholarship Council. We thank Humane Technologies for support with the 578 MicrobeMeter, the Melbourne Mass Spectrometry and Proteomics Facility of the Bio21 579 Institute at the University of Melbourne, Palika Abayakoon and Janice Mui for reagents, and 580 Dr Shuai Nie, Yunyang Zhang and Alex Chen (Thermo Fisher) for technical support. Thermo 581 Fisher Scientific Australia are acknowledged for access to the TSQ Altis triple quadrupole 582 mass spectrometer.

583 **References**

584 1. Hu ZY, Zhao, F.J. & McGrath, S.P. 2005. Sulphur fractionation in calcareous soils and 585 bioavailability to plants. Plant Soil 268:103-109. 586 2. Wilhelm Scherer H. 2009. Sulfur in soils. J Plant Nutr Soil Sci 172:326-335. 587 Scherer HW. 2001. Sulphur in crop production — invited paper. Eur J Agronomy 3. 588 14:81-111. 589 4. Tabatabai MA. 1984. Importance of Sulphur in Crop Production. Biogeochemistry 590 1:45-62. 591 5. Kertesz MA, Mirleau P. 2004. The role of soil microbes in plant sulphur nutrition. J 592 Exp Bot 55:1939-45. 593 6. Kertesz MA. 2000. Riding the sulfur cycle - metabolism of sulfonates and sulfate 594 esters in Gram-negative bacteria. FEMS Microbiol Rev 24:135-175. 595 Autry AR, Fitzgerald JW. 1990. Sulfonate S: A major form of forest soil organic sulfur 7. 596 Biol Fertil Soils 10:50-56. 597 Harwood JL, Nicholls RG. 1979. The plant sulpholipid - a major component of the 8. 598 sulphur cycle. Biochem Soc Trans 7:440-447. 599 9. Goss R, Nerlich J, Lepetit B, Schaller S, Vieler A, Wilhelm C. 2009. The lipid 600 dependence of diadinoxanthin de-epoxidation presents new evidence for a 601 macrodomain organization of the diatom thylakoid membrane. J Plant Physiol 602 166:1839-1854. 603 10. Strickland TC, Fitzgerald JW. 1983. Mineralization of sulphur in sulphoquinovose by 604 forest soils. Soil Biol Biochem 15:347-349. 605 11. Goddard-Borger ED, Williams SJ. 2017. Sulfoquinovose in the biosphere: occurrence, 606 metabolism and functions. Biochem J 474:827–849. 607 Denger K, Weiss M, Felux AK, Schneider A, Mayer C, Spiteller D, Huhn T, Cook AM, 12. 608 Schleheck D. 2014. Sulphoglycolysis in Escherichia coli K-12 closes a gap in the 609 biogeochemical sulphur cycle. Nature 507:114-117. 610 13. Felux AK, Spiteller D, Klebensberger J, Schleheck D. 2015. Entner-Doudoroff pathway 611 for sulfoquinovose degradation in Pseudomonas putida SQ1. Proc Natl Acad Sci USA 612 112:E4298-305. 613 14. Denger K, Huhn T, Hollemeyer K, Schleheck D, Cook AM. 2012. Sulfoquinovose 614 degraded by pure cultures of bacteria with release of C₃-organosulfonates: complete 615 degradation in two-member communities. FEMS Microbiol Lett 328:39-45. 616 Drew EA, Ballard RA. 2010. Improving N2 fixation from the plant down: Compatibility 15. 617 of Trifolium subterraneum L. cultivars with soil rhizobia can influence symbiotic 618 performance. Plant Soil 327:261-277. 619 16. Melino VJ, Drew EA, Ballard RA, Reeve WG, Thomson G, White RG, O'Hara GW. 2012. 620 Identifying abnormalities in symbiotic development between Trifolium spp. and 621 Rhizobium leguminosarum bv. trifolii leading to sub-optimal and ineffective nodule 622 phenotypes. Ann Bot 110:1559-72. 623 17. Reeve W, Drew E, Ballard R, Melino V, Tian R, De Meyer S, Brau L, Ninawi M, Teshima 624 H, Goodwin L, Chain P, Liolios K, Pati A, Mavromatis K, Ivanova N, Markowitz V, 625 Woyke T, Kyrpides N. 2013. Genome sequence of the clover-nodulating Rhizobium 626 leguminosarum bv. trifolii strain SRDI565. Stand Genomic Sci 9:220-31. 627 18. Udvardi M, Poole PS. 2013. Transport and metabolism in legume-rhizobia symbioses. 628 Annu Rev Plant Biol 64:781-805.

629 19. Poole P, Ramachandran V, Terpolilli J. 2018. Rhizobia: from saprophytes to 630 endosymbionts. Nat Rev Microbiol 16:291-303. 631 20. Stowers MD. 1985. Carbon metabolism in Rhizobium species. Annu Rev Microbiol 632 39:89-108. 633 Abayakoon P, Lingford JP, Jin Y, Bengt C, Davies GJ, Yao S, Goddard-Borger ED, 21. 634 Williams SJ. 2018. Discovery and characterization of a sulfoquinovose mutarotase 635 using kinetic analysis at equilibrium by exchange spectroscopy. Biochem J 475:1371-636 1383. 637 22. Bergersen FJ. 1961. The growth of rhizobium in synthetic media. Aust J Biol Sci 638 14:349-360. 639 Goldschmidt L, Cooper DR, Derewenda ZS, Eisenberg D. 2007. Toward rational 23. 640 protein crystallization: A Web server for the design of crystallizable protein variants. 641 Protein Sci 16:1569-76. 642 24. Abayakoon P, Jin Y, Lingford JP, Petricevic M, John A, Ryan E, Wai-Ying Mui J, Pires 643 DEV, Ascher DB, Davies GJ, Goddard-Borger ED, Williams SJ. 2018. Structural and 644 Biochemical Insights into the Function and Evolution of Sulfoquinovosidases. ACS 645 Cent Sci 4:1266-1273. 646 25. Speciale G, Jin Y, Davies GJ, Williams SJ, Goddard-Borger ED. 2016. YihQ is a 647 sulfoquinovosidase that cleaves sulfoquinovosyl diacylglyceride sulfolipids. Nat Chem 648 Biol 12:215-217. 649 26. Roy AB, Hewlins MJE. 1997. Sulfoquinovose and its aldonic acid: their preparation 650 and oxidation to 2-sulfoacetaldehyde by periodate. Carbohydr Res 302:113-117. 651 Abayakoon P, Epa R, Petricevic M, Bengt C, Mui JWY, van der Peet PL, Zhang Y, 27. 652 Lingford JP, White JM, Goddard-Borger ED, Williams SJ. 2019. Comprehensive 653 synthesis of substrates, intermediates and products of the sulfoglycolytic Embden-654 Meyerhoff-Parnas pathway. J Org Chem 84:2910-2910. 655 28. Roy AB, Ellis AJ, White GF, Harwood JL. 2000. Microbial degradation of the plant 656 sulpholipid. Biochem Soc Trans 28:781-3. 657 Roy AB, Hewlins MJ, Ellis AJ, Harwood JL, White GF. 2003. Glycolytic breakdown of 29. 658 sulfoquinovose in bacteria: a missing link in the sulfur cycle. Appl Environ Microbiol 659 69:6434-6441. 660 30. Martelli HL, Benson AA. 1964. Sulfocarbohydrate metabolism. I. Bacterial production 661 and utilization of sulfoacetate. Biochim Biophys Acta 93:169-171. 662 31. Davidson AL, Dassa E, Orelle C, Chen J. 2008. Structure, Function, and Evolution of 663 Bacterial ATP-Binding Cassette Systems. Microbiol Mol Biol Rev 72:317-364. 664 32. McKay IA, Glenn AR, Dilworth MJ. 1985. Gluconeogenesis in Rhizobium 665 leguminosarum MNF3841. Microbiology 131:2067-2073. 666 Sasidharan K, Martinez-Vernon AS, Chen J, Fu T, Soyer OS. 2018. A low-cost DIY 33. 667 device for high resolution, continuous measurement of microbial growth dynamics. 668 bioRxiv doi:10.1101/407742:407742. 669 Hughes CS, Moggridge S, Müller T, Sorensen PH, Morin GB, Krijgsveld J. 2019. Single-34. 670 pot, solid-phase-enhanced sample preparation for proteomics experiments. Nat 671 Protoc 14:68-85. 672 35. Rappsilber J, Mann M, Ishihama Y. 2007. Protocol for micro-purification, enrichment, 673 pre-fractionation and storage of peptides for proteomics using StageTips. Nat Protoc 674 2:1896-1906.

- 675 36. Cox J, Mann M. 2008. MaxQuant enables high peptide identification rates,
 676 individualized p.p.b.-range mass accuracies and proteome-wide protein
 677 quantification. Nat Biotechnol 26:1367-1372.
- 678 37. Cox J, Hein MY, Luber CA, Paron I, Nagaraj N, Mann M. 2014. Accurate Proteome679 wide Label-free Quantification by Delayed Normalization and Maximal Peptide Ratio
 680 Extraction, Termed MaxLFQ. Mol Cell Proteomics 13:2513.
- 681 38. Tyanova S, Temu T, Sinitcyn P, Carlson A, Hein MY, Geiger T, Mann M, Cox J. 2016.
- 682 The Perseus computational platform for comprehensive analysis of (prote)omics683 data. Nat Methods 13:731.
- Masukagami Y, Nijagal B, Mahdizadeh S, Tseng CW, Dayalan S, Tivendale KA,
 Markham PF, Browning GF, Sansom FM. 2019. A combined metabolomic and
 bioinformatic approach to investigate the function of transport proteins of the
- 687 important pathogen Mycoplasma bovis. Vet Microbiol 234:8-16.

688

689 Figure 1: Proposed sulfoglycolytic Entner-Doudoroff (sulfo-ED) pathway in *Rhizobium*

690 *leguminosarum* bv. trifolii SRDI565. (a) Operon encoding sulfo-ED pathway. (b) Proposed

- 691 sulfo-ED pathway. (c) Comparison with the Entner-Doudoroff pathway.
- 692

693 Figure 2: Growth of Rhizobium leguminosarum by. trifolii SRDI565 on SQ produces SL 694 as the major terminal metabolite. a) Growth of *Rl*-SRDI565 on 5% yeast extract media 695 containing 5 mM SQ (•) or 5 mM mannitol (•). This data is representative of 2 independent experiments. b) ¹³C NMR (126 MHz) spectra of (top) SQ, (middle) 5 mM SQ in 5% yeast 696 697 extract media and (bottom) spent culture media from growth of *Rl*-SRDI565 on 5 mM SQ. c) 698 ¹³C NMR (126 MHz) spectrum of spent culture media from growth of *Rl*-SRDI565 on 5 mM 699 $(^{13}C_6)$ -SQ. The signal at δ 38.7 ppm is present in control experiments of *Rl*-SRDI565 grown on mannitol and is believed to derive from yeast extract. d) Tabulated ¹³C NMR (126 MHz) 700 data for ¹³C₃-SL from (c). All samples contain 10% D₂O, added to allow frequency lock. e) 701 702 Quantitative proteomics was undertaken to identify proteins associated with sulfoquinovose 703 catabolism versus mannitol. Examination of proteins observed to increase in abundance 704 greater than four-fold revealed 17 proteins including alpha-dehydro-beta-deoxy-D-glucarate 705 aldolase (WP 017967308.1). f) Growth in sulfoquinovose leads to the increase of multiple 706 proteins associated with the TCA cycle including NAD(P)-dependent oxidoreductase 707 (WP_017965793.1), NADH-quinone oxidoreductase subunit NuoH (WP_017963854.1), 708 NAD-dependent succinate-semialdehyde dehydrogenase (WP_017967313.1) and citrate 709 synthase/methylcitrate synthase (WP_017964386.1) highlighted in blue.

710

711 Figure 3: *Rhizobium leguminosarum* SRDI565 produces a functional sulfoquinovosidase

that can be detected in cell lysates. a) pH profile of *Rl*SQase*. Specific activities were
determined for hydrolysis of PNPSQ at the isosbestic point, 348 nm. b) Michaelis Menten

plot of kinetic parameters for *Rl*SQase* for hydrolysis of PNPSQ at 400 nm. c) Analysis of
sulfoquinovosidase activity of *Rl*-SRDI565 lysate grown on sulfoquinovose and mannitol.
Cell lysates of soluble proteins derived from growth on SQ or mannitol was standardized for
equal protein and SQase activity measured using the chromogenic substrate PNPSQ at 400
nm. SQase activity was confirmed by inhibition by the azasugar inhibitor SGIFG. Error bars
denote standard error of the mean.

720

721 Figure 4: Detection of sulfoglycolytic intermediates and end-products in cytosolic 722 extracts of *RI*-SRDI565. RI-SRDI565 was grown on Y_{5%}SQ media and metabolically-723 quenched by rapid cooling to 4 C, followed by extraction of cellular metabolites and lc/ms 724 analysis. Detection of sulfoglycolytic and glycolytic/neoglucogenic intermediates A) SQ, B) 725 SG, C) SL, D) SF, E) DHPS. In each case the upper panel corresponds to the collision-726 induced dissociation mass spectrum of chemically-synthesized standard, while the lower 727 panel is the equivalent mass spectrum for the metabolite identified in the cytosolic extract. F) 728 Relative mass spectrometric intensities of metabolites from cells grown on Glc or SQ.

729

Figure 5: Proposed pathway for SQ metabolism in *Rhizobium leguminosarum* SRDI565.

Table 1. Analysis of sulfonate metabolites detected in spent culture media of *Rl*-SRDI565
grown on 5.0±0.5 mM SQ or SQGro (standard error estimate). Measurements were
performed in triplicate using LC/MS-MS. Errors listed in the table are standard error mean.







Figure 1



-			
	¹³ C chemical shift (δ ppm)	coupling (Hz)	assignment
	53.2	${}^{1}J_{\rm C1,C2} = 37.1$	C3
	64.7	${}^{1}J_{C2,C3} = 54.8$ ${}^{1}J_{C1,C2} = 37.1$	C2
	178.6	${}^{1}J_{C2,C3} = 54.8$	C1





Figure 3







259.0124

241.0015

Relative intensity (millions)

Figure 4



Figure 5