

1 Isolation of sophorose during sophorolipid production and 2 studies of its stability in aqueous alkali: epimerisation of 3 sophorose to 2-O- β -D-glucopyranosyl-D-mannose 4

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33 **Abstract**

34 NMR and anion exchange chromatography analysis of the waste streams generated during the
35 commercial production of sophorolipids by the yeast *Candida bombicola* identified the presence of
36 small but significant quantities (1 % w/v) of free sophorose. Sophorose, a valuable disaccharide,
37 was isolated from the aqueous wastes using a simple extraction procedure and was purified by
38 chromatography on a carbon celite column providing easy access to large quantities of the
39 disaccharide. Experiments were undertaken to identify the origin of sophorose and it is likely that
40 acetylated sophorose derivatives were produced by an enzyme catalysed hydrolysis of the
41 glucosyl-lipid bond of sophorolipids; the acetylated sophorose derivatives then undergo hydrolysis
42 to release the parent disaccharide.

43 Treatment of sophorose with aqueous alkali at elevated temperatures (0.1M NaOH at 50 °C)
44 resulted in C2-epimerisation of the terminal reducing sugar and its conversion to the corresponding
45 2-O- β -D-glucopyranosyl-D-mannose which was isolated and characterised. In aqueous alkaline
46 solution β -(1,2)-linked glycosidic bonds do not undergo either hydrolysis or peeling reactions.

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49 Key words:

50 Sophorose, Sophorolipids, β -(1,2)-linked disaccharides, 2-O- β -D-glucopyranosyl-D-mannose

51

52 1. Introduction

53 The disaccharide sophorose, 2-O- β -D-glucopyranosyl-D-glucose, is an important biologically active
54 molecule. As a free disaccharide, sophorose is a potent inducer of cellulases¹ and is used to
55 generate enzymes for use in bioethanol production. Sophorose is a structural component of the
56 biologically active glycosides saponins², including ginsenosides³. Sophorose is also the
57 carbohydrate component of sophorolipids⁴⁻⁶ which are increasingly being manufactured on a large
58 scale for use as surface active agents.

59 The synthesis and derivatisation of sophorose was first reported in the 1920s by Freudenberg *et*
60 *al*⁷ and subsequently the methods were improved in the 1960s by Coxon and Fletcher⁸. Interest in
61 sophorose grew in the 1950s when it was found to be present in trace amounts in commercial
62 supplies of D-glucose that had been obtained by the mineral acid catalysed hydrolysis of starch⁹
63 and this led to increased efforts to find a more convenient route to sophorose. In the 1950 and
64 1960s a number of authors reported the isolation of sophorose from naturally occurring sophoryl-
65 glycosides including kaempferol sophoroside isolated from pods of *Sophora japonica*¹⁰ and from
66 the sweet ester glycoside stevioside from *Stevia rebaudiana*¹¹. More recently sophorose and
67 activated sophorose donors have been produced from sophorolipids. Hoffmann *et al*¹² have
68 reported the isolation of per-O-acetyl-sophorosyl bromide from the reaction of acetylated
69 sophorolipids with acetic acid and hydrogen bromide. Jourdir and Ben¹³ have reported the acid
70 catalysed hydrolysis of sophorolipids generating a mixture of glucolipids, glucose and sophorose
71 from which, after enzyme treatment to remove glucose, they were able to isolate sophorose.

72 Despite the relatively wide interest in sophorose there is only limited information about its chemical
73 reactivity and that of 1,2-linked glycosides in general; this is partially due to the fact that they are
74 not readily accessible. In this paper we report the isolation of sophorose from a waste stream
75 generated during the commercial production of sophorolipids, provide evidence for its synthesis by
76 a combination of enzyme and base catalysed hydrolysis reactions and we describe the results of
77 our studies of the reaction of sophorose with aqueous alkali.

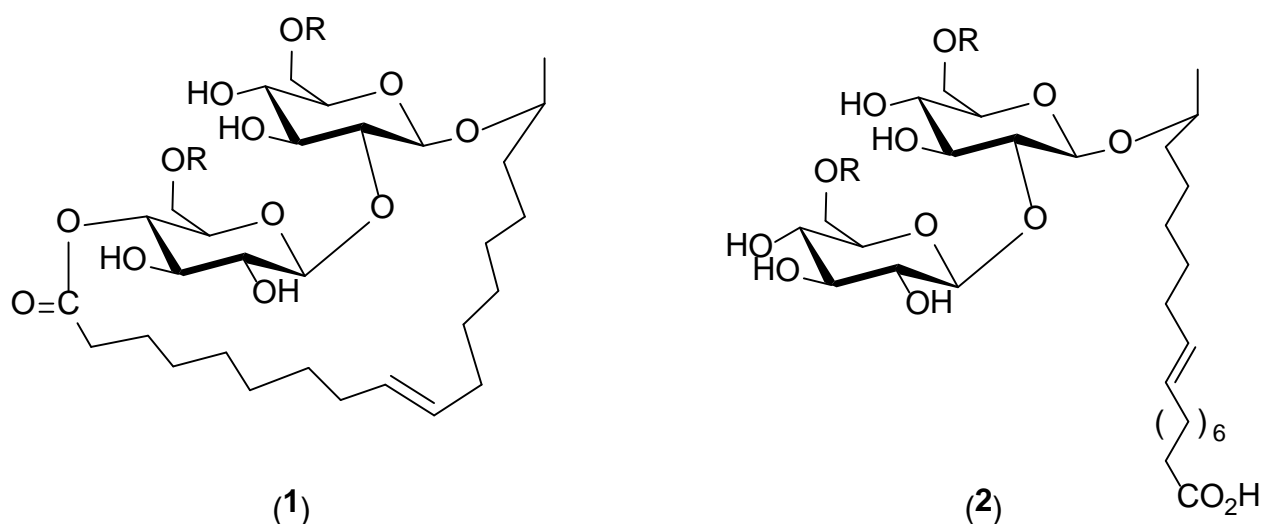
78 **2. Results and Discussion.**

79 **2.1 Isolation of sophorose from sophorolipid process waste streams.**

80 The yeast *Candida bombicola* produces sophorolipids in a biosynthetic pathway^{14,15} involving the
81 hydroxylation of a fatty acid (normally C18) to generate a (ω -1)-hydroxyfatty acid which is then
82 glucosylated twice, initially at the (ω -1)-hydroxyl-group and then at the 2'-position of the newly
83 added glucose residue to generate a sophorolipid. The pathway is concluded with the partial
84 acetylation of the sugar primary hydroxyls (6' and 6'') and by partial lactonisation of the 4''-OH with
85 the lipid fatty acid to give a mixture of lactone (**1**, **R=H or OAc**) and acid (**2**, **R=H or OAc**) forms of
86 the sophorolipids (Scheme 1).

87 Scheme 1 here.

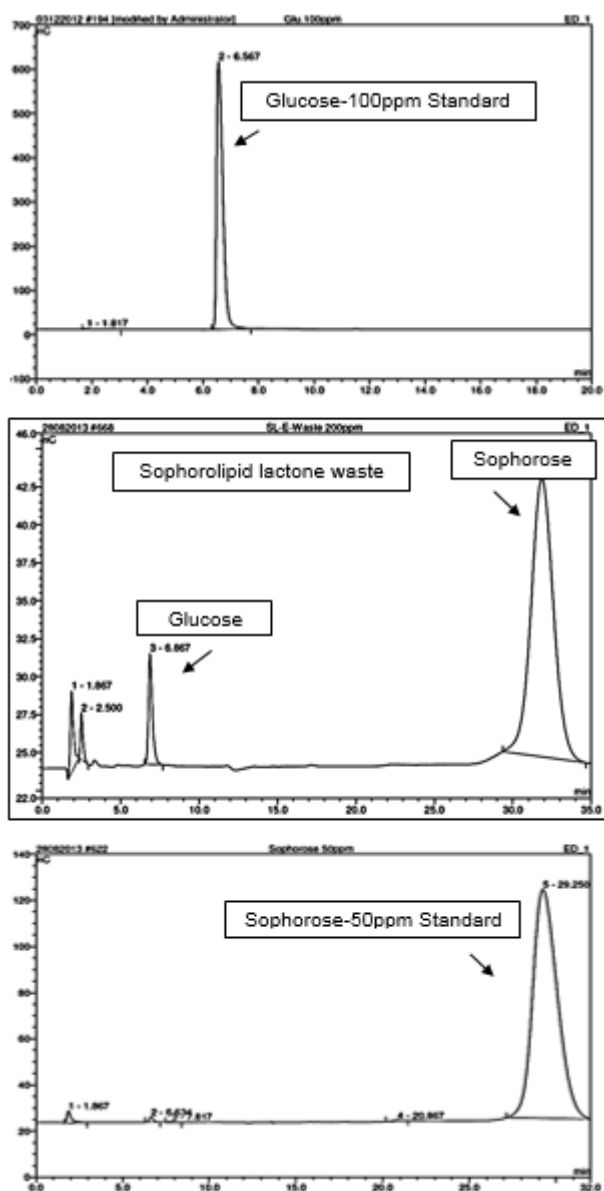
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90 During a typical fermentation, sophorolipid production is allowed to proceed until the pH of the
91 fermentation mixture falls to approximately 3.5 and the mixture settles into two phases: a lipid rich
92 bottom layer containing the sophorolipids¹⁶ and an upper aqueous phase (spent liquors). To isolate
93 the deacetylated acid form of the sophorolipid (**2 R=H**) it is normal practice to recover the pure
94 lactone and then to convert it to the acid. The lactone is recovered from the lipid rich layer by
95 extraction of acidic impurities by adjustment of the pH to above the pKa of the acid group using
96 cold aqueous alkali (pH 5.5). After separation of the layers, the lipid rich layer contains
97 predominately the lactone form and the aqueous phase (lactone waste) which is brown in colour
98 contains a number of impurities. To isolate the acid form the lactone is converted into the acid in a

99 hydrolysis step involving heating with aqueous alkali¹⁷. In an attempt to determine the composition
100 of the different phases and to understand the extraction and hydrolysis reaction the different
101 phases were analysed by NMR and HPAEC. HPAEC analysis of the lactone waste identified the
102 presence of free sugars, interestingly the HPAEC chromatograph (Fig 1) contained peaks
103 consistent with the presence of both glucose and sophorose (**3**) (ratio 1: 10 w/w). NMR analysis
104 confirmed the presence of sugars and acid sophorolipids.



105
106 **Figure 1:** High Performance Anion Exchange Chromatographs (HPAEC) chromatographs recorded
107 using a Dionex ICS3000 and employing a CarboPac PA20 column. Analytes (50-200 ppm) were
108 eluted with a mobile phase composed of 20 mM KOH with a flow rate of 0.3 ml.min⁻¹. Top: A 100
109 ppm glucose standard; Middle: A sample of the lactone waste; Bottom: A 50 ppm sophorose
110 standard.

112 As the commercial process generates large volumes of lactone waste (1000s of litres) an attempt
113 was made to isolate the sophorose. Initially, the waste (98g, 100mL) was diluted with an equal
114 volume of ultrapure water and the sugars were extracted into the aqueous phase leaving the lipid
115 components as a separate organic phase. After separation of the two layers, the aqueous phase
116 was evaporated to leave a brown solid residue which contained a range of salts as well as the
117 desired sugars (5.5g, 5.6% w/w of the original waste). NMR analysis of the solid (Fig 2a) confirmed
118 that the organic component of the material was a mixture of glucose and sophorose and that no
119 lipids were present. Comparison of the integrals for the anomeric protons (α plus β -anomers) for
120 the two sugars indicated that the extracted mixture contained a 1: 2.3 (w/w) ratio of glucose to
121 sophorose and HPAEC analysis suggested a slightly higher ratio of 1:2.9 (w/w). The solid was
122 dissolved in ultra pure water and applied to a carbon-celite column and the column was eluted
123 initially with water to remove salts and then with an aqueous mobile phase containing increasing
124 concentrations of ethanol. Sophorose eluted as a single peak when the eluent was composed of
125 20% ethanol; after evaporation sophorose was recovered as a crystalline white solid (1.0g). NMR
126 analysis (Fig 2b) indicated that sophorose with a purity of greater than 95% had been recovered
127 and this corresponds to approximately one percent of the original waste. Given the large volume of
128 lactone waste available this represents a potentially large source of sophorose.

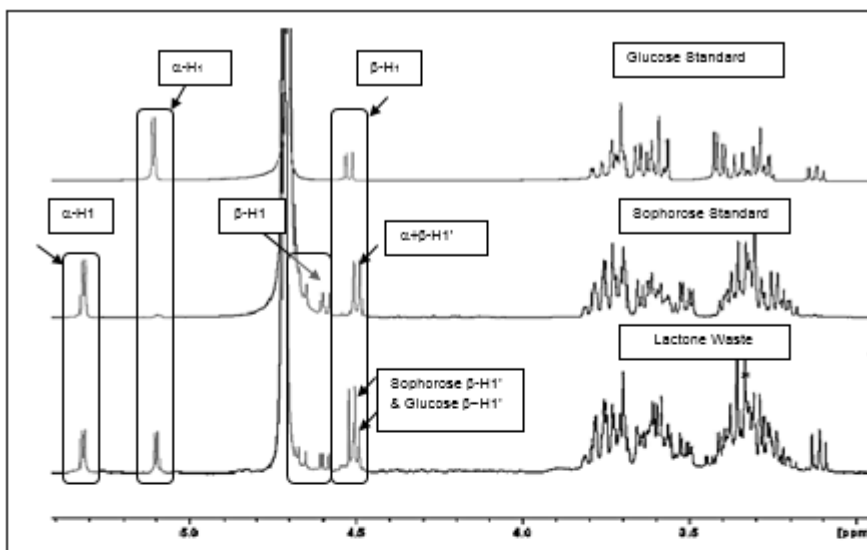
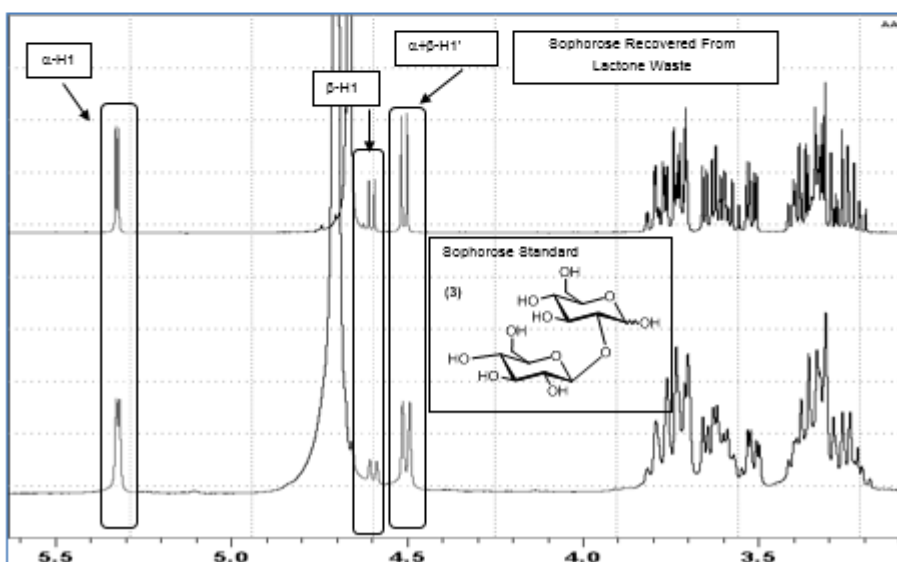


Figure 2b



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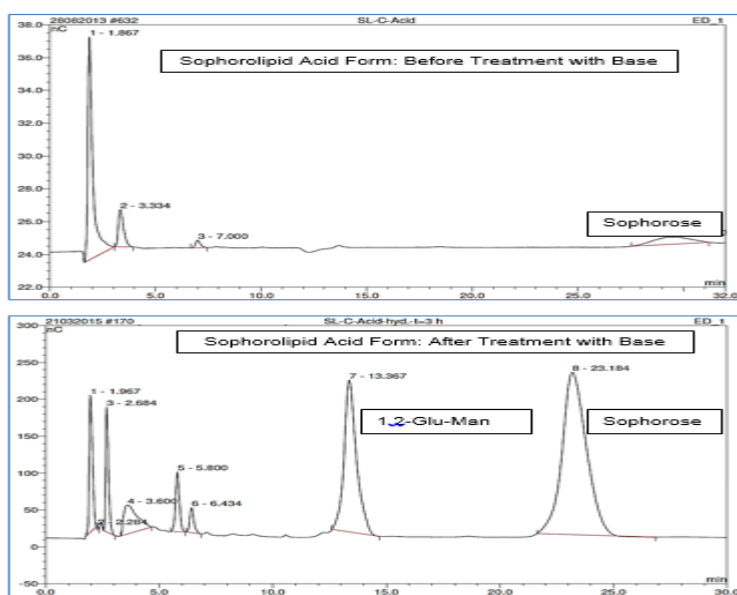
130 **Figure 2:** $^1\text{H-NMR}$ spectra recorded at $70\text{ }^\circ\text{C}$ in D_2O on a Bruker 500MHz Avance Spectrometer (a)
 131 comparison of NMRS of a glucose standard, a sophorose standard and a sample of the lactone
 132 waste: a) top: glucose standard; middle: sophorose standard; bottom: sugar sample extracted from
 133 lactone waste (b) top: NMR of the sophorose isolated from a carbon-celite column; bottom: NMR of
 134 a sophorose standard.

135 As the biosynthetic pathway involves sequential addition of glucose residues to the lipid chain¹⁵,
 136 free sophorose was not expected to be a product of the fermentation process.

137 2.2 Attempt to observe base catalysed hydrolysis of sophorolipids.

138 In an attempt to determine if sophorose was being produced by a base catalysed hydrolysis of the
 139 sophorolipid during the processing of the crude fermentation products, an experiment was
 140 undertaken to see if sophorose was released from the acid form of the sophorolipid on heating with

141 aqueous alkali. The acid form of the sophorolipid was chosen because of its solubility in aqueous
142 alkali and because of the absence of acetyl-ester groups, the hydrolysis of which would consume
143 base. The reaction of the acid sophorolipid with aqueous alkali (pH 12, 50 °C) was monitored by
144 HPAEC and NMR over a period of 24 h. No reaction was observed by NMR, however, analysis of
145 the HPAEC chromatographs (Fig 3) identified the rapid production of very small quantities of
146 glucose, small quantities of sophorose and a second disaccharide which was identified as the C2-
147 epimer 2-O- β -D-glucopyranosyl-D-mannose (see Section 2.4). These additional sugars were
148 produced in the first three hours after which time no further reaction was observed and the
149 combined peak area of the new sugars suggested that less than 0.5% of the starting material had
150 been converted to sophorose. Under the relatively mild alkaline conditions employed, sophorolipids
151 were not expected to undergo glycosidic bond hydrolysis or to participate in either substitution or
152 elimination processes centred at the lipid ω -1 carbon; it is more likely that the very small amount of
153 sophorose that is produced is generated through the rapid hydrolysis of small quantities of the
154 corresponding acetylated-sugars that are present as minor impurities in the starting acid. This
155 result suggests that under moderately basic conditions hydrolysis of sophorolipids does not lead to
156 the production of sophorose.



157 **Figure 3:** High performance
158 anion exchange chromatographs recorded using a Carbobac PA20 column and eluting analytes (50-
159 200 ppm) with a mobile phase composed of 20 mM KOH with a flow rate of 0.3 ml.min⁻¹. Top: Acid
160 form of the sophorolipid; bottom: reaction mixture obtained after treatment of the acid sophorolipid
161 with aqueous alkali (10 mM, 3 h, 50 °C).

162 2.3 Observation of an enzyme catalysed hydrolysis of sophorolipids. One of the reasons why the
 163 yeast *C. bombicola* is thought to synthesise sophorolipids is to allow it to sequester available fatty
 164 acids. It has previously been suggested that under conditions of starvation the yeast is able to
 165 metabolise sophorolipids¹⁴. In order to investigate the possibility that the yeast releases an enzyme
 166 which is able to catalyse the hydrolysis of the gluco-lipid bond of the acid or lactone forms of the
 167 sophorolipids, two experiments were undertaken in which the spent liquors recovered at the end of
 168 a large scale fermentation were filtered to remove any biomass and then immediately added to a
 169 solution containing either the diacetylated-lactone sophorolipid (**1**, **R=Ac**) or the deacetylated acid
 170 form (**2**, **R=H**) of the sophorolipid.

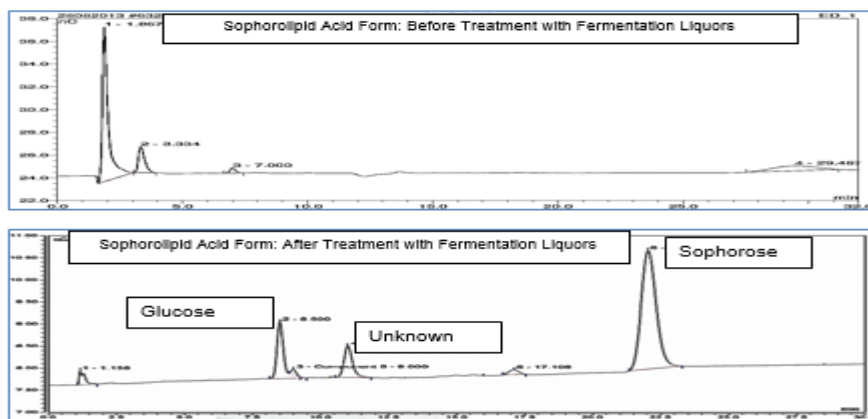
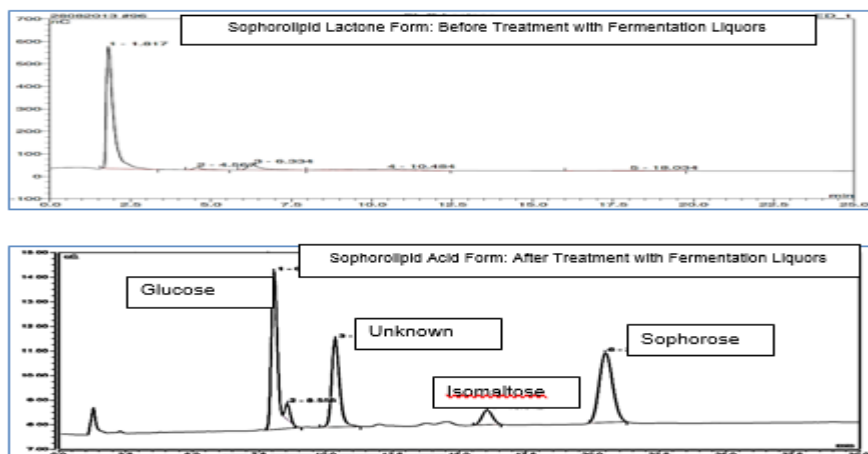


Figure 4b



171

172 **Figure 4:** Use of high performance anion exchange chromatography to monitor the enzyme
 173 catalysed hydrolysis of sophorolipids¹

174 (a) Acid form of the sophorolipid; top: before addition of fermentation liquor and bottom: after
 175 treatment with spent fermentation liquor

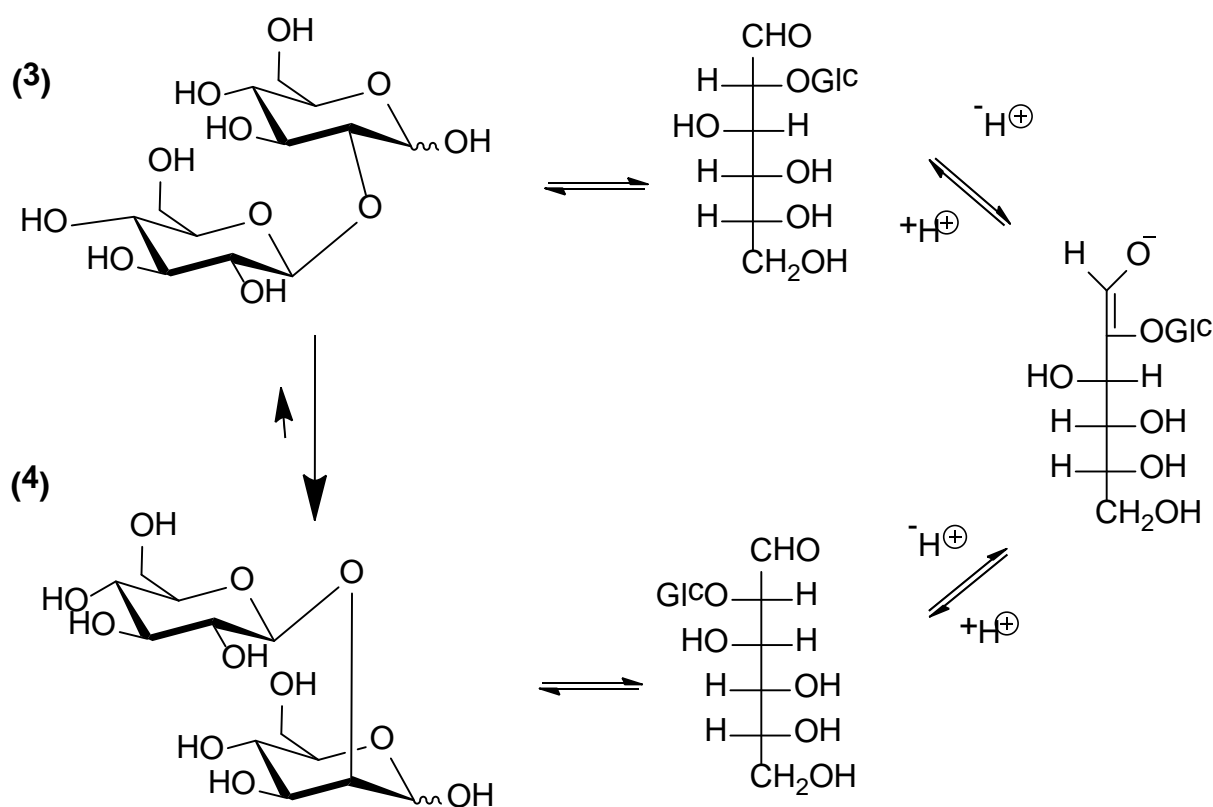
176 (b) Lactone form of the sophorolipid; top: before addition of fermentation liquor and bottom: after
 177 treatment with spent fermentation liquors.

178

179 Chromatographs generated during HPAEC analysis of the products (Fig 4a Acid after treatment
180 and 4b Lactone after treatment) showed the presence of sophorose and glucose along with a small
181 amount of isomaltose and an unknown product (potentially glucolipid). The amount of sophorose
182 was significantly higher when the acid form of the sophorolipid was used as a substrate. The
183 results are consistent with an enzyme catalysed hydrolysis of the sophorolipids with the acid form
184 having generated sophorose, in comparison, the product formed from the lactone would be
185 expected to be diacetylated-sophorose. The latter result, whilst not being conclusive evidence, is
186 consistent with the presence of enzyme/s that are able to cleave the 1,2-glycosidic bond and the
187 lipid-glycosidic bond. An enzyme catalysed cleavage of the 1,2-glycosidic link would, in the first
188 instance, generate a gluco-lipid and a glucosyl-enzyme intermediate. Attack by water on the
189 glucosyl-enzyme intermediate would generate glucose and release the enzyme. Attack by a
190 primary hydroxyl of a second glucose molecule on the glucosyl-enzyme intermediate would lead to
191 production of isomaltose and release the enzyme. Production of isomaltose requires the
192 transferase activity of an inverting enzyme.

193 *2.4 Studies of the chemical reactivity of sophorose and isolation of 2-O-β-D-glucopyranosyl-D-*
194 *mannose.* Very little is known about the chemical reactivity of 1,2-linked glycosides. Treatment of
195 1-3, 1-4 and 1-6-linked glycosides with aqueous alkali results in the observation of peeling
196 reactions and the generation of multiple small chain fatty acids including saccharinic acids^{18,19}. As
197 peeling proceeds through enolisation and migration of the carbonyl carbon to C2 and beyond (a
198 process that is not possible in C2-linked glycosides) it was of interest to study the stability of
199 sophorose in aqueous alkali. After treatment of sophorose with sodium hydroxide solution using
200 conditions which would promote peeling of glycosides, the HPAEC chromatographs contained
201 signals identifying the presence of two carbohydrates: a late eluting peak for sophorose and a
202 second disaccharide peak. NMR analysis of the reaction mixture indicated that deuterium
203 exchange was occurring at the C2 position of the starting material and, as was the case in the
204 HPAEC analysis, a second disaccharide was present. These results were consistent with the
205 epimerisation of the reducing sugar to form the corresponding 2-O-β-D-glucopyranosyl-D-mannose
206 (4, Scheme 2).

207



209

210 The partial epimerisation of sophorose has already been reported but in this earlier study the

211 product 2-*O*- β -D-glucopyranosyl-D-mannose was not isolated and was not fully characterised²⁰. The

212 NMR spectra recorded here suggested that under the conditions of the current experiments 75% of

213 the starting sophorose was converted to 2-*O*- β -D-glucopyranosyl-D-mannose (4). The mechanism

214 for such epimerisation reactions was first proposed by Lobry de Bruyn and Alberda van

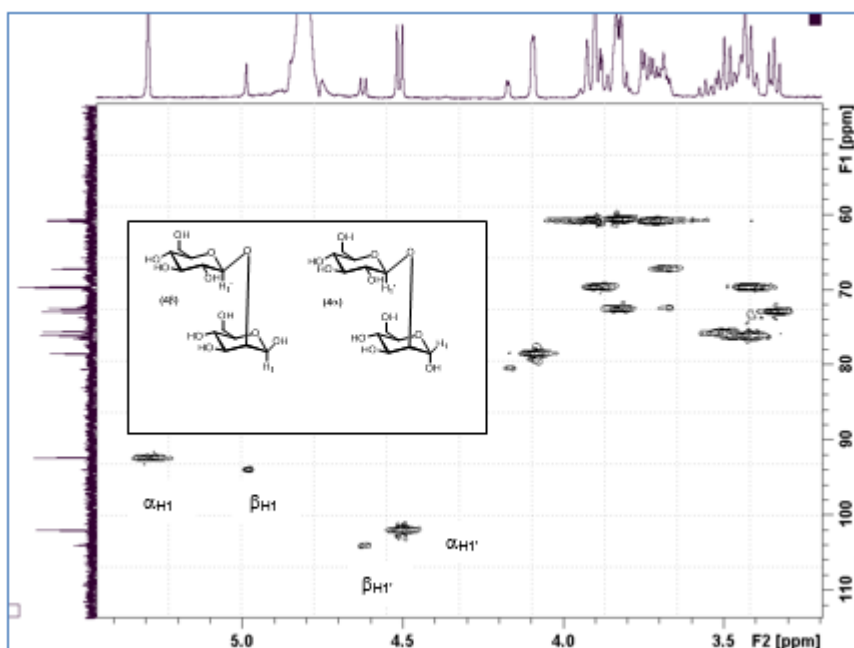
215 Ekenstein²¹ and involves ring-opening of the reducing glucose and base catalysed reversible

216 enolate formation between C2 and C1. Reprotonation of the enolate anion on C2 can take place on

217 either the top Re-face to regenerate glucose or from the bottom Si-face to generate mannose. At

218 equilibrium, and under the conditions employed in the current experiments, the thermodynamically

219 more favourable mannose epimer dominates (see scheme 2).



220

221 **Figure 5.** Selected region of the HSQC NMR spectrum of 2-O- β -D-glucopyranosyl- α/β -D-mannose.
 222 The F1-axis provides the ^{13}C -spectrum and the F2-axis the ^1H -spectrum. The location of the
 223 individual resonances for the different protons and carbons are listed in the table. The spectrum was
 224 recorded at 70 °C in D_2O .

225

226 In an attempt to isolate the 2-O- β -D-glucopyranosyl-D-mannose a small quantity of sophorose was
 227 treated with aqueous alkali and after HPAEC analysis confirmed that the majority of the starting
 228 material had been converted to 2-O- β -D-glucopyranosyl-D-mannose the reaction was quenched by
 229 acidification with acetic acid. The crude products were then applied to a carbon-celite column and
 230 the new disaccharide was isolated by elution with aqueous ethanol. The product eluted as a single
 231 peak in 20% ethanol and, after evaporation of the solvent, 2-O- β -D-glucopyranosyl-D-mannose was
 232 recovered as a white powder. The proton NMR (Fig 5 F2 (x)-axis) of an aqueous solution of 2-O- β -
 233 D-glucopyranosyl-D-mannose indicated that the product was present as a pair of anomers. At 30 °C
 234 and in an aqueous solution of D_2O 2-O- β -D-glucopyranosyl-D-mannose exists as a mixture
 235 containing 77% of the alpha-anomer and 23% of the beta-anomer. A number of 1D and 2D-NMRs
 236 were recorded (^1H , ^{13}C and HSQC spectra are shown in Fig 5.) and a full assignment for the proton
 237 and carbons of the alpha anomer and for the protons of the beta-anomer (for the beta-anomer
 238 carbon signals were weak and only the anomeric signals were clearly visible above the spectral
 239 noise); the chemical shifts for both the protons and carbons are listed in the table. It should be

240 noted that the proton assignment for the anomeric glucose (H1')-protons are different to those in
 241 the literature²⁰ (the values in the literature for the H1' proton for alpha and beta have been
 242 inverted). Even after prolonged exposure to high pHs there was no evidence for hydrolysis of the
 243 glycosidic bond and there was no evidence for decomposition via peeling reactions.
 244 In summary, the valuable β -(1,2)-linked disaccharide sophorose was isolated from waste streams
 245 generated during the commercial manufacture of sophorolipids and we have provided evidence
 246 which indicates that sophorose is generated by an enzyme catalysed hydrolysis of the parent
 247 sophorolipid. When treated with aqueous alkali, sophorose did not undergo either hydrolysis or a
 248 peeling reaction and instead underwent epimerisation at C2 to generate 2-O- β -D-glucopyranosyl-D-
 249 mannose which was isolated and characterised by NMR.

Sugar/Position	H1	H2	H3	H4	H5	H6s	C1	C2	C3	C4	C5	C6s
Reducing	5.29	4.17	3.88	3.81	3.69	3.93 & 3.72	92.3	78.6	69.6	72.4	67.2	60.7
Non-reducing	5.51	3.34	3.49	3.43	3.41	3.83	101.9	72.9	75.6	76.0	69.6	60.4

β -anomer

Sugar/Position	H1	H2	H3	H4	H5	H6s	C1
Reducing	4.98	4.09	3.68	3.55	3.41	ND	93.9
Non-reducing	4.62	3.41	3.48	ND	ND	ND	104.1

250

251 **Table:** Individual chemical shifts for protons and carbons for 2-O- β -D-glucopyranosyl- α -D-mannose
 252 (top) and specific resonances for 2-O- β -D-glucopyranosyl- α -D-mannose (bottom) (recorded in D₂O
 253 at 70 °C on a Bruker Avance 500 MHz spectrometer).

254

255 **Experimental**

256 **3.1 Materials and Chemicals**

257 Monosaccharides and disaccharide standards were purchased from Carbosynth (Berkshire, UK).
 258 All other reagents were purchased from Aldrich (Poole, UK) unless otherwise stated and were
 259 used as supplied.

260 **3.1.1 Sophorolipid and sophorolipid process wastes.** Samples of sophorolipids and sophorolipid
 261 process wastes were provided by Croda Biotechnology Group (Widnes, UK) and were generated
 262 from a large scale fermentation employing an appropriate lipid and glucose as feeds.
 263 Fermentations (>10,000 L) were initiated by inoculation of the media with the yeast strain *Candida*

264 *bombicola* and at the end of the fermentation the fermentation broth was allowed to settle
265 generating two phases: a lipid-rich phase containing sophorolipids and an aqueous phase.
266 Adjustment of the pH of the sophorolipid layer by addition of aqueous alkali to 5.5 provided a
267 sophorolipid lactone and generated an aqueous phase containing sophorose and which will be
268 subsequently referred to as 'lactone-waste'. Pure samples of the lactone form (1) and the acid form
269 of the sophorolipids (2) were also supplied by Croda.

270 3.2 General Analytical Procedures.

271 Analysis of monosaccharides and disaccharides was performed using high pressure anion
272 exchange chromatography coupled to a pulsed amperometric detector (HPAEC-PAD). Standards
273 and unknowns were separated on a Dionex ICS3000 HPAEC system incorporating a Carbowac PA
274 20 column operating at 30 °C and using a mobile phase containing 20mM KOH running at a flow
275 rate of 0.3 ml.min⁻¹. Standards of glucose, isomaltose and sophorose were prepared in ultra-pure
276 water (50-200 ppm).

277 NMR analyses of sophorolipids, sophorose, isomaltose and 2-O-β-D-glucopyranosyl-D-mannose
278 were performed on a Bruker DPX500 NMR spectrometer, a series of 1D and 2D-NMR spectra
279 were recorded in D₂O at a probe temperature of 70 °C and employing acetone as either an internal
280 or external standard. The 2D-spectra recorded included homonuclear ¹H-¹H-COSY and TOCSY
281 (80 ms mixing time) and heteronuclear ¹H-¹³C HSQC, ¹H-¹³C- HMBC, ¹H-¹³C- HSQC-TOCSY and
282 finally a NOESY spectrum was recorded for 2-O-β-D-glucopyranosyl-α/β-D-mannose .

283 3.3 Isolation of disaccharides.

284 3.3.1. Isolation of sophorose (2-O-β-D-glucopyranosyl-α/β-D-glucose). Lactone waste (100 ml, 98
285 g) was added to ultra-pure water (100 ml) and was left to stir at 35-40 °C for 1h. The resulting
286 brown suspension was centrifuged (4200 rpm for 10 mins) and the supernatant was collected.
287 After completion of the separation process, the aqueous phase was rotary evaporated at reduced
288 pressure (10 mmHg) and at a temperature of 45°C to give a crude mixture of salts and sugars as a
289 brown syrup (5.5g). The salt and sugar mix (5.5 g) was redissolved in UPW (3 ml) and the sample
290 was sonicated for 5 mins before being applied to the top a carbon-celite column (5 x 50 cm

291 prepared from Darco G 60, 100 mesh carbon, 200 g, and Celite 535, 200 g). Elution of sugars was
292 carried out with an increasing concentration of aqueous ethanol (from 0 to 20% ethanol in UPW in
293 5% incremental steps of 2 L), HPAEC analysis of the different fractions identified salt plus glucose
294 in the fractions eluting with 5% ethanol (4.0 g) isomaltose in the fractions eluting with 10% ethanol
295 (0.5g) and sophorose from the fractions eluting with 20% ethanol (1.0g).

296 *3.3.2 Isolation of 2-O- β -D-glucopyranosyl- α / β -D-mannose.*

297 Sophorose (50 mg) was treated with aqueous sodium hydroxide (100 mM, 20 ml
298 at pH 13) in a pressure tube and was held at 50 °C for 24 h. The progress of the reaction was
299 monitored using HPAEC; samples (1 ml) were removed every 3 h and the pH of the sample was
300 adjusted to 6 using glacial acetic acid. Samples were then transferred to small glass vials and
301 injected into the HPAEC using sodium hydroxide (50 mM) as the mobile phase. HPAEC analysis
302 suggested that the reaction had reached equilibrium after 24 h at which point the remaining
303 solution was neutralised with glacial acetic acid and the solvent removed under vacuum to give a
304 crude mixture of sugars and sodium acetate as a yellow solid. The sample was redissolved in
305 ultrapure water (1 ml) and was applied to the top of a carbon-celite column (2.5 x 20 cm). Elution of
306 sugars was carried out with an increasing concentration of aqueous ethanol (from 0 to 20% ethanol
307 in UPW in 5% incremental steps each of 200 ml). The desired product eluted with 20% aqueous
308 ethanol and the combined fractions were rotary evaporated to give a pure sample of 2-O- β -D-
309 glucopyranosyl- α / β -D-mannose as a white solid (10 mg, 40% overall yield based on the volume of
310 solution remaining at the end of the reaction). The identity and anomeric purity of the product was
311 determined by 1D-and 2D-NMR (see Section 2).

312

313 *3.4 Monitoring the base catalysed hydrolysis of sophorolipids*

314 Acid sophorolipid (200 mg, 0.322 mM) was dissolved in aqueous sodium hydroxide
315 (10 mM, 20 ml at pH 12) and stirred in a pressure tube at a temperature of 50 °C for 270 min.
316 Samples (1 ml) were taken at 30 min intervals and the pH of the samples was adjusted to pH 6

317 using glacial acetic acid. Samples were then transferred into small glass vials and injected into the
318 HPAEC.

319

320 *3.5 Monitoring the stability of sophorose in aqueous alkaline solution.*

321 Sophorose (5 mg, 0.015 mM) was treated with aqueous sodium hydroxide (100 mM, 20 ml at pH
322 13) in a pressure tube at room temperature for 32 h. Samples (1 ml) were taken out every 1 h and
323 the pH of the sample was adjusted to 6 using glacial acetic acid. Samples were then transferred
324 into small glass vials and injected into the HPAEC using sodium hydroxide (50 mM) as the mobile
325 phase.

326 The same experiment was carried out on sophorose (8 mg), but in an alkaline solution of NaOD
327 (0.1M, 600 μ L in D₂O) at room temperature. The progress of the reaction was followed by running
328 ¹H-NMR spectra after 1, 12 and 24 h.

329

330 *3.6. Hydrolysis of sophorolipids by spent fermentation broth*

331 A sample of a fermentation broth (100 ml) recovered at the end of the fermentation process was
332 supplied by Croda Biotechnology and was stored in a cold-room for 24 h before being used to test
333 for the presence of enzyme activity that may hydrolyse sophorolipids (pH 3.5). In separate
334 experiments, both lactone and acid sophorolipids (0.5 g) were added to a 250 ml clean and
335 autoclaved conical flask and a third flask without any sophorolipid was used as a control. The
336 newly prepared mixtures were kept in a shake-flask incubator. The incubator conditions were set
337 as follows; 30°C, under agitation at 120 rev/min for 3 days (Stuart Orbital Incubator, SI500). After
338 24 h intervals, samples (1 ml) were taken and filtered through (0.45 μ l disk filter) and the
339 supernatant was diluted with UPW (2 ml) then transferred into small vials. The sugar composition
340 of the flasks was assayed by HPAEC-PAD.

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