A novel galacturonide from Xanthomonas campestris

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Enzyme preparations from Xanthomonas campestris incubated in the presence of UDP-[14C]GlcA and Mg2+ produced a lipophilic galacturonide with unusual properties. It was easily degraded by both mild acid treatment (0.01 M-HCl, 100 °C, 10 min) and mild alkali treatment (0.06 M-NaOH, room temperature, 5 min) releasing free [14C]galacturonic acid. The galacturonide appeared to be a single compound with one negative charge, as judged by TLC, paper electrophoresis and chromatography, LH-20 gel filtration and DEAE-cellulose column chromatography. Competition experiments indicated that the true glycosyl donor was UDP-GalA, in agreement with the detection of UDP-GlcA-4-epimerase activity in the crude enzyme preparation. The transglycosidase activity was located mainly in the membrane fraction. UDP inhibited the reaction and even produced some loss of label, suggesting an easily reversible reaction. UMP had almost no effect.

Introduction

Xanthomonas campestris, a plant pathogen, produces an exopolysaccharide, xanthan gum, of extensive industrial application (Sandford, 1979; Sutherland, 1985). Work in this laboratory has shown that, in vitro, the repeating unit of this polysaccharide, which contains glucose, mannose and galacturonic acid, is sequentially assembled on a prenyl diphosphate derivative and subsequently polymerized into xanthan gum (Ielpi et al., 1981). The sugar donors are UDP-Glc, GDP-Man and UDP-GlcA. These studies involved the characterization of a series of radiolabelled prenyl phosphosugars. It was observed that when the label was in UDP-[14C]GlcA, a highly lipophilic compound was formed that did not participate in the biosynthesis of the polysaccharide.

In this paper, some properties of this 'lipid-bound uronic acid' derivative and its synthesizing system are described. The word 'lipid' simply means that the compound is soluble in organic but not in aqueous solvents.

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Abbreviations: GlcA, glucuronic acid; GalA, galacturonic acid; ManA, mannuronic acid; UDP-Glc, uridine diphosphate glucose; GDP-Man, guanosine diphosphate mannose; UDP-GlcA, uridine diphosphate glucuronic acid; UDP-GalA, uridine diphosphate galacturonic acid; EDC, 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide.

Methods

Chemicals. UDP-[14C]glucuronic acid [285–329 mCi mmol-1 (10.5–12.2 GBq mmol-1)] was prepared as previously described (Couso et al., 1982). A mixture of UDP-[14C]glucuronic acid (75%) and UDP-[14C]galacturonic acid (25%) [200 mCi mmol-1 (74 GBq mmol-1)] was prepared with enzymes from Xanthomonas campestris or Rhizobium melliloti 131 (M. Bravo and M. A. Dankert, unpublished results) and is referred to as UDP-[14C]GlcA-GalA. [14C]Galactose-P-prenol and [14C]Glucose-P-P-prenol were synthesized with enzymes from Acetobacter xylinum as reported previously (Romero et al., 1977; Garcia et al., 1974). [14C]Glucose-P-dolichol was a kind gift from Dr N. Behrens, of this Institute. [14C]Palmitic acid was purchased from New England Nuclear. Other chemicals were obtained from commercial sources.

Enzyme preparations. The standard enzyme preparation consisted of cells of Xanthomonas campestris strain NRRL B-1459, grown, harvested and treated with EDTA as previously described (Ielpi et al., 1981). A membrane fraction obtained from fresh cells (about 500 mg) was resuspended in 10 ml 70 mM-Tris/HCl buffer, pH 8.2, and fractionated with an X-press (Biox, Sweden). The disrupted cell suspension was brought to 10 mM-MgCl2 and 3 mM-EDTA in 70 mM-Tris/HCl, pH 8.2, and incubated at 40 °C for 15 min in the presence of deoxyribonuclease II (Sigma, D4138) (0.1 mg ml-1). Coarse fragments were removed by centrifugation at 8700 g for 20 min and the supernatant was centrifuged at 130000 g for 3 h. Two fractions were obtained: a supernatant, termed the 'soluble fraction', which contained 25–30 mg protein ml-1, and a pellet, which, resuspended in 70 mM-Tris/HCl buffer, pH 8.2, was termed 'insoluble membrane fraction' and contained 30–33 mg protein ml-1. Protein concentration was determined by the Lowry method. The enzyme preparations contained endogenous donors as well as acceptors of galacturonic acid residues, in varying amounts for the different batches.

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Standard incubation assays. The reaction mixture contained the following components: 70 mM-Tris/HCl buffer, pH 8.2; EDTA-treated whole cells (or soluble or insoluble membrane fractions, where indicated) (0.9-1 mg protein), 16 mM-MgCl₂ and 0.01 mM-UDP-[¹⁴C]GlcA. The reaction mixtures (final volume 70 µl) were incubated at 15 °C for 30 min; the reaction was stopped by adding 0.1 ml butanol-1-ol and mixing by vortexing. The mixture was centrifuged in a bench-top Eppendorf centrifuge at 14000 r.p.m. for 2 min and the upper layer, containing the lipophilic incorporation products, carefully removed. This extraction was repeated three times and the combined butanol layers were washed with water in a similar way (4 x 0.1 ml each). Radioactivity incorporated into the butanol extract was counted in Bray's solution (Bray, 1960) with a Tri-Carb scintillation counter (Packard).

Chemical treatments. Mild acid hydrolysis, at pH 2 (0.01 M-HCl) and 100 °C for 10 min, mild alkaline treatment [0.06 M-NaOH in chloroform/methanol (4:1, v/v) at room temperature for 5 min] and phenol treatment (50%, w/v, phenol at 68-70 °C for 2 h) were done as previously described (Garcia et al., 1974).

Reduction of carboxyl groups to the respective alcohols was done by the method of Taylor & Conrad (1972) modified in the following way. The butanolic extract was treated with 10 mg solid EDC [1-ethyl-3-(3-dimethylaminopropyl) carbodiimide] for 2 h at room temperature maintaining a constant pH of 4-7 with 0.5 M-HCl before slowly adding 0.1 ml 3 M-sodium borohydride in butanol. During this operation the pH was maintained at 7 by addition of 4 ml-HCl. This procedure was done at 0 °C to avoid hydrolysis of the product, before leaving the reaction mixture at room temperature for 4 h. After this period, 0.1 ml water and sufficient Dowex 50 (H⁺ form) resin were added to lower the aqueous phase to pH 4. The two phases were separated by centrifugation. About 90% of the radioactivity was recovered in the butanolic phase.

Catalytic reduction was done by bubbling H₂ gas into the butanolic extract containing (2-4 mg) finely divided Pt, PtO₂ (Adam's catalyst, Aldrich) or 10₂₀ Pd on activated carbon (Aldrich) at room temperature done at 0°C to avoid hydrolysis of the product, before leaving the reaction mixture at room temperature for 4 h. After this period, 0.1 ml water and sufficient Dowex 50 (H⁺ form) resin were added to lower the aqueous phase to pH 4. The two phases were separated by centrifugation. About 90% of the radioactivity was recovered in the butanolic phase.

Chromatography and electrophoresis. Gel-filtration column chromatography on Sephadex LH-20 (1.2 x 160 cm) in 0.1 M-ammonium acetate in 99% methanol and ion-exchange column chromatography on DEAE-cellulose (1.2 x 58 cm) in 99% methanol with a 0-0.4 M-ammonium acetate gradient (in a total volume of 400 ml) were done as previously described (Dankert et al., 1966; Garcia et al., 1974) collecting 150 fractions of 0.5 ml and 3.0 ml respectively. Where indicated, the samples were desalted by LH-20 gel-filtration with 99% methanol.

Paper chromatography and electrophoresis were done on Schleicher and Schuell 2043 paper as previously reported (Garcia et al., 1974) using the following solvents: A, 1.2 M-pyridinium acetate buffer, pH 6.5; B, 0.1 M-sodium molybdate buffer, pH 5.0 (Bourne et al., 1961); C, nitromethane/acetic acid/ethanol/boric acid saturated solution (8:1:1:1, by vol.); D, ethanol/concentrated ammonia (7:3, v/v); E, propan-2-ol-acetic acid/water (27:4:9, by vol.) (Tung & Nordin, 1968). TLC was done on Silica Gel G plates (250 µm thick, Merck) with chloroform/methanol/water (60:25:4, by vol.) as solvent (Pont Lezica et al., 1975).

Enzyme treatments: Galactose oxidase assays (Sigma, type V) were done as described previously (Garcia et al., 1974); the oxidation product (Rₛᵤₐ₉, 0.52) was detected by paper chromatography in solvent E.

Results

Incorporation of ¹⁴C-label into a lipidic compound

Incorporation of radioactivity from UDP-[¹⁴C]GlcA into a butanol-soluble compound was dependent on the presence of Mg²⁺; no incorporation was observed in its absence and the maximum incorporation was reached at 16 mM. Under the standard conditions used the incorporation reached a plateau after 30 min incubation. Incorporation increased linearly with increased amounts of enzyme until a plateau was reached at about 1 mg protein. A plot of incorporated radioactivity versus UDP-[¹⁴C]GlcA concentration indicated that under the conditions of the assay (10 µM) there was a large excess of lipid acceptor in the preparation.

Increasing amounts of UDP inhibited incorporation of radioactivity into the butanolic extract and at 0-75 mM-UDP a 75-80% inhibition plateau was reached. UMP had almost no effect. This difference in behaviour is clearly shown in a time-course experiment in which UMP was added after 15 min incubation; the radioactivity incorporated was only slightly decreased (Fig. 1). If UDP was substituted for UMP, incorporation of radioactivity was abruptly interrupted, and even some delabelling could be observed (Fig. 1). This result suggests that UDP is possibly one of the reaction products and, as such, can reverse incorporation.
Fig. 2. Paper chromatography and electrophoresis of the incorporation product. A, Paper electrophoresis with buffer A of a butanol extract of the standard incubation mixture without treatment (I) and after mild acid hydrolysis (0.01 M-HCl, 100 °C, 10 min) (II). B, Paper electrophoresis with buffer B of the butanol extract after mild acid hydrolysis (I) or mild alkaline treatment (0.06 M-NaOH, room temperature, 5 min) (II). C, Paper electrophoresis with buffer C of ‘carboxyl reduced lipid-bound [14C]GalA’ without treatment (I) and after mild acid hydrolysis (II) or mild alkaline treatment (III). D, Paper chromatography with buffer C of ‘carboxyl reduced lipid-bound [14C]GalA’ after mild acid hydrolysis (I) or mild alkaline treatment (II). The arrows indicated the mobility of the respective standards and the dotted arrow, the solvent front. The maximum point on the radioactivity scale corresponds to 100 c.p.m.

**Isolation and chemical properties of ‘lipid-bound uronic acid’**

Analysis of the butanolic extract by paper chromatography with solvent D showed only one radioactive component, Rf 0.84, indicative of a lipophilic compound (not shown) (Garcia et al., 1974). This was confirmed by paper electrophoresis with buffer A in which all the radioactivity remained at the origin, as expected for a water-insoluble substance (Fig. 2A) (Couso et al., 1982).

**Fig. 3. LH-20 column chromatography of ‘lipid-bound uronic acid’ and ‘reduced lipid-bound uronic acid’.** The arrows indicate the elution position of [14C]glucuronic acid and [14C]palmitic acid (Palm) used as standards. The ‘lipid-bound [14C]uronic acid’ sample (40000 c.p.m.) consisted of a butanol extract of a standard incubation scaled up 10 times. Samples (0.1 ml) were counted for radioactivity (A). The ‘reduced lipid-bound [14C]uronic acid’ sample (20000 c.p.m.) was obtained by esterification with EDC and sodium borohydride reduction, as indicated in Methods. Samples (0.1 ml each) were counted for radioactivity (B).

TLC showed a single radioactive compound with a mobility intermediate between [14C]glucose-P-dolichol and [14C]galactose-P-prenol (not shown). The profile obtained by gel filtration on a Sephadex LH-20 column showed one peak with a small shoulder eluting slightly ahead of a GlcA standard (Fig. 3A).

DEAE-cellulose column chromatography gave only one component, eluting at 0.12 m-ammonium acetate (Fig. 4A). Under the same conditions [14C]galactose-P-prenol is eluted at 0.11 m- and [14C]glucose-P-P-prenol at 0.23 m-ammonium acetate. These results suggest that the radioactive compound formed should have only the negative charge of the uronic acid since the galactose derivative, which has a phosphate residue with one primary hydroxyl group, is eluted at about the same concentration. The diphosphate bridge of the glucose derivative, which was retained for longer in the column, has two primary hydroxyl groups.

Mild acid hydrolysis (pH 2, 100 °C, 10 min) of the butanol-soluble compound released practically all the radioactivity into the water phase (Table 1). Unexpect-
Fig. 4. DEAE-cellulose column chromatography of 'lipid-bound uronic acid' and 'reduced lipid-bound uronic acid'. The ammonium acetate gradient (---) was started at fractions 13 and 30 respectively (small arrow). The large arrows indicate the elution positions of [14C]Gal-P-prenol and [14C]Glc-P-P-prenol used as standards. The 'lipid-bound [14C]uronic acid' sample (30000 c.p.m.) consisted of a butanol extract of a standard incubation mixture scaled up 5 times. Samples (0.5 ml) were counted for radioactivity (A). The 'reduced lipid-bound [14C]uronic acid' sample (12000 c.p.m.) was obtained by esterification with EDC and sodium borohydride reduction, as indicated 'in Methods. Samples (0.5 ml each) were counted for radioactivity (B).

Fig. 5. Catalytic reduction of 'lipid-bound [14C]GalA. Butanol extracts (20000-30000 c.p.m.) were treated with H₂ in the presence of PtO₂ (2 mg) (○, ●) or 10% Pd on activated carbon (2 mg) (△, ▲) at room temperature. Samples (0-25 ml each) were taken at the times indicated and partitioned with water (0-25 ml) as indicated in Methods. Radioactivity in the butanolic (○, △) and aqueous phases (●, ▲) was counted in Bray's solution.

edly, when submitted to paper electrophoresis with buffer A this material behaved as galacturonic acid (Fig. 2A). This result was confirmed by paper electrophoresis with buffer B, a system that clearly distinguishes between galacturonic, glucuronic and mannuronic acids (Fig. 2B).

Phenol treatment, a procedure that splits allylic phosphate–prenol linkages (Garcia et al., 1974) releasing the phosphosugar moiety, had almost no effect on lipid-bound galacturonic acid ('lipid-bound GalA') even after 2 h treatment at 70 °C (Table 1).

Mild alkaline hydrolysis (0-06 M-NaOH, room temperature, 5 min) liberated all the radioactivity (Table 1) as galacturonic acid, as judged by paper electrophoresis with buffers A and B (Fig. 2B).

Table 1. Lability of different samples to chemical degradation treatments

<table>
<thead>
<tr>
<th>Sample</th>
<th>Percentage radioactivity released after treatment with:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>H⁺</td>
</tr>
<tr>
<td>Prenol-P-[14C]Gal</td>
<td>100</td>
</tr>
<tr>
<td>'Lipid-bound-[14C]GalA'</td>
<td>87</td>
</tr>
<tr>
<td>'Carboxyl reduced lipid-bound [14C]GalA'</td>
<td>67</td>
</tr>
<tr>
<td>H₂-Pt treated 'lipid-bound [14C]GalA*'</td>
<td>80</td>
</tr>
<tr>
<td>H₂-PtO₂ treated 'lipid-bound [14C]GalA*'</td>
<td>80</td>
</tr>
<tr>
<td>H₂-Pd treated 'lipid-bound [14C]GalA*'</td>
<td>82</td>
</tr>
<tr>
<td>H₂-Pt treated 'carboxyl reduced lipid-bound [14C]GalA*'</td>
<td>62</td>
</tr>
</tbody>
</table>

* Samples were treated as indicated for 3 h.
Table 2. Effect of unlabelled UDP-GlcA and UDP-GalA on incorporation of radioactivity into the butanolic extract

Conditions were as for the standard incubation assay, except for the addition of the unlabelled sugar donors at 0.1 mM or 0.5 mM final concentration, as indicated in each case. ND, Not detectable.

<table>
<thead>
<tr>
<th>Substrate donor</th>
<th>Radioactivity incorporated [c.p.m. (mg protein)-¹]</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>UDP-[¹⁴C]GlcA</td>
<td>1552</td>
<td>100</td>
</tr>
<tr>
<td>UDP-[¹⁴C]GlcA + UDP-GalA (1:10)</td>
<td>117</td>
<td>7</td>
</tr>
<tr>
<td>UDP-[¹⁴C]GlcA + UDP-GalA (1:50)</td>
<td>ND</td>
<td>0</td>
</tr>
<tr>
<td>UDP-[¹⁴C]GlcA + UDP-GlcA (1:10)</td>
<td>620</td>
<td>40</td>
</tr>
<tr>
<td>UDP-[¹⁴C]GlcA + UDP-GlcA (1:50)</td>
<td>152</td>
<td>10</td>
</tr>
</tbody>
</table>

Table 3. Enzyme location and substrate donor specificity

The standard assay was done as described in Methods except for the enzyme preparation or the sugar donor, which were as indicated in each case.

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Substrate donor (10 μM)*</th>
<th>Radioactivity incorporated [c.p.m. (mg protein)-¹]</th>
</tr>
</thead>
<tbody>
<tr>
<td>EDTA-treated cells</td>
<td>UDP-[¹⁴C]GlcA</td>
<td>2150</td>
</tr>
<tr>
<td>Membrane fraction</td>
<td>UDP-[¹⁴C]GlcA</td>
<td>1838</td>
</tr>
<tr>
<td>Soluble fraction</td>
<td>UDP-[¹⁴C]GlcA-GalA</td>
<td>4036</td>
</tr>
<tr>
<td>EDTA-treated cells</td>
<td>UDP-[¹⁴C]GlcA-GalA</td>
<td>170</td>
</tr>
<tr>
<td>Membrane fraction</td>
<td>UDP-[¹⁴C]GlcA-GalA</td>
<td>4036</td>
</tr>
<tr>
<td>Soluble fraction</td>
<td>UDP-[¹⁴C]GlcA-GalA</td>
<td>170</td>
</tr>
</tbody>
</table>

* The UDP-[¹⁴C]GlcA-GalA mixture contained 75% and 25% of the glucuronic and galacturonic acid derivatives, respectively.

'Lipid-bound GalA' was resistant to hydrogenolysis. After 3 h treatment with H₂ using either Pt, PtO₂ or 10% Pd on activated carbon as catalysts about 85% of the radioactivity still remained in the butanolic phase (Fig. 5). These treatments did not change substantially the acid and alkali lability of the butanol-soluble galacturonide (Table 1). Longer treatments (6 h) with any of the catalysts released 80–85% of the radioactivity into the water phase (Fig. 5). This material was identified as free galacturonic acid by paper electrophoresis with buffers A and B, which distinguishes galacturonic acid from galacturonic acid.

These results taken together indicated that either the true sugar donor was UDP-GalA or that once transferred to the acceptor lipid the glucuronic acid residue was epimerized to galacturonic acid. This latter possibility was ruled out by competition experiments with unlabelled UDP-GalA (Table 2). In the presence of unlabelled UDP-GlcA or UDP-GalA incorporation was lowered to 40% and 7%, respectively, for a 1:10 molar ratio; for a 1:50 molar ratio incorporation of radioactivity was lowered to 10% with UDP-GlcA and was not detectable with UDP-GalA. Furthermore, incorporation from the mixture of UDP-[¹⁴C]GlcA and UDP-[¹⁴C]GalA (UDP-[¹⁴C]GlcA-GalA) was higher than from pure UDP-[¹⁴C]GlcA (Table 3). This approach was confirmed by the characterization of a UDP-GlcA-4-epimerase activity in these enzyme preparations (M. Bravo & M. A. Dankert, unpublished results).

Carboxyl reduced lipid-bound GalA

'Lipid-bound [¹⁴C]GalA', purified by DEAE-cellulose column chromatography and desalted by LH-20 gel filtration, was esterified in the presence of EDC and treated with sodium borohydride in order to reduce the carboxyl groups to primary alcohols. More than 87% of the label remained butanol soluble. Gel filtration of this material through a Sephadex LH-20 column gave a profile similar to that obtained with 'lipid-bound GalA' (Fig. 3B).

'Carboxyl reduced lipid-bound GalA', chromatographed on a DEAE-cellulose column, was eluted before starting the ammonium acetate elution gradient, instead of at 0.12 M as observed for the non-reduced product,
indicating that the negative charge had been lost (Fig. 4B). This was confirmed by mild acid hydrolysis of the 'carboxyl reduced lipid-bound GalA' which liberated a neutral compound, 14C-labelled, as judged by paper electrophoresis with buffer A (Fig. 2C), which had the mobility of [14C]galactose by paper chromatography with solvent C (Fig. 2D). Its identity was confirmed by incubation with galactose oxidase. Consistently, the acid lability of the 'carboxyl reduced lipid-bound GalA' was slightly lowered (Table 1).

'Carboxyl reduced lipid-bound GalA' was not affected by phenol treatment nor by catalytic reduction, but was degraded to [14C]galactose by mild alkaline hydrolysis (Table 1 and Fig. 2D).

**Enzyme location**

Experiments done with disrupted cells and UDP-[14C]GlcA as donor indicated that galacturonic acid transfer activity in the 'membrane fraction' was ten times higher than in the 'soluble fraction'. Nevertheless, the activity was maximal with EDTA-treated cells (Table 3). When the UDP-[14C]GlcA–GalA mixture was substituted for UDP-[14C]GlcA the maximum incorporation was obtained with the 'membrane fraction'; EDTA-treated cells incorporated 72% and the 'soluble fraction' 4-2% of the radioactivity present in the 'membrane fraction' (Table 3). These results agree with the fact that most of the UDP-GlcA-4-epimerase activity is recovered in the 'soluble fraction' (M. Bravo & M. A. Dankert, unpublished results).

**Discussion**

The results presented above show that EDTA-treated *X. campestris* cells synthesize in vitro a highly lipophilic galacturonide, with unusual chemical properties. The enzyme preparation also contains UDP-GlcA-4-epimerase activity since UDP-[14C]GlcA can be used as sugar donor. The sequence of reactions can be represented as follows.

\[
\text{UDP-GlcA} \rightleftharpoons \text{UDP-GalA} \quad (1)
\]

\[
\text{UDP-GalA} + \text{lipid} \rightleftharpoons \text{lipid-bound GalA} + \text{UDP} \quad (2)
\]

Reaction (1) has been described for other systems (Fan & Feingold, 1972; Feingold et al., 1960; Gaunt et al., 1974) and a study of the present enzyme, which is Mg2+-independent, will be reported elsewhere.

Reaction (2), on the contrary, has an absolute requirement for Mg2+, the optimal concentration being 16 mM. The reaction is reversible, as suggested by the loss of label observed in the presence of excess UDP (Fig. 1). Only the sugar moiety is transferred since UMP has almost no effect on the reaction (Fig. 1) and no radioactivity was detected in the butanol extract when [β-32P]UDP-GlcA was used as donor (not shown). Furthermore no evidence for the presence of a phosphate group in 'lipid-bound GalA' was detected (see below).

The transferase activity is located mainly in the insoluble 'membrane fraction' (Table 3) and could not be totally separated from reaction (1). For this reason, no further studies were made.

The properties of the galacturonide formed, called 'lipid-bound GalA' are quite unusual and differ from those of prenyl-phosphosugars and other glycosides so far described (Ballou, 1984; Pont Lezica et al., 1975).

'Lipid-bound GalA' seems to be fairly homogeneous. TLC analysis showed only one radioactive substance, although the profile obtained by LH-20 gel filtration could indicate either some heterogeneity or partial adsorption to the gel. The carboxyl reduced derivative demonstrates a similar pattern indicating that this group is not the cause of tailing.

DEAE-cellulose column chromatography of 'lipid-bound GalA' produced a single sharp peak in the position expected for a compound with only one negative charge (as the Gal-P-lipid used as standard) attributable to the galacturonic acid moiety. This interpretation was confirmed by reduction of the carboxyl group to obtain the galactose derivative: the DEAE-cellulose elution pattern of 'carboxyl reduced lipid-bound GalA' showed that the negative charge had been lost.

'Lipid-bound GalA' was very sensitive to mild acid, as expected for a prenyl-phosphosugar but was degraded neither by phenol treatment nor by catalytic reduction in contrast to allylic prenols (Table 1). In this respect it behaves as a dolichol derivative (Pont Lezica et al., 1975).

Most surprisingly, 'lipid-bound GalA' was degraded by mild alkali, liberating free galacturonic acid. This was an unexpected property. Sugar monophosphate prenols, both allylic and dolicolic, are resistant to this treatment, and the diphosphate derivatives are either resistant or produce the 1,2-cyclic phosphate ester of the sugar, provided the hydroxyl groups at carbons 1 and 2 are in the cis-position and the latter is not substituted (Garcia et al., 1974). For the 'lipid-bound GalA' these possibilities are ruled out because it does not contain phosphate residues. Regular glycosides are relatively acid- or alkali-resistant (Ballou, 1984) and the only reported derivatives easily degraded under both conditions are some enolic (or vinylic) glycosides (Mead et al., 1958; Wakabayashi et al., 1964). To test this possibility and to gain some insight into the nature of the aglycone the 'lipid-bound' derivatives were treated with H2 in the presence of...
different catalysts (Rylander, 1969). ‘Lipid-bound GalA’ and ‘carboxyl-reduced lipid-bound GalA’ were resistant for short reduction times (Table 1). Longer treatments led to liberation of free galacturonic acid or galactose, respectively. The Pd catalyst is reputed to favour reduction of double bonds prior to hydrogenolysis (Alden & Davies, 1968). This would cause a glycosidic-enol linkage to become a regular hydroxylidy linkage, providing greater stability to acid and/or alkali degradation. No such changes were observed with the ‘lipid-bound GalA’ (Table 1). Since the nature of the lipid or aglycone moiety is unknown, the structure of this part of the molecule is currently under study.

To our knowledge this is the first acid- and alkali-labile lipid-linked galacturonide reported. The synthesis of related lipid-bound glucuronides has been described in chicken liver (Cummings & Roth, 1982) and in Acetobacter xylinum (Couso et al., 1986). Some of their properties coincide with those of ‘lipid-bound GalA’, for example acid- and alkali-lability, but both compounds have an extra negative charge, attributed to a phosphate group in the chicken liver substance and of unknown origin in the A. xylinum derivative.

Very little can be said about the function of ‘lipid-bound GalA’. Its lipophilic nature and the lability of its glycosidic linkage would make it an adequate galacturonyl donor at the cytoplasmic membrane level. It could also be a lipophilic uronic acid reservoir. We are tempted to suggest that, together with the chicken liver and the A. xylinum component of a phenol-soluble lipopolysaccharide from X. campestris, the A. xylinum glucuronides, the X. campestris gluconoxide belongs to a new type of ‘activated sugar’.

D-Galacturonic acid has been reported to be a minor component of a phenol-soluble lipopolysaccharide from X. campestris (Hickman & Ashwell, 1966) and a galacturonyl phosphate ester of unknown function has also been described in this organism (Volk, 1968). We have no evidence to relate these compounds to the ‘lipid-bound GalA’ and the UDP-GalA here described, but so far they are the only GalA-containing derivatives reported in X. campestris.

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References