Effect of replacing the general energy-coupling proteins of the PEP:sugar phosphotransferase system of Salmonella typhimurium with their fructose-inducible counterparts on utilization of the PTS sugar glucitol

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A strain of Salmonella typhimurium in which the genes encoding the general phosphoenolpyruvate:sugar phosphotransferase system (PTS) proteins HPr and Enzyme I have been deleted, the normally cryptic gene encoding the fructose-inducible Enzyme I (EI* or Ef\text{fructose}) is expressed, and the fructose repressor protein is inactive \((fruR\) or \(cra\) mutant) was studied. This strain lacks HPr and EI, but expresses FPr (DTP) and EI\text{fructose} constitutively. Since FPr and Ef\text{fructose} can substitute for HPr and EI, the strain grew in minimal liquid medium supplemented with the PTS sugars glucose, fructose, \(N\)-acetylglucosamine, mannitol or mannose. However, it showed very poor to negligible growth on the PTS sugar glucitol. It also grew very poorly on the non-PTS sugars maltose, melibiose and especially glycerol. Adding cAMP to the medium allowed growth on glucitol, but did not affect growth on glycerol. We suggest that poor phosphorylation of the regulatory molecule Enzyme IIA\text{glucose} by FPr is responsible for these effects.

**Keywords:** phosphoenolpyruvate:sugar phosphotransferase system, carbon catabolite control, gene regulation, carbohydrate transport

INTRODUCTION

The bacterial phosphoenolpyruvate (PEP):sugar phosphotransferase system (PTS) catalyses the uptake and concomitant phosphorylation of a number of carbohydrates (for reviews see Postma et al., 1993; Meadow et al., 1990; Saier, 1989). In Escherichia coli and Salmonella typhimurium, PTS sugars include glucose, mannose, fructose, mannitol, glucitol, galactitol and \(N\)-acyetylglucosamine. The general energy-coupling proteins Enzyme I (EI) and HPr interact with sugar-specific Enzymes II (EIIsugar), with the exception of the fructose system, which has its own HPr-like domain, FPr. EI autophosphorylates, using PEP as the phosphate donor; the phosphoryl group is then transferred successively to HPr, then to the A domain of a sugar-specific EII, then to the B domain of the same EII, then to the incoming sugar as it enters the cell via the C domain of the EII. Other carbohydrates are taken up by other mechanisms. These non-PTS sugars include melibiose, glycerol and maltose in both \(E\). coli and \(S\). typhimurium, as well as lactose in \(E\). coli.

The fructose PTS is structurally unique in that its HPr counterpart is a domain of a larger protein, DTP (diphosphoryltransfer protein); the other two domains are the sugar-specific phosphocarrier domain EIIA\text{Fruc} and a third domain M (modulator) which is thought to play a regulatory role (Geerse et al., 1989; Wu et al., 1990). It has been long established that FPr can substitute for HPr in PTSs for other sugars besides fructose both \textit{in vivo} and \textit{in vitro} (Saier et al., 1970, 1976). In complementation assays it was observed that purified FPr, whether as part of the larger DTP or in the form of a 9 kDa separate protein (free FPr), worked about 5% as well as HPr in the PTSs for mannose, \(N\)-acyetylglucosamine, mannitol and glucitol, but only 0.5% as well as HPr in the glucose-specific PTS (Sutrina et al., 1988). More recently, \textit{in vitro} assays by a different group indicated that both DTP and free FPr accepted a phosphate group...
from EI about 40% as well as HPr, and donated the phosphate group to EIIA\(^{\text{Glu}}\) about as well as HPr (Lux et al., 1995). These results taken together suggest that FPr, molecule for molecule, does not interact with EI as effectively as HPr, and that it also interacts relatively poorly with the A domain of the glucose-specific Enzyme effectivly as HPr, and that it also interacts relatively molecule for molecule, does not interact with EI as poorly as HPr. In a recent study, DTP was also shown to substitute for HPr in both the activation of the antiterminator BglG of the \(\beta\)-glucosidases (\(\beta\)-gl) operon of \(E. coli\) and in the negative regulation of this same protein by phospho-Enzyme IIBCA\(^{\text{Glu}}\) (Görke & Rak, 1999).

In \(E. coli\) and \(S. typhimurium\), EIIA\(^{\text{Glu}}\) mediates control of utilization of non-PTS sugars by the PTS (reviewed by Saier, 1989; Postma et al., 1993; Meadow et al., 1990). A model to account for this was proposed. When in its phosphorylated form, EIIA\(^{\text{Glu}}\)-P activates adenylate cyclase (Reddy & Kamireddi, 1998). When in its unphosphorylated form, EIIA\(^{\text{Glu}}\) prevents the uptake of non-PTS sugars by binding to and inhibiting the permeases for these sugars, or, in the case of glycerol, by binding to and inhibiting glycerol kinase, a phenomenon called induced exclusion. In the absence of PTS sugars, the ratio of EIIA\(^{\text{Glu}}\)-P to EIIA\(^{\text{Glu}}\) is relatively high, leading to relatively high levels of cAMP and to active non-PTS permeases. This allows for high level expression of catabolite activator protein (CAP)–cAMP-dependent operons in general and to induction of the genes for the uptake and utilization of the non-PTS sugars in particular. In the presence of a PTS sugar, phosphate is drained off EIIA\(^{\text{Glu}}\)-P, and inducer exclusion/catabolite repression results. Recent studies challenge the model just described; it has been proposed, based on these studies, that the role of cAMP in glucose/lactose diauxie in \(E. coli\) is to enhance inducer exclusion by activating the membrane component of the glucose PTS, EIIIC\(^{\text{Glu}}\), at the transcriptional level (Inada et al., 1996; Kimata et al., 1997). In contrast, another recent study suggests that, in \(E. coli\), cAMP-dependent catabolite repression, rather than inducer exclusion, may be the dominant mechanism by which glucose represses glycerol kinase levels during diauxic growth (Holtman et al., 2001). In \(S. typhimurium\), both mechanisms may be important in glucose/glycerol diauxie (Novotny et al., 1985).

In \(S. typhimurium\) strain LJ705 the structural genes for EI and HPr have been deleted, and the \(fruR\) gene encoding the fructose repressor protein FruR, or Cra, has been disrupted by a \(Tn10\) insertion; this strain also expresses the normally cryptic EI-like protein EIIIC\(^{\text{Glu}}\) (EI\(^{\text{Glu}}\)), which is part of the \(fru\) regulon (Chin et al., 1987). This strain is able to grow on PTS sugars, since the constitutively expressed FPr (DTP) and EI\(^{\text{Glu}}\) substitute for the missing HPr and EI. In this study, we found that growth of LJ705 on the PTS sugar glucitol as its sole carbon source is poor to negligible and that its growth on the non-PTS sugars melibiose, maltose and especially glycerol is also poor. Addition of cAMP to the growth medium markedly improved growth on glucitol, but not on glycerol. We propose that underphosphorylation of EIIA\(^{\text{Glu}}\) by FPr in this strain is responsible for these effects; low cAMP leads to low activation of the CAP–cAMP-dependent glucitol operon (Yamada & Saier, 1988) and a high EIIA\(^{\text{Glu}}\)/EIIA\(^{\text{Glu}}\)-P ratio causes inducer exclusion of non-PTS sugars even in the absence of PTS sugars. In this study we have focussed primarily on the lack of ability of LJ705 to grow on glucitol. Its poor growth on non-PTS sugars is currently under further investigation.

**METHODS**

**Growth rate studies.** Bacterial strains were grown in either Luria–Bertani (LB) broth or minimal medium 63 (Saier et al., 1970) supplemented with tryptophan (50 \(\mu\)g ml\(^{-1}\)) and carbohydrates (at concentrations indicated). Other supplements and antibiotics were included where indicated. Strains were incubated at 37 °C, at 175–180 r.p.m. Growth was monitored by measuring the optical density at 540 nm, using a Beckman DU-65 spectrophotometer with a 1 cm cuvette.

**Strains.** *Salmonella typhimurium* strains used were LJ705 (AcysK ptsH141 fruRS1::Tn10 ptsJ52), SB1475 (ptsH15), SB3507 (trpB223), SB1667 (mal) and SB1744 (malA61). The ptsJ52 mutation in LJ705 allows expression of the normally cryptic EIIIC\(^{\text{Glu}}\). All of these strains are derivatives of wild-type strain LT-2. *Escherichia coli* strains used were K12 (wild-type) and ESK147 (F\(^{-}\)ompT tph- mfr- pylS\(^{-}\) cm\(^{-}\)). Strains ESK147 (E. B. Waygood, Department of Biochemistry, University of Saskatchewan), SB3507 (Cordaro & Roseman, 1972) and SB1667 (M. H. Saier, Jr, University of California at San Diego) are wild-type with respect to the PTS.

**Preparation of crude extracts.** Cultures were grown to late exponential phase in LB broth and harvested by centrifugation. The pellets were washed twice with 0.15 M NaCl and once with buffer (50 mM Tris–HCl, pH 7.5, 0.1 M PMSF, 0.2 M DTT), then resuspended in a small volume of the same buffer. The suspensions were passed through the French press three times at 10000 p.s.i. (69 MPa). Unbroken cells and debris were removed by centrifugation (10000 r.p.m., 10 min, JA17 rotor).

**Preparation of washed membranes.** Crude extracts were subjected to centrifugation for 4 h at 15000 r.p.m. in a JA17 rotor. The membrane pellets were resuspended in the above buffer and the centrifugation step was repeated. The washed membrane pellets were resuspended in a minimal volume of buffer.

**Assays.** Complementation assays for PTS components were conducted as previously described, using \(^{14}\)C-labelled sugar substrates and PEP (Reizer et al., 1992). In assays of crude extracts for the sugar-specific PTS components, partially purified HPr and EI were added to ensure an excess of these general energy-coupling proteins.

Transphosphorylation assays for the glucitol-specific membrane component EI(BC)\(^{\text{Glu}}\) were conducted as previously described (Saier et al., 1977). Assay mixtures contained 10 \(\mu\)M [\(^{14}\)C]gluconolactone (5 \(\mu\)Ci \(\mu\)mol\(^{-1}\); 1.85 kBq \(\mu\)mol\(^{-1}\)) as substrate and 10 \(\mu\)M glucitol 6-phosphate as phosphate donor, as well as 50 mM KPO\(_4\), pH 6.0, 25 mM KF, 12.5 mM MgCl\(_2\), 2.5 mM MgCl\(_2\) and the membrane preparation being assayed; the total volume was 100 \(\mu\)l. After incubation for 30 min at 37 °C, the reactions were stopped by adding 1 ml water. The [\(^{14}\)C]gluconolactone product was collected using anion exchange columns and quantified with a scintillation counter (Kündig &
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Roseman, 1971). The Lowry assay was used to determine total protein concentration.

**RESULTS AND DISCUSSION**

**Growth of LJ705 versus wild-type strains**

*S. typhimurium* strain LJ705 lacks the general PTS component proteins HPr and EI but produces their normally fructose-inducible counterparts FPr and EFru constitutively. The latter proteins substitute for the former, allowing the strain to utilize PTS sugars (Chin *et al.*, 1987). We compared growth of this strain under various conditions to that of two wild-type *S. typhimurium* strains, SB3507 and SB1667, and two wild-type *E. coli* strains, ESK147 and K12.

On minimal medium, growth of LJ705 was poor to negligible, with respect to both rate and extent, on the PTS sugar glucitol relative to its growth on other PTS sugars (glucose, fructose, N-acetylglucosamine, mannose and mannitol) or galactose. Growth on mannitol was also somewhat slow compared to the other PTS sugars, and growth on glucose (and galactose, which is not shown) somewhat fast (Fig. 1b). The growth rate did not increase when the concentration of glucitol was increased from the 0-2% used in most of our studies to 0-5% or 1% ; decreasing the concentration also did not promote growth. This strain also grew very poorly on the non-PTS sugars maltose, melibiose and especially glycerol (not shown). We normally included tetracycline (20 µg ml⁻¹) in the medium for LJ705; omission of the drug did not affect the growth rate.

The wild-type *S. typhimurium* strain SB3507 grew faster than LJ705 on all the PTS sugars; it also grew more slowly, and with an initial lag, on glucitol than on other PTS sugars, but was clearly able to utilize glucitol (Fig. 1a). Both of the wild-type *E. coli* strains also grew on glucitol, as did another wild-type *S. typhimurium* strain, SB1667. Like SB3507, strain SB1667 grew relatively slowly (with a similar lag period) on glucitol relative to other PTS sugars.

On rich medium, little difference in growth rate or extent was observed when LJ705 was grown on unsupplemented LB or on glucose, fructose, mannitol or glucitol or combinations of glucitol and one of the other sugars, and there was no large difference between growth rates of LJ705 and the wild-type strain SB3507. Supplementing minimal medium with tryptone (0-5 %) also minimized the differences in growth rate. On
fermentation plates, glucitol effects were subtle; LJ705 generally showed a slightly slower positive response on EMB or MacConkey plates containing glucitol than did wild-type strains. This was not reported previously. All strains tested showed a weak response on glucitol plates relative to other PTS sugars.

Since glucitol did not hinder growth of LJ705 on rich medium or on medium supplemented with other sugars, the effect did not appear to be due to toxicity due, for example, to build-up of glucitol-6-phosphate as a result of defective glucitol-6-phosphate dehydrogenase. Rather, the strain seemed to be unable to utilize glucitol as a growth substrate.

**Growth of a *ptsH* mutant versus LJ705**

One possible explanation for the inability of LJ705 to grow on glucitol is that FPr interacts so poorly with the glucitol-specific PTS protein EIIA<sup>Gal</sup> that glucitol cannot be taken up fast enough to support growth. This seemed unlikely since a previous *in vitro* study indicated that FPr substitutes equally well for HPr in the PTSs for mannitol, mannose, glucitol and N-acetylglucosamine, and relatively (10×) poorly only in the glucose system (Sutrina et al., 1988). On the other hand, the *in vitro* study was done under conditions such that crosstalk between the glucitol and mannitol systems could have confused the results (Saier et al., 1977). To investigate this possibility, we compared growth of LJ705 to that of *S. typhimurium* strain SB1475 (Saier et al., 1976), which carries a point mutation in the structural gene for HPr, *ptsH*. The latter strain has negligible HPr activity, and because the *ptsH* mutation is pleiotropic, it also has low EI activity; in contrast to LJ705, it does not express EI<sup>Py</sup> and production of FPr (DTP) is not constitutive.

As expected, SB1475 grew relatively well (usually slightly more slowly than LJ705) on fructose, but not on most other PTS sugars, as the sole carbon source; it grew, slowly, on glucose (Fig. 1c). As expected, on minimal medium supplemented with a mixture of fructose (0.02%) and glucitol, mannitol, mannose or N-acetylglucosamine (0.2%), this strain continued to grow on the other PTS sugar after the fructose should have been consumed (or grew on both sugars simultaneously), using fructose-induced FPr to substitute for HPr in the uptake of the other PTS sugars (Fig. 1d). Thus, poor interaction of FPr with the glucitol PTS cannot be the sole explanation for the failure of LJ705 to grow on glucitol. Interestingly, glucitol and mannitol seemed to slow down growth of SB1475 on fructose, N-acetylglucosamine may have increased the growth rate on fructose slightly, and growth on a mixture of fructose and glucose was monophasic and considerably more rapid than on fructose alone. The latter result was surprising, since FPr works poorly with EIIA<sup>Glc</sup>, but was consistent with the finding that LJ705 grew better on glucose than on other PTS sugars. The fact that glucose is capable of entering *S. typhimurium* cells via routes other than the glucose-specific PTS, e.g., it is a good substrate of the mannose PTS and is also a substrate of the non-PTS galactose permease (Postma & Stock, 1980), probably accounts for these results.

**Effect of cAMP on growth of LJ705**

Although FPr substitutes for HPr, it interacts relatively poorly with EI (Lux et al., 1995; Sutrina et al., 1988), and, as mentioned above, the latter study suggested that, relative to other EIAs, its interaction with EIIA<sup>Glu</sup> is particularly poor. Thus this mediator of inducer exclusion/catabolite repression may be underphosphorylated in LJ705. It is possible that the substitution of EI<sup>P24</sup> for EI also contributes to this effect; *in vitro* studies suggested that EI<sup>P24</sup> behaves very similarly to EI, but these studies were not extensive (Chin et al., 1987). This protein remains something of a mystery, since it has not been identified as any of the known paralogues of EI in the sequences of *E. coli* or *S. typhimurium*. Since the phosphorylated form of EIIA<sup>Gal</sup> mediates catabolite repression by activating adenylate cyclase, its underphosphorylation, by FPr, in LJ705 could lead to low levels of cAMP, and thus to low levels of expression of cAMP-dependent operons. The *gut* operon absolutely requires cAMP for expression (Yamada & Saier, 1988). The non-PTS operons for glycerol, melibiose and maltose utilization are also under CAP–cAMP control.

To test our hypothesis, we added 1 mM cAMP to the medium of either LJ705 or the wild-type strain SB3507 growing on 0.2% glucitol. The cAMP had little effect on the growth rate of the wild-type strain (Fig. 2a). The growth rate of LJ705 increased markedly, after a lag; when the experiment was extended further than the one shown in Fig. 2(b), the final density of the culture approached that of the wild-type strain. We also added cAMP to cultures of the wild-type strain, LJ705 and the mutant SB1475 growing on a mixture of 0.2% glucitol and 0.02% fructose. The wild-type strain kept growing on glucitol after the fructose should have been exhausted (or grew on both sugars simultaneously; there was no obvious lag phase); cAMP had no effect on the growth rate (Fig. 2a). After exhaustion of fructose, the growth rate of LJ705 dropped to its usual negligible level; cAMP allowed a continuation of growth on glucitol, after a lag, and a final density approaching that of the wild-type culture (Fig. 2b). As above, glucitol appeared to slow down the growth of SB1475 on fructose, but growth continued, at a slow rate, after the fructose should have been exhausted. Addition of cAMP did not appear to affect the growth rate of this mutant strongly; in the experiment shown, a slight positive effect was observed (Fig. 2c), while a duplicate experiment showed a slight negative effect.

The experiment was repeated with cultures of SB3507 and LJ705 growing on minimal medium supplemented with 0.2% glycerol or with a mixture of 0.2% glycerol and 0.02% fructose (not shown). The wild-type strain grew somewhat slowly on glycerol compared to glucose, with an initial lag; LJ705 grew slowly after a very long initial lag (8–12 h). On the mixture, the wild-type strain...
Growth of *S. typhimurium* on glucitol

![Graphs showing growth of *S. typhimurium* strains on glucitol](image)

**Fig. 2.** Effect of cAMP on growth of wild-type (SB3507), LJ705 and HPr mutant on glucitol. *S. typhimurium* strains were grown in minimal medium supplemented with 50 µg tryptophan ml⁻¹ and 0-02% fructose or 0-2% glucitol, or a mixture of 0-02% fructose and 0-2% glucitol. Where indicated, 1 mM cAMP was added at 2 h. (a) SB3507 (wild-type), (b) LJ705, (c) SB1475. ○, Fructose; •, glucitol; □, glucitol + cAMP; ▲, fructose + glucitol; △, fructose + glucitol + cAMP.

**Table 1.** Specific activities of glucitol-specific Enzyme II(BC) in membrane preparations

<table>
<thead>
<tr>
<th>Growth conditions*</th>
<th>Strain</th>
<th>SB3507 (WT)</th>
<th>LJ705</th>
<th>SB1475 (HPr⁻)</th>
<th>SB1677 (WT)</th>
<th>SB1744 (mtlA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No sugar</td>
<td></td>
<td>16.2±4.8</td>
<td>8.2</td>
<td>25.6±12.5</td>
<td>79.4</td>
<td>31.8</td>
</tr>
<tr>
<td>Gut</td>
<td></td>
<td>370±59</td>
<td>160±19</td>
<td>52.9±15.1</td>
<td>427</td>
<td>232</td>
</tr>
<tr>
<td>Gut + cAMP</td>
<td></td>
<td>539±132</td>
<td>315±31</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Gut + Glc</td>
<td></td>
<td>129±18.1</td>
<td>77.3±2.9</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
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<tr>
<td>Gut + Glc + cAMP</td>
<td></td>
<td>100</td>
<td>108</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Gut + Mtl</td>
<td></td>
<td>22.0±3.6</td>
<td>18.1±9.1</td>
<td>ND</td>
<td>ND</td>
<td>176</td>
</tr>
<tr>
<td>Gut + Mtl + cAMP</td>
<td></td>
<td>22.4</td>
<td>15.5</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Gut + Fru</td>
<td></td>
<td>278±56</td>
<td>66.7±8.6</td>
<td>161±24</td>
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<td>ND</td>
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<tr>
<td>Gut + low Fru</td>
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<td>390</td>
<td>183</td>
<td>155</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Gut + GlcNac</td>
<td></td>
<td>117</td>
<td>36.7</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Glc</td>
<td></td>
<td>32.4</td>
<td>24.5</td>
<td>ND</td>
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<tr>
<td>Mtl</td>
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<td>21.6</td>
<td>25.5</td>
<td>ND</td>
<td>ND</td>
<td>30.6</td>
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<tr>
<td>Fru</td>
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<td>29.5</td>
<td>14.6</td>
<td>17.5</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>GlcNac</td>
<td></td>
<td>27.5</td>
<td>2.55</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>

*0.5% sugar, except ‘low fructose’ (0.02%). Gut, glucitol; Glc, glucose; Mtl, mannitol; Fru, fructose; GlcNac, N-acetylglucosamine.
extracts were then prepared and assayed as previously
difficult to interpret due to interference by the
mannitol-specific PTS component EIICBA
et al
glucitol PTS assay (Grenier
To determine the levels of glucitol-specific PTS components in LJ705
levels of cAMP for full activation than do the man, mtl, glc and
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compared to membranes from glucitol-grown wild-type strain SB3507. Another wild-type strain, SB1667, was also found to have high specific activity relative to LJ705. In the presence of 1 mM cAMP, specific activities of EII(BC)\textsuperscript{Gut} in both SB3507 and LJ705 membranes increased, but the increase was more substantial for the mutant strain; values for the mutant strain grown with cAMP approached those for the wild-type strain grown without. Thus the rather small deficiency could be responsible for the inability of LJ705 to grow on glucitol.
Membrane preparations from SB1475 were also assayed for EII(BC)\textsuperscript{Gut}. Maximal induction by glucitol was seen only in the presence of both glucitol and fructose. Membranes from cells grown in the presence of 0·5% glucitol and either 0·5% or 0·02% fructose showed EII(BC)\textsuperscript{Gut} specific activities similar to glucitol-grown L705. Thus it is not clear why this strain but not LJ705 continued to grow on glucitol after fructose should have been exhausted, while LJ705 showed a long lag before resuming slow growth on glycerol. Addition of cAMP during the lag period had negligible effect on the growth rate of LJ705 on glycerol or on the mixture of fructose and glycerol.
The results suggest that low expression of the gut operon due to low levels of cAMP is responsible for poor growth of LJ705 on glucitol. On glycerol, addition of cAMP should promote catabolite derepression, but a relatively high ratio of unphosphorylated EIIA\textsuperscript{Glc}-P may still cause inducer exclusion (EIIA\textsuperscript{Glc} inhibits glycerol kinase). Our results suggest that the gut operon of S. typhimurium requires higher levels of cAMP than do the man, mtl, glc and nag PTS operons.

**Levels of glucitol-specific PTS components in LJ705 versus other strains**

To determine the levels of the glucitol-specific PTS components, strains of *S. typhimurium* were grown in liquid LB broth supplemented with 0·5% glucitol or with 0·5% glucitol plus another PTS sugar (usually 0·5%) and/or cAMP (1 mM, added at 2·5 h); crude extracts were then prepared and assayed as previously described (Reizer et al., 1992). The results (not shown) were difficult to interpret due to interference by the mannitol-specific PTS component EIICBA\textsuperscript{Gut} in the glucitol PTS assay (Grenier et al., 1985; Saier et al., 1977). To eliminate this problem, washed membranes were prepared and the transphosphorylation assay was carried out using [\(^{14}\)C]glucitol as substrate and glucitol 6-phosphate as phosphate donor (Saier et al., 1977). Results, expressed as pmol [\(^{14}\)C]glucitol 6-P formed min\(^{-1}\) (mg membrane protein\(^{-1}\)), are shown in Table 1. The specific activity of the membrane component of the glucitol PTS, EII(BC)\textsuperscript{Gut}, was consistently lower (by 50–70%) in membranes from glucitol-grown LJ705 compared to membranes from glucitol-grown wild-type strain SB3507. Another wild-type strain, SB1667, was also found to have high specific activity relative to LJ705. In the presence of 1 mM cAMP, specific activities of EII(BC)\textsuperscript{Gut} in both SB3507 and LJ705 membranes increased, but the increase was more substantial for the mutant strain; values for the mutant strain grown with cAMP approached those for the wild-type strain grown without. Thus the rather small deficiency could be responsible for the inability of LJ705 to grow on glucitol.

Membrane preparations from SB1475 were also assayed for EII(BC)\textsuperscript{Gut}. Maximal induction by glucitol was seen only in the presence of both glucitol and fructose. Membranes from cells grown in the presence of 0·5% glucitol and either 0·5% or 0·02% fructose showed EII(BC)\textsuperscript{Gut} specific activities similar to glucitol-grown L705. Thus it is not clear why this strain but not LJ705 continued to grow on glucitol after fructose should have been exhausted when exposed to a mixture of the two sugars.

Other PTS sugars appeared to repress expression of the glucitol PTS in the wild-type strain; interestingly, mannitol repressed most strongly, fructose least, and glucose and N-acetylglucosamine repressed to a similar intermediate extent. Addition of cAMP did not relieve repression by glucose or mannitol. In strain SB1744 (Saier et al., 1976), which has a defective mannitol PTS, mannitol did not strongly repress expression of the glucitol PTS. Growth of wild-type strain SB3507 in the presence of PTS sugars other than glucitol generally resulted in a level of the membrane component EII(BC)\textsuperscript{Gut} about twice that of membranes from cells grown on unsupplemented LB, while glucitol induced about 20×. Results were similar for LJ705, although repression by fructose appeared to be stronger than in the wild-type strain. Fructose, normally a Class B, or weakly repressing, PTS sugar, may be a Class A, or strongly repressing, PTS sugar in *fruR* (cra) strains (Crasnier-Mednansky et al., 1997).
Repression of the gut operon by mannitol in wild-type S. typhimurium

To investigate these observations, further growth studies were conducted. When wild-type strain SB3507 was grown on a mixture of 0.02% mannitol and 0.2% glucitol, an obvious lag phase was observed between growth on mannitol and resumption of growth on glucitol (Fig. 3). Such a marked lag was not observed in the presence of 0.02% glucose, fructose or N-acetylglucosamine and 0.2% glucitol, and is consistent with strong repression of the glucitol operon in the presence of mannitol, and with early studies reporting diauxic growth of E. coli on mannitol and glucitol (Lengeler & Lin, 1972).

Conclusions

A mutant strain (LJ705) of S. typhimurium in which the normally fructose-inducible (and, in the case of EI$_{Fru}$, normally cryptic) PTS proteins FPr (DTP) and EI$_{Fru}$ must substitute for the general PTS proteins HPr and EI showed poor to negligible growth on the PTS sugar glucitol as its sole carbon source. Since addition of cAMP to the medium allowed LJ705 to grow on glucitol, we hypothesized that low cAMP, which could result from underphosphorylation of the mediator of catabolite repression/inducer exclusion, EI$_{G}$, could be responsible. Determination of the levels of the glucitol PTS protein EI$_{I}(BC)$G$_{out}$ in membranes from wild-type strains and LJ705 supported the hypothesis; induced levels in the mutant were consistently lower (50–70%) than in the wild-type strains, and levels in the mutant grown with glucitol and cAMP in the medium approached those in the wild-type grown with glucitol but without cAMP.

Another possible explanation for the failure of LJ705 to grow on glucitol could be that FPr works too poorly with the glucitol-specific PTS proteins to support growth. To investigate this possibility, mutant strain SB1475, which lacks active HPr and has low EI activity but is wild-type with respect to the fructose-specific PTS, i.e., does not express EI$_{Fru}$ at all and does not express FPr (DTP) constitutively, was grown on a mixture of fructose and glucitol. This mutant was able to continue growing on the mixture to a density much higher than on fructose alone, suggesting that it must have been able to take up glucitol using FPr. Thus it seems unlikely that poor interaction of FPr with the glucitol PTS is the sole explanation for the poor growth of LJ705 on glucitol.

Membranes from SB1475 contained maximal levels of EI$_{I}(BC)$G$_{out}$ only when the strain was grown in the presence of both glucitol and fructose. Surprisingly, this maximal level was similar to that of LJ705 rather than to that of the wild-type strains. It should also be noted that SB1475 has been reported to produce levels of cAMP about 50% those of wild-type strains (Feldheim et al., 1990). Thus it is not clear why this strain was able to grow on glucitol, apparently using FPr, while LJ705 was not. Addition of cAMP to the medium did not strongly affect growth of SB1475 on a mixture of fructose and glucitol. Possibly relatively poor interaction of FPr and EI$_{G}$ is a contributing factor to the poor growth of LJ705 on glucitol, in addition to low EI$_{I}(BC)$G$_{out}$. It has been reported that increasing cAMP may lead to increased levels of fructose PTS proteins even in fruR strains (Crasnier-Mednansky et al., 1997). Growing LJ705 in the presence of cAMP could thus increase the levels of both FPr and the glucitol PTS components, as well as EI$_{Fru}$, and a combination of these effects may allow growth on glucitol.

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