Glycollate Inhibition of Growth of *Pseudomonas aeruginosa* on Lactate Medium

By P. R. BROWN* AND RENÉE TATA

Department of Biochemistry, King's College London, Strand, London WC2R 2LS, UK

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Glycollate inhibited growth of *Pseudomonas aeruginosa* in media containing either pyruvate or lactate as carbon sources. Glycollamide, but not glyoxylate, showed similar effects. Spontaneous mutants (L/G strains) were isolated that were able to grow on lactate medium in the presence of glycollate: their growth in pyruvate medium was still inhibited by glycollate. Synthesis of membrane-bound NAD*-independent d(−)- and l(+)-lactate dehydrogenases (iLDHs) was inducible by D- or L-lactate in the parent strain but was constitutive in the L/G strains. Glycollate inhibited induction of the synthesis of iLDHs in the parent strain growing in succinate medium but had no effect under the same conditions on strain L/G1. Glycollate was a competitive inhibitor of l(+)-iLDH ($K_i = 11$ mM). No differences were found in the kinetic properties of l(+)-iLDH in cell-free extracts from strain L/G1 and the parent organism. Glycollate appears to inhibit growth on lactate medium predominantly through prevention of lactate induction of iLDH synthesis.

INTRODUCTION

The metabolism of DL-lactate by *Pseudomonas aeruginosa* (Kemp, 1972), *Pseudomonas citronellis* (O'Brien, 1977a) and *Pseudomonas putida* (O'Brien, 1977b) seems to depend on the induction of NAD*-independent l(+) and d(−)-lactate dehydrogenases (iLDHs) (EC 1.1.1.27 and EC 1.1.1.28, respectively) (for a review of LDHs see Garvie, 1980) that catalyse the conversion of lactate to pyruvate. Kemp (1972) showed that in *P. aeruginosa* these enzymes were membrane-bound and interacted with the electron transport chain. The iLDHs of *P. citronellis* (O'Brien, 1977a; O'Brien and Taylor, 1977) and *P. putida* (O'Brien, 1977b) appear to be similar to those in *P. aeruginosa* although the pattern of induction differs slightly for the different species. In *P. putida* and *P. aeruginosa* either D-lactate or L-lactate induces synthesis of both enzymes but, in *P. citronellis*, L-lactate induces both iLDHs, whereas D-lactate induces only d(−)-iLDH.

With lactate as carbon source no growth of *P. aeruginosa* occurs if glycollamide is used as a nitrogen source though the organism synthesizes an aliphatic amidase catalysing the hydrolysis of glycollamide to give ammonia and glycollate (Brown & Tata, 1987; this paper). In a taxonomic study of the aerobic pseudomonads, Stanier et al. (1966) reported that although a few *Pseudomonas* spp. are able to metabolize glycollate, *P. aeruginosa* strains cannot use it as a carbon source. It seemed possible therefore that glycollate prevented growth on lactate. In this paper we describe the results of experiments to confirm this and to elucidate the mechanism involved. A preliminary report of this work has appeared previously (Brown & Tata, 1981).

METHODS

**Bacterial strains.** The strains used (obtained from NCIB, Aberdeen, UK) were *Pseudomonas aeruginosa* PAC 1, described originally as strain 8602 by Kelly & Clarke (1962) and strain L10, a mutant derived from strain PAC 1

**Abbreviations:** iLDH, NAD*-independent lactate dehydrogenase; DCPIP, dichlorophenolindophenol; PMS, phenazine methosulphate.
that harbours mutations affecting the regulation of synthesis of amidase (Brown & Clarke, 1970; Smyth & Clarke, 1975).

Growth media. Minimal media, containing as carbon sources sodium DL-lactate (0.3%, w/v), sodium pyruvate (0.3%, w/v), sodium acetate (0.3%, w/v) or potassium glycollate (0.1%, w/v), were prepared by adding the appropriate solution, sterilized by filtration, to autoclaved minimal salts medium (Brammar & Clarke, 1964) containing 0.1% (w/v) (NH₄)₂SO₄ as nitrogen source. Succinate medium was prepared in a similar way but sodium succinate (0.3%, w/v) was autoclaved with the medium. Solid media contained 1% (w/v) Difco Noble Agar. Lactate/glycollate solid medium contained sodium DL-lactate (0.05%, w/v) and potassium glycollate (0.2%, w/v).

Growth inhibition studies. The effects of glycollate and glycollamide on bacterial growth in liquid media were investigated by growing bacteria in 250 ml culture medium at 37 °C to early exponential phase (OD₆₁₀ = 0.15) in 1 l conical flasks, and then transferring 45 ml portions to 250 ml flasks containing 5 ml sterile glycollate or glycollamide solution. Flasks were shaken in a Gallenkamp orbital shaker at 37 °C and samples removed at intervals to monitor growth by measuring OD₆₁₀ in a Unicam SP600 spectrophotometer.

Isolation of L/G mutants. A portion (5 ml) of overnight broth culture of strain L10 was centrifuged and the bacteria were washed in 5 ml dilution buffer (Brammar et al., 1967). After centrifugation, bacteria were resuspended in 5 ml dilution buffer and portions (0-1 ml) were spread on lactate/glycollate medium and incubated for 48 h at 37 °C. Mutant colonies were purified by restreaking on the isolation medium to obtain single colonies.

Enzyme assays. iLDHs were assayed using an adaptation of the method of Bennett et al. (1966). The assay mixture contained the following: dichlorophenolindophenol (DCPIP), 0.13 mM, phenazine methosulphate (PMS), 0.65 mM, and potassium cyanide, 0.4 mM, in 0.06 m-potassium phosphate buffer, pH 6.8. Bacterial extracts were added to 1 ml of reaction mixture in cuvettes equilibrated at 37 °C and the A₆₁₀ was monitored using a Unicam SP800 recording spectrophotometer for 2–3 min before starting the reaction by adding either lithium L-lactate (7.5 mM) or lithium D-lactate (7.5 mM). The reaction was monitored by measuring the rate of decrease in A₆₁₀. Enzyme activities are expressed as nmols DCPIP reduced min⁻¹ (mg protein)⁻¹ using a molar absorption coefficient ε₆₁₀ = 1.9 × 10⁴ l mol⁻¹ cm⁻¹ for DCPIP.

Preparation of cell-free extracts. Bacteria were harvested by centrifugation, washed in dilution buffer and then resuspended in 1/100th of the original medium volume of 100 mM-potassium phosphate buffer, pH 6.8, at 0 °C. Cells were broken by sonication using an MSE Mullard ultrasonic disintegrator (power output 50 W) for 2 min at 0 °C. Cell debris was removed by centrifugation at 22000 g in an MSE Angle 18 centrifuge at 4 °C for 15 min. Supernatant solutions were stored on ice before assay.

Protein estimations. These were done by the method of Lowry using bovine serum albumin as a standard.

Chemicals. Chemicals were of analytical grade. DCPIP and PMS were obtained from Sigma. Glycollic acid was recrystallized from ethanol twice and was neutralized with potassium carbonate before use in growth media. Glycollamide was prepared by the ammoniolysis of glycollic acid ethyl ester obtained from Eastman-Kodak (Schmuck, 1924) and was recrystallized twice from ethanol before use.

RESULTS

Growth inhibition by glycollate

Stanier et al. (1966) reported that glycollate is unable to support growth of P. aeruginosa. This was confirmed for strains PAC 1 and L10: neither strain produced visible colonies on solid minimal medium containing glycollate as sole carbon source even after 2–3 d incubation at 37 °C, and neither strain grew in liquid medium containing glycollate.

Plate tests showed that inclusion of 0.1% glycollate in pyruvate or lactate minimal media prevented growth of strains L10 and PAC 1. On succinate medium the presence of glycollate made no apparent difference to colony size after overnight incubation. Glyoxylate, the immediate product of glycollate metabolism in glycollate-utilizing Pseudomonas spp. (Kornberg & Gottlo, 1961) had no discernible effect at a concentration of 0.1% on growth on lactate or pyruvate media, which suggested that glycollate itself, rather than an intermediate in a blocked glycollate metabolic pathway, was the growth inhibitor.

Spontaneous mutants (L/G strains) were isolated from strain L10 on lactate/glycollate medium at a frequency of about 1 per 10⁶ cells plated. The growth properties of 10 of these mutants, picked at random, were examined on solid media. None grew on glycollate as sole carbon source but all were able to grow in the presence of glycollate plus lactate or glycollate plus succinate; none grew on pyruvate medium containing glycollate. The susceptibilities to glycollate inhibition of growth of the parent and one mutant strain, L/G1, were compared in more detail in liquid media (Fig. 1): 52 mM-glycollate strongly repressed growth of strain L10 in lactate medium but had no effect on strain L/G1.
Glycollate inhibition of *P. aeruginosa*

Fig. 1. Effects of glycollate on the growth of *P. aeruginosa* strains L10 (a) and L/G1 (b) in lactate medium. Arrows indicate the time at which glycollate was added (see Methods). Glycollate concentrations were as follows: 0 (○); 6.5 mM (▽); 13 mM (●); 52.5 mM (▲).

Fig. 2. Effects of glycollamide on the growth of *P. aeruginosa* strains L10 (a) and L/G1 (b) in lactate medium. Glycollamide was added (see Methods) at the time indicated by the arrows. Concentrations of glycollamide were as follows: 0 (○); 13.3 mM (▽); 33 mM (▲).

To examine the possibility that the mutation in strain L/G1 reduced the susceptibility to glycollate inhibition by preventing glycollate uptake, the effects of glycollamide on growth of the parent and mutant strains in lactate medium were compared. Farin (1977) concluded that *P. aeruginosa* PAC 1 is freely permeable to aliphatic amides and that amide uptake depends on the activity of an aliphatic amidase that catalyses the hydrolysis of the amide and includes glycollamide within its range of substrates (Kelly & Clarke, 1962). Glycollamide at the same concentration had a more pronounced effect than glycollate on strain L10. The mutant L/G1 under the same growth conditions was less susceptible to inhibition by glycollamide although some residual susceptibility remained: 33 mM-glycollamide completely suppressed growth of strain L10 and initially slowed growth of strain L/G1 (Fig. 2). Parent and mutant strains had the same level of amidase activity (data not shown) so the observation indicated that the relative resistance of the mutant strain to glycollate inhibition was not due to impermeability to glycollate.

In pyruvate medium no differences were observed between strains L10 and L/G1 in their susceptibilities to growth inhibition either by glycollate or glycollamide. Concentrations of glycollamide and glycollate that suppressed growth completely (8.3 and 26 mM respectively) were lower than those required to prevent growth of strain L10 in lactate medium. No effects on
Table 1. *iLDH* activities in *P. aeruginosa* strains L10 and L/G1 grown in various media

Activities are expressed as nmol lactate oxidized min⁻¹ (mg protein)⁻¹ with DCPIP as electron acceptor. Assays were done on exponential phase cultures except on those grown in succinate and acetate media when stationary phase cells were used. Activities represent means of duplicate assays done in two separate experiments. ND, Not determined.

<table>
<thead>
<tr>
<th>Carbon source for growth</th>
<th>Strain L10</th>
<th></th>
<th>Strain L/G1</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>L(+)iLDH</td>
<td>D(−)-iLDH</td>
<td>L(+)iLDH</td>
<td>D(−)-iLDH</td>
</tr>
<tr>
<td>Succinate</td>
<td>4</td>
<td>ND</td>
<td>96</td>
<td>ND</td>
</tr>
<tr>
<td>Acetate</td>
<td>9</td>
<td>ND</td>
<td>368</td>
<td>ND</td>
</tr>
<tr>
<td>Pyruvate</td>
<td>9</td>
<td>0</td>
<td>211</td>
<td>224</td>
</tr>
<tr>
<td>DL-Lactate</td>
<td>210</td>
<td>77</td>
<td>480</td>
<td>252</td>
</tr>
<tr>
<td>L-Lactate</td>
<td>310</td>
<td>207</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>

growth in succinate medium were observed for either strain with glycollate concentrations up to 19.7 mM and glyoxylate (16 mM) had no effect on growth in lactate medium confirming the observations made on solid media.

**Synthesis of iLDHs**

*iLDH* activities were measured in extracts of strain L10 grown in a variety of media. *iLDH* activity was found in cells grown in the presence of lactate but only very low levels were detectable in the absence of lactate (Table 1). D-Lactate (3 mM) and L-lactate (1-13 mM) were tested separately as inducers of *iLDH* synthesis in cells of strain L10 growing in succinate medium: both resulted in synthesis of L(+)-*iLDH* and D(−)-*iLDH*. Variations in the concentration of L-lactate had no effect on the level of *iLDH* (data not shown).

In striking contrast to the parent strain, synthesis of *iLDHs* in all the L/G mutants occurred both in the presence and in the absence of lactate. Data are shown in Table 1 only for strain L/G1. For several, though not all, of the L/G strains tested, *iLDH* activities of cells grown in lactate medium were higher than those for strain L/G1. *iLDH* activities were lowest in cells grown in succinate medium but were still at least 20-fold higher in the L/G strains than in strain L10.

The apparently constitutive synthesis of both lactate dehydrogenases in the L/G mutant strains suggested that glycollate might normally exert its inhibitory effect on growth on lactate medium by repressing induction of lactate dehydrogenases. To test this possibility, the effect of glycollate on induction by lactate of *iLDHs* in strains L10 and L/G1 growing in succinate medium was tested. Succinate was chosen because glycollate exhibited no detectable growth inhibitory effects in succinate medium. The results, given in Table 2, showed that for strain L10, as the glycollate concentration in the medium was increased so the level of *iLDHs* induced by lactate was reduced. In strain L/G1, in contrast, glycollate at a concentration that caused complete inhibition of induction of *iLDHs* in the parent showed no effect on *iLDH* synthesis.

**Inhibition of iLDHs by glycollate**

The structural similarities between lactate and glycollate suggested that glycollate might act as an inhibitor of *iLDH* activity; this could therefore be a contributory factor in growth inhibition by glycollate. Preliminary experiments using cell extracts as sources of enzyme showed that glycollate inhibited both D(−)-*iLDH* and L(+)-*iLDH*. The effects of varying the concentration of glycollate on L(+)-*iLDH* activity at different fixed concentrations of L-lactate were investigated and values for $K_i$, determined using a Dixon plot (not shown), were 12.5 mM for strain L10 and 11 mM for strain L/G1. Replots (not shown) of the data by the method of Cornish-Bowden (1974) gave parallel lines, indicating competitive inhibition. Determinations of the $K_m$ of L(+)-*iLDH* for L-lactate using cell free extracts gave linear double reciprocal plots and values of 0.25 mM for strain L10 and 0.27 mM for strain L/G1.
**Table 2. Effect of glycollate on induction of L(+)−iLDH synthesis in P. aeruginosa cells growing in succinate medium**

Cultures were grown to early exponential phase in succinate medium, then divided into four flasks containing glycollate and/or L-lactate at the concentrations shown. Cells were harvested at an OD$_{600}$ of about 0.5 and assayed for L(+)−iLDH activity. Activity is expressed as nmol L-lactate oxidized min$^{-1}$ (mg protein)$^{-1}$ with DCPIP as electron acceptor.

<table>
<thead>
<tr>
<th>Strain</th>
<th>L-Lactate concn (mM)</th>
<th>Glycollate concn (mM)</th>
<th>L(+)−iLDH activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>L10</td>
<td>1.02</td>
<td>0</td>
<td>68.7</td>
</tr>
<tr>
<td></td>
<td>0.89</td>
<td>1.76</td>
<td>21</td>
</tr>
<tr>
<td></td>
<td>0.89</td>
<td>4.42</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>0.89</td>
<td>8.84</td>
<td>0</td>
</tr>
<tr>
<td>L/G1</td>
<td>0</td>
<td>0</td>
<td>154</td>
</tr>
<tr>
<td></td>
<td>0</td>
<td>8.84</td>
<td>144</td>
</tr>
<tr>
<td></td>
<td>0.89</td>
<td>8.84</td>
<td>123</td>
</tr>
<tr>
<td></td>
<td>0.89</td>
<td>0</td>
<td>160</td>
</tr>
</tbody>
</table>

**DISCUSSION**

The properties of the L/G mutants indicate that a reduction in the sensitivity of *P. aeruginosa* to inhibition of growth by glycollate in lactate medium is associated with mutations resulting in constitutive synthesis of iLDHs. This suggested that the primary action of glycollate is through repression of iLDH induction, and evidence is presented showing that glycollate normally inhibits induction of iLDH synthesis by lactate. The mechanism of this proposed inhibition is not clear. It could be caused through competition of glycollate with lactate for entry into the cell. Matin & Konings (1973) showed that uptake rates of lactate into cells of a freshwater *Pseudomonas* (species unidentified) were inhibited nearly 60% by 1 mM-glycollate. However, it seems unlikely that iLDHs themselves are directly involved in lactate uptake since entry of DL-lactate into membrane vesicles is not dependent on iLDH (Matin & Konings, 1973), and O’Brien (1977b) concluded that uptake of D- and L-lactic acids was independent of iLDH activity in *P. putida*. It is feasible that synthesis of a lactate permease could be coordinately regulated with transcription of iLDH genes and constitutive synthesis would allow lactate entry in the presence of glycollate by providing more binding sites. Alternatively, glycollate inhibition of iLDH induction may be an intracellular event, glycollate competing with lactate for binding to a hypothetical regulatory protein that controls transcription of iLDH genes. We have screened several hundred L/G mutants to identify any that are able to grow on the selection medium at 40 °C but not at 25 °C, but without success (C. L. Crossley & P. R. Brown, unpublished), which argues against the presence of a regulatory protein requiring lactate binding to activate transcription.

The inhibitory effects of glycollamide on growth with lactate depend on its conversion to glycollate since amidase-negative mutants show no such susceptibility (Brown & Tata, 1987), but this does not necessarily imply that glycollate inhibition is exerted intracellularly since glycollate might be exported after formation from glycollamide then taken up by an active transport process as Farin (1977) has suggested for acetate formed from acetamide. Using [1$^{14}$C]glycollate we have found that the label is taken up into the cell by an azide-sensitive process (A. Allsop & P. R. Brown, unpublished).

No large differences were found between the kinetic parameters of the iLDHs of parent and mutant strains, suggesting that the differences in enzyme activities are unlikely to be due to structural changes in the enzymes. The inhibitory action of glycollate on enzyme activity is relatively weak, reflected in a high $K_v$ value, but could contribute to growth inhibition depending on the relative intracellular concentrations of glycollate and lactate.

The continued sensitivity of L/G mutants to glycollate inhibition of growth on pyruvate medium seems to indicate an effect of glycollate on transport of pyruvate into the cell, unless pyruvate formed by iLDH activity towards lactate is somehow differentiated by, for example, channelling or chemical modification, from pyruvate entering the cell.
The induction of both iLDHs by either D- or L-lactate suggests that the enzymes are coordinately regulated although the possibility of a racemase interconverting the two isomers has not been eliminated. Since regulation of both enzymes was similarly affected in the L/G strains [and in L/G mutants isolated on L-lactate/glycollate medium (P. R. Brown, unpublished), this too strongly indicates that the synthesis of the enzymes is coordinately controlled. The high frequency at which L/G mutants arise makes unlikely the possibility that each is the result of more than one mutation.

REFERENCES


