Iron Transport in *Mycobacterium smegmatis*:
Uptake of Iron from Ferriexochelin

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Exochelins are a group of extracellular iron chelators produced by mycobacteria. Iron uptake by washed suspensions of iron-deficiently grown *Mycobacterium smegmatis* from $^{55}$Fe(III)-exochelin fractions (at about 1 $\mu$M) was greatest from the fractions containing the compounds that naturally predominate in culture filtrates. Uptake from the major fraction, as well as from combined exochelins, had a $K_m$ of about 6 $\mu$M and was unaffected by the presence of a large excess of desferriexochelin; it was inhibited by more than 90% by electron transport inhibitors, uncouplers of oxidative phosphorylation, thiol reagents and by anaerobiosis and low temperature. Uptake of iron from $^{55}$Fe-salicylate, which is mediated by mycobactin, was insensitive to these inhibitors and a 10-fold excess of ferric salicylate did not inhibit $^{55}$Fe-exochelin uptake. Thus mycobactin is probably not involved in the transport of iron from ferriexochelin at physiological concentrations. The rate of uptake of iron from $^{55}$Fe-exochelin into iron-sufficiently grown cells, which contain less than 0.05% of the concentration of mycobactin found in iron-deficiently grown cells, was only slightly lower than the rate of uptake into iron-deficiently grown cells.

Uptake of ferri$[^3]$H$\varepsilon$xochelin, which could only be carried out at high and probably non-physiological concentrations (about 60 $\mu$M), was also extremely sensitive to metabolic inhibitors suggesting that the whole complex was being transported. At these high external concentrations of ferriexochelin a second, non-saturable, inhibitor-insensitive iron uptake process occurred. This process was inhibited in iron-deficiently grown cells by ferric salicylate and may therefore involve mycobactin. A similar but not identical second system which was not sensitive to ferric salicylate was found in iron-sufficiently grown cells; this might indicate yet another pathway of iron uptake from ferriexochelin.

INTRODUCTION

Ratledge & Marshall (1972) proposed a mechanism for the uptake of iron into *Mycobacterium smegmatis* which involved the mediation of mycobactin (Snow, 1970), a lipid-soluble iron-binding agent, in shuttling iron across the boundary layers of the cell. The loading of iron on to mycobactin was presumed to be via salicylic acid, a compound which had previously been isolated from culture filtrates of iron-deficiently grown *M. smegmatis* (Ratledge & Winder, 1962). As salicylate was subsequently shown not to fulfil this role in the presence of phosphate ions, due to the formation of insoluble ferric phosphate (Ratledge et al., 1974), we searched for, and found, an extracellular iron chelator which would hold iron in solution in the presence of competing ions. This chelator was subsequently shown to consist of a group of compounds which were termed exochelins (Macham & Ratledge, 1975; Macham et al., 1975). Exochelins from *M. smegmatis* can solubilize iron from ferric phosphate and ferritin and mediate the uptake of iron by *M. smegmatis* (Macham et al., 1977). We have now investigated this process of iron transport from ferriexochelin. A preliminary communication of part of this work has appeared elsewhere (Stephenson & Ratledge, 1978).
METHODS

Abbreviations. CCCP, Carbonyl cyanide m-chlorophenylhydrazone; DNP, 2,4-dinitrophenol; NEM, N-ethylmaleimide.

Organism, growth and preparation of washed suspensions. Mycobacterium smegmatis NCIB 8548 was grown in 100 ml medium (Ratledge & Hall, 1971) in 250 ml conical flasks, with shaking, at 37 °C. For iron-deficient growth 0.05 μg Fe²⁺ ml⁻¹ was added to the medium; for iron-sufficient growth 2.0 μg Fe²⁺ ml⁻¹ was added. Cells were harvested after 3 d by filtering through Whatman GF/C (70 mm) glass-fibre filters and were washed with 0.85% (w/v) NaCl containing 1.0% (v/v) Tween 80. They were then resuspended in an identical solution at approximately 60 mg dry wt ml⁻¹ for iron-deficiently grown cells and 80 mg dry wt ml⁻¹ for iron-sufficiently grown cells and were stored on ice until needed. Rates of iron transport were stable over 3 h storage.

Exochelins. Exochelin MS was isolated and purified as described previously (Macham et al., 1977), as were the preparations of desferri- and ³²Fe-exochelin. Tritiated exochelin was prepared by adding β-[³H]-alanine (sp.act. 49 Ci mmol⁻¹) to iron-deficient growth medium at 1.02 nmol per 100 ml medium, at the time of inoculation. After 3 d growth, the exochelin was isolated as described previously for unlabelled exochelin and the purification was taken to the stage of preparation of the 'purified but unfractionated material' (Macham et al., 1977).

Preparation of ³²Fe-salicylate. To a solution of ³²FeCl₃ containing 0.254 μmol Fe (sp.act. 93-5 μCi μmol⁻¹), salicylic acid (8.5 μmol) was added. (This was the equivalent of an 11-fold excess of salicylate since salicylate and iron bind in the molar ratio of 3:1.) MgSO₄ was then added to give a final concentration of 1 mM, the solution was adjusted to pH 7-0 with 0.1 M-Tris and the volume was adjusted to 20 ml, so that the initial concentration of Fe(salicylate)₃ was 12.7 μM. The solution was filtered through a Whatman GF/A (25 mm) glass-fibre filter and used immediately for transport assays. Solutions were prepared daily.

Uptake of ³²Fe from ³²Fe-exochelin. Initial rates of iron uptake into washed suspensions of 3 d-old cultures of M. smegmatis were assayed in a solution containing 50 mM-KH₂PO₄/NaOH, pH 7.1 (2.45 ml) and water or other component (0.35 ml). This solution was incubated for 5 min in a 25 ml Erlenmeyer flask in a shaking waterbath at 25 °C. The washed suspension (0.35 ml, see above) was then added and the incubation was continued for a further 10 min before ³²Fe-exochelin (0.35 ml), pre-warmed to 25 °C, was added by syringe. Using an automatic pipette, samples (0.5 ml) were run into modified Millipore Swinnex-25 filter units holding a Whatman GF/A (25 mm) glass-fibre filter and 2 ml 50 mM-Na₂EDTA, pH 6.0, at room temperature. Suction was immediately applied to the filter disc to drain it completely. The filter was washed with a further 2 × 1 ml of EDTA solution, removed immediately, placed in a glass scintillation vial and dried in an oven at 108 °C. Soluene 100 (0.5 ml, Packard Instruments) was added to each vial before incubating at 60 °C for 5 h to digest the bacteria. Glacial acetic acid (50 μl) and scintillation fluid (10 ml) were added to each vial before counting (see below). Initial rates of uptake, expressed as nmol min⁻¹ (g dry wt)⁻¹, were measured in duplicate from four samples taken during the first 2 min after the addition of ³²Fe-exochelin.

Uptake of ³²Fe from ³²Fe-salicylate. Experiments were performed as described above for ³²Fe-exochelin except that the KH₂PO₄/NaOH buffer was replaced by 0.1 M-Tris/Cl⁻ buffer, pH 7.0.

Uptake of ³H-exochelin. Experiments were performed as described above for ³²Fe-exochelin except that initial rates were measured in duplicate from four samples taken during the first 15 min after the addition of ferri³H-exochelin. The washed filters were placed directly into 10 ml scintillation fluid (see below). Rates of ³H uptake were converted to rates of iron transport on the basis of the ferriexochelin complex contains equimolar concentrations of iron and exochelin.

Counting of radioactivity. (i) ³²Fe. Aqueous samples for the determination of specific activities were dried in glass scintillation vials and treated in an identical manner to the dried filters from transport assays (see above). Samples were counted in a scintillation fluid comprising 10% (w/v) naphthalene and 1% (w/v) 2,5-diphenyloxazole in toluene. (ii) ³H. Aqueous samples for the determination of specific activities and filters from transport assays were counted directly in a scintillation fluid comprising: xylene, 726 ml; Triton N-101, 264 ml; 2-(4'-tert-butylphenyl)-5-(4'-biphenylyl)-1,3,4-oxadiazole, 10 g.

RESULTS

Iron uptake from various exochelin fractions

The uptake of ³²Fe from unfractonated but purified ³²Fe-exochelin (1.25 μM) into washed suspensions of iron-deficiently grown M. smegmatis, held in 50 mM-KH₂PO₄/NaOH buffer, pH 7.1, reached a constant value after 12 min (Fig. 1). The amount of iron accumulated at
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Fig. 1. Uptake of $^{55}$Fe from ferriexochelin by iron-deficiently grown Mycobacterium smegmatis. Transport was assayed as described in Methods. Unfractionated $^{55}$Fe-exochelin was added at a final concentration of 1.25 $\mu$M.

the plateau represented about a 60-fold increase over the external iron concentration, assuming an intracellular volume of 2 ml (g dry wt)$^{-1}$ (a typical value for bacteria; see, for example, Midgley & Dawes, 1973).

Exochelin MS is a mixture of at least seven components which are separable by ion-exchange chromatography (Macham et al., 1977). When the ability of the different fractions to transport $^{56}$Fe (1.7 $\mu$M) into M. smegmatis was compared, fractions 1 and 2 combined had the same activity as the unfractionated exochelins [i.e. 34 nmol min$^{-1}$ (g dry wt)$^{-1}$] whereas fraction 3 had a slightly higher activity [46 nmol min$^{-1}$ (g dry wt)$^{-1}$] and fraction 4 a slightly lower activity [21 nmol min$^{-1}$ (g dry wt)$^{-1}$]. Fractions 5 and 6 were inactive or almost so [0 and 3-6 nmol min$^{-1}$ (g dry wt)$^{-1}$, respectively]. Thus the highest activity was shown by the major fraction 3 which constitutes about 75% of the total exochelin. Fraction 3A, obtained by further purification of fraction 3, exhibited an almost identical activity to fraction 3 itself. The relative inactivity of fractions 5 and 6 may indicate that they are breakdown products of the major exochelins.

Uptake in the presence of desferriexochelin

The concentration of ferriexochelin used in these experiments (1.7 $\mu$M) was equivalent to an exochelin concentration of about 1.3 $\mu$g ml$^{-1}$, assuming a molecular weight of 800. This is similar to the physiological concentrations of ferriexochelin which are found in the medium during growth of M. smegmatis under iron-deficient conditions. Although the amount of desferriexochelin isolated from spent growth medium is about 50 $\mu$g ml$^{-1}$, the amount of ferriexochelin present during growth will be limited by the supply of iron. As the total iron concentration (added plus residual) is about 0.1 $\mu$g ml$^{-1}$, ferriexochelin cannot exceed about 1.4 $\mu$g ml$^{-1}$.

There was no inhibition of iron uptake when $^{55}$Fe-exochelin (at 1.1 $\mu$M) was added to bacterial suspensions to which a large excess of desferriexochelin (50 $\mu$g ml$^{-1}$) had been added to mimic the probable growth conditions. Thus there was no competition between ferriexochelin and desferriexochelin for a possible receptor site.

Sensitivity of iron uptake to inhibitors

At these low, physiological, external ferriexochelin concentrations, $^{55}$Fe uptake into iron-deficiently grown cells was inhibited by more than 90% by the electron transport inhibitors NaN$_3$ and KCN, by the uncouplers CCCP and DNP, and by the thiol reagent HgCl$_2$ (Table 1). Uptake was also virtually eliminated by anaerobiosis and low temperature (4°C). Similar levels of inhibition occurred whether the purified fraction 3 (or 3A) or the unfrac-
Table 1. Effect of various potential inhibitors on iron transport from ferriexochelin, at low external concentrations, by Mycobacterium smegmatis

Transport was assayed as described in Methods. Iron-deficiently and iron-sufficiently grown cells were incubated with the potential inhibitors for 10 min before adding the various 55Fe-exochelin fractions (Unf, unfractionated exochelin) at the concentrations shown. The results were compiled from several experiments in which 100% activity with iron-deficiently grown cells was in the range 17 to 44 nmol min⁻¹ (g dry wt)⁻¹ for fraction 3A, 28 to 52 nmol min⁻¹ (g dry wt)⁻¹ for fraction 3 and 37 to 72 nmol min⁻¹ (g dry wt)⁻¹ for the unfractionated exochelin at the various concentrations used; 100% activity with the iron-sufficiently grown cells was in the range 18 to 39 nmol min⁻¹ (g dry wt)⁻¹.

<table>
<thead>
<tr>
<th>Exochelin fraction</th>
<th>Inhibitor</th>
<th>Concen (mM)</th>
<th>3A (0.36 μM)</th>
<th>3 (1.3 μM)</th>
<th>Unf. (1.3 μM)</th>
<th>Unf. (1.8 μM)</th>
<th>Unf. (1.8 μM)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>None</td>
<td>—</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>KCN</td>
<td>10</td>
<td>5</td>
<td>3</td>
<td>4</td>
<td>6</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>NaN₅₋₋</td>
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<td>4</td>
<td>3</td>
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<td></td>
<td>0.10</td>
<td>5</td>
<td>2</td>
<td>3</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>DNP</td>
<td>2</td>
<td>7</td>
<td>9</td>
<td>3</td>
<td>8</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>NEM</td>
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<td>49</td>
<td>56</td>
<td>51</td>
<td>52</td>
<td>58</td>
</tr>
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<td>3</td>
<td>7</td>
<td>4</td>
<td>11</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>N₂ atmosphere</td>
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<td>2</td>
<td>5</td>
<td>3</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>Temp. 4°C</td>
<td>—</td>
<td>0</td>
<td>2</td>
<td>0</td>
<td>4</td>
<td>3</td>
</tr>
</tbody>
</table>

Difficulties in producing large quantities of the purified fraction 3A precluded using it in large numbers of routine assays. When iron-sufficiently grown cells were used, initial rates of 55Fe uptake from ferriexochelin (1.8 μM) were only two- to threefold lower than those of the iron-deficiently grown cells and inhibition studies revealed essentially similar results with both types of organism (Table 1).

Investigation of the role of mycobactin

The inhibitor-sensitivity of iron uptake from 55Fe-salicylate was compared with that of iron uptake from ferriexochelin since the uptake of iron from 55Fe-salicylate by iron-deficiently grown M. smegmatis is mediated by mycobactin (Ratledge & Marshall, 1972). However, the uptake of 55Fe from 55Fe-salicylate (1.27 μM) in 0.1 M-Tris/HCl buffer, pH 7.0 (phosphate buffer cannot be used as insoluble ferric phosphate is quickly produced) was markedly different from iron uptake from ferriexochelin, being insensitive to KCN (10 mM), CCCP (0.1 mM) and HgCl₂ (0.1 mM) in both iron-deficiently and iron-sufficiently grown cells. (Initial rates of 55Fe uptake from 55Fe-salicylate in iron-sufficiently grown cells were about threefold lower than those in the iron-deficiently grown cells.) These results suggest that different mechanisms operate for the uptake of iron from ferric salicylate and ferriexochelin. Supporting this suggestion, no inhibition of 55Fe uptake from 55Fe-exochelin (1.25 μM) by iron-deficiently grown cells was observed when unlabelled ferric salicylate (12.5 μM) was added, as a possible competitor for the receptor site, 15 s prior to adding the labelled substrate.
Iron transport in *M. smegmatis*

Fig. 2. Uptake of [3H]exochelin by *Mycobacterium smegmatis* grown iron-deficiently (●) or iron-sufficiently (▲). Transport was assayed as described in Methods. Unfractionated ferri[3H]exochelin was added at a final concentration of 71 μM.

Fig. 3. Uptake of 55Fe by iron-deficiently grown *Mycobacterium smegmatis* in response to the external ferriexochelin concentration, in the absence (●) or presence (▲) of NaN₃. Transport was assayed as described in Methods. Cells were preincubated with NaN₃ (30 mM) for 10 min prior to the addition of unfractionated 55Fe-exochelin.

Thus we concluded that iron uptake from ferriexochelin at physiological concentrations was a process in some way dependent on metabolism, as evidenced by its sensitivity to a wide range of metabolic inhibitors and anaerobiosis, which did not involve mediation by the lipid-soluble iron-binding compound mycobactin.

Studies on the uptake of ferri[3H]exochelin

Iron uptake from ferriexochelin could involve uptake of the entire complex or release of iron from the complex without necessitating uptake of the chelator into the cell. To investigate the uptake process, attempts were made to prepare radioisotopically labelled exochelins by growing *M. smegmatis* in the presence of a suitably labelled precursor of exochelin. Exochelin fraction 3A contains moieties of lysine, β-alanine and threonine (L. P. Macham, M. C. Stephenson & C. Ratledge, unpublished work). We therefore initially tried adding L-[1-14C]lysine (sp.act. 0.1 mCi mmol⁻¹) at 1 pmol ml⁻¹ to cultures of iron-deficiently growing *M. smegmatis*, but the recovered exochelin had too low a specific activity [about 150 c.p.m. (μg exochelin)⁻¹] to allow useful experiments to be conducted. Addition of β-[3-3H]alanine (sp.act. 49 mCi mmol⁻¹) to the growth medium at 10-2 pmol ml⁻¹ produced a purified but unfractionated exochelin with a specific activity of 650 c.p.m. (μg exochelin)⁻¹, showing that considerable dilution with exogenous β-alanine had occurred. This exochelin preparation, though of lower specific activity than had been hoped for, was sufficiently active to enable assays of exochelin uptake to be conducted provided that the external exochelin concentration was increased to about 60 μM. At such high external concentrations [3H]exochelin was accumulated from ferri[3H]exochelin by both iron-deficiently and iron-sufficiently grown *M. smegmatis* (Fig. 2), the rate of iron uptake by the former being about three times that by the latter, in keeping with previous data for 55Fe uptake from ferri-exochelin (see Table 1). However, a comparison of the absolute rates of 3H uptake using ferri[3H]exochelin and 55Fe uptake using 55Fe-exochelin (both at 62 μM) revealed that rates of 55Fe uptake were 10- to 20-fold higher than rates of 3H uptake. Thus the amount of iron transported as the ferriexochelin complex accounted for only 5 to 10% of the total iron accumulated with both iron-deficiently and iron-sufficiently grown cells. An explanation for this discrepancy became apparent from the work described below.
Table 2. Effect of various potential inhibitors on iron transport from ferriexochelin, at high external concentrations, by Mycobacterium smegmatis

The assays were performed as described in the legend to Table 1. The unfractionated $^{55}$Fe-exochelin concentration was 50 $\mu$M in both sets of experiments and 100% activity was in the range 105 to 168 nmol min$^{-1}$ (g dry wt)$^{-1}$ for iron-deficiently grown cells and 48 to 89 nmol min$^{-1}$ (g dry wt)$^{-1}$ for iron-sufficiently grown cells.

<table>
<thead>
<tr>
<th>Inhibitor</th>
<th>Concen (mM)</th>
<th>Relative initial rate of uptake</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>Iron-deficient</td>
</tr>
<tr>
<td>None</td>
<td>—</td>
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</tr>
<tr>
<td>KCN</td>
<td>10</td>
<td>42</td>
</tr>
<tr>
<td>NaN$_4$</td>
<td>30</td>
<td>68</td>
</tr>
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<td>CCCP</td>
<td>0.1</td>
<td>84</td>
</tr>
<tr>
<td>DNP</td>
<td>2</td>
<td>73</td>
</tr>
<tr>
<td>HgCl$_2$</td>
<td>0.1</td>
<td>84</td>
</tr>
</tbody>
</table>

At such high concentrations of ferri[$^3$H]exochelin, uptake of $^3$H was sensitive to the compounds previously shown to inhibit $^{55}$Fe uptake (see Table 1); i.e. NaN$_4$ (30 mM), KCN (10 mM), CCCP (0.1 mM), DNP (2 mM) and HgCl$_2$ (0.1 mM) all produced immeasurably low rates of $^3$H uptake which were therefore less than 5% of the uninhibited activity. However, the external ferriexochelin concentration had been raised approximately 80-fold to assay $[^3$H]exochelin uptake, and when the inhibitor studies of $^{55}$Fe uptake from ferriexochelin were repeated at similar external concentrations, a markedly lower sensitivity to the various inhibitors was observed (Table 2).

The data therefore suggest the operation of at least two processes for iron transport from ferriexochelin. The simplest interpretation suggests that one process operates at low external ferriexochelin concentrations, is inhibitor-sensitive and involves the uptake of the ferriexochelin complex. A second process predominates at high external ferriexochelin concentrations and this is inhibitor-insensitive and does not involve the uptake of exochelin.

Evidence for alternative routes for iron uptake

To seek further evidence for the existence of more than one component for iron transport, the initial rate of $^{55}$Fe uptake from ferriexochelin was measured in response to increasing external substrate concentration from 1.25 to 125 $\mu$M. The results (Fig. 3) indicate that probably two components are operating for iron transport from ferriexochelin by iron-deficiently grown cells. One process is saturable (at about 12.5 $\mu$M-ferriexochelin) and the other fails to saturate within the range of concentration used. In the presence of NaN$_4$ (30 mM) the saturable component was eliminated (Fig. 3). In addition, at high external ferriexochelin concentration (63.1 $\mu$M) the uptake of $^{55}$Fe was inhibited by 80% when a 10-fold excess of ferric salicylate was added 15 s before the $^{55}$Fe-exochelin. These results, which are in contrast to those obtained using 1.25 $\mu$M-$^{55}$Fe-exochelin, suggest that ferric salicylate interferes with the second, inhibitor-insensitive component for iron transport from ferriexochelin by iron-deficiently grown cells.

This last result suggested that mycobactin may be involved in the second component for iron uptake. Iron-sufficiently grown cells, which contain only minimal amounts of mycobactin (see Discussion), were similar to iron-deficient cells in possessing a second, inhibitor-insensitive component for iron uptake at high concentrations of $^{55}$Fe-exochelin (Table 2). However, unlike iron-deficiently grown cells, they were not inhibited in the uptake of $^{55}$Fe from $^{55}$Fe-exochelin (63 $\mu$M) when unlabelled ferric salicylate was added in 10-fold excess even though $^{55}$Fe-salicylate is transported at such concentrations by iron-sufficiently grown cells. These results suggest that mycobactin was not involved in the second process.
Iron transport in *M. smegmatis*

![Diagram](image)

**Fig. 4.** Mechanisms of iron transport by *Mycobacterium smegmatis*: Sal, salicylate; Exo, exochelin; M, mycobactin.

(A) Salicylate–mycobactin system operating in iron-deficiently grown *M. smegmatis*, previously characterized by Ratledge & Marshall (1972).

(B) Ferriexochelin transport system operating in both iron-deficiently and iron-sufficiently grown *M. smegmatis* at low, physiological ferriexochelin concentrations.

(C) Iron transport from ferriexochelin at high external concentrations, probably mediated by mycobactin in iron-deficiently grown *M. smegmatis*.

Thus, the second component for iron transport operating at high concentrations of ferriexochelin has different characteristics in iron-sufficiently grown cells from those in iron-deficiently grown cells. Both types of cell, however, apparently share a common process for the uptake of iron from ferriexochelin at low external concentrations.

**DISCUSSION**

Iron uptake into *M. smegmatis* is a more complex process than hitherto believed and a number of separate routes may co-exist, as they do in *Escherichia coli* (Braun, 1978). Unlike *E. coli*, which synthesizes only one iron-chelating agent (enterochelin) and makes use of the presentation of other naturally occurring forms of iron for its alternative pathways, *M. smegmatis* and other mycobacteria produce three iron-chelating agents – salicylic acid, exochelin and mycobactin. Only mycobactin occurs exclusively in the cell (Ratledge & Marshall, 1972). The previously described route of iron uptake via mycobactin (see Fig. 4A, and Ratledge & Marshall, 1972) depended upon ‘soluble’ iron, in the form of ferric salicylate, being presented to cells. Though other aromatic acids can substitute for salicylate and also transfer iron to mycobactin, neither ferric citrate, ferric EDTA nor ferric oxalate
can do so (Ratledge et al., 1974). Iron from ferrous sulphate is also rapidly taken up by mycobactin (oxidation may occur prior to or upon complex formation) when freshly prepared solutions are added to established iron-deficient cultures of M. smegmatis, that is, before polymeric forms of iron have been able to form (McCready & Ratledge, 1978). However, since inhibition studies suggest that mycobactin is not involved in the uptake of iron from ferriexochelin at low concentrations, a novel and independent route for iron uptake is now recognized.

This novel iron transport system was seen in both iron-deficiently and iron-sufficiently grown cells despite the fact that exochelin synthesis is repressed during iron-sufficient growth. Although culture filtrates from iron-sufficiently grown cells have a small but detectable capacity to solubilize iron (Macham & Ratledge, 1975), the amount of exochelin present has not been measured. Iron acquisition is, however, necessary no matter what the concentration of extracellular iron and thus the exochelin-mediated system is deduced to be prevalent at all concentrations of extracellular iron.

As a result of investigating the inhibitor-sensitivity of both $^{59}$Fe and $[^{3}H]$exochelin uptake, we propose that iron uptake from ferriexochelin at low external concentrations involves simultaneous uptake of both ligand and metal. In this respect, the uptake would be similar to the uptake of other ferric siderophores in other bacteria, for example, ferrischizokinen transport in Bacillus megaterium, ferriaerobactin in Aerobacter (Klebsiella) aerogenes 62/1, and ferrienterochelin in Escherichia coli (Arceneaux et al., 1973; Frost & Rosenberg, 1973). (This could be confirmed if $[^{14}C]$- or $[^{3}H]$exochelin of high specific activity could be prepared as this would permit double-labelling experiments at low ferriexochelin concentrations.) The mechanism of uptake is also similar to other ferric siderophore transport systems in showing sensitivity to a wide range of metabolic inhibitors. The sensitivity to uncouplers and electron transport inhibitors is similar to that shown for ferrienterochelin transport in E. coli (Pugsley & Reeves, 1977), suggesting that iron transport is dependent upon the creation and maintenance of an energized membrane state. Lack of uptake at 4°C may also reflect the need for metabolism. Similarly, under anaerobic conditions, when respiration and oxidative phosphorylation are prevented, no uptake is observed. The ability of thiol reagents to block uptake suggests the involvement of $-SH$ groups in the overall process, although in view of the lack of specificity of these compounds it is not possible to define their site of action. However, Ernst & Winkelman (1977) have reported that the binding sites for ferricoprogen on the surface of intact cells of Neurospora crassa can be labelled to a high degree by $[^{14}C]NEM$. If the similarity between the ferriexochelin system and other siderophore uptake systems is extended, one would expect that there will also be specific binding proteins on the mycobacterial cell surface as for N. crassa, E. coli (Hancock et al., 1976; Braun, 1978) and Salmonella typhimurium (Luckey & Neilands, 1976; Neilands, 1977).

The uptake of iron from ferriexochelin at low external concentrations is considered to be characteristic of the in vivo physiological situation, and occurs via a high-affinity system, with a $K_m$ of approximately 6 $\mu$M. (This value, obtained from the data of Fig. 3, is only approximate due to the difficulty in estimating the saturating ferriexochelin concentration in the presence of interference by the non-saturable, inhibitor-insensitive system.) This $K_m$ is similar to that reported in other systems (Lankford, 1973; Wiebe & Winkelman, 1975). The mechanism of iron release and the fate of the desferriexochelin are unknown. The intact ligand may be excreted, as occurs with schizokinen in B. megaterium (Arceneaux et al., 1973), aerobactin in A. aerogenes (Arceneaux et al., 1973) and ferrichrome in E. coli (Leong & Neilands, 1976). Alternatively, exochelin could be hydrolysed, as occurs with enterochelin and the fusarinsines (O’Brien et al., 1971; Emery, 1976). In these cases the degradation products can be detected in the medium. If this occurs in M. smegmatis it could explain the number of exochelins found in the culture filtrates, some of which are not active in iron transport though they do still bind iron. A scheme for this high-affinity, inhibitor-sensitive process of iron uptake is depicted in Fig. 4B.
Iron transport in M. smegmatis

A further, different process for iron uptake from ferriexochelin at high, and probably non-physiological, concentrations was recognized by its insensitivity to the various inhibitors. As ferri[3H]exochelin uptake at the same high concentration was extremely sensitive to these metabolic inhibitors the process cannot involve simultaneous uptake of metal and ligand. The inhibition of iron uptake by ferric salicylate under these conditions, using iron-deficiently grown cells, suggests that mycobactin may be the receptor for the transported iron (see Fig. 4C). This inhibition was not observed when iron-sufficiently grown cells were used, suggesting the operation of two different systems. However, since iron-sufficiently grown cells contain at most only 0.05% of the mycobactin of iron-deficiently grown cells (Ratledge & Marshall, 1972), it is possible that the location and accessibility of this small concentration of mycobactin prevents observation of the salicylate inhibition phenomenon, especially if a binding site for ferriexochelin specifically facilitates exchange of iron between exochelin and mycobactin. It must, however, be stressed that this second exochelin system, for which parallels exist in other organisms, viz. uptake of ferric citrate by Esherichia coli (Winkelmann & Zahner, 1973), may be non-physiological as it is unlikely that very high concentrations of ferriexochelin could suddenly arise in vivo.

Two major questions arise from this present work: the first, and more obvious, is what is the detailed mechanism of iron uptake from exochelin? This could be elucidated with a suitably isotopically labelled substrate. The second, more enigmatic, is what is the role of mycobactin if the physiological iron-sequestering agent, exochelin, does not transfer its iron to the cell via mycobactin?

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REFERENCES


