Aromatic components of two ferric enterobactin binding sites in *Escherichia coli* FepA

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Summary

Ferric enterobactin is a catecholate siderophore that binds with high affinity ($K_d \approx 10^{-10}$ M) to the *Escherichia coli* outer membrane protein FepA. We studied the involvement of aromatic amino acids in its uptake by determining the binding affinities, kinetics and transport properties of site-directed mutants. We replaced seven aromatic residues (Y260, Y272, Y285, Y289, W297, Y309 and F329) in the central part of FepA primary structure with alanine, individually and in double combinations, and determined the ability of the mutant proteins to interact with ferric enterobactin and the protein toxins colicins B and D. All the constructs showed normal expression and localization. Among single mutants, Y260A and F329A were most detrimental, reducing the affinity between FepA and ferric enterobactin 100- and 10-fold respectively. Double substitutions involving Y260, Y272 and F329 impaired (100- to 2500-fold) adsorption of the iron chelate more strongly. For Y260A and Y272A, the drop in adsorption affinity caused commensurate decreases in transport efficiency, suggesting that the target residues primarily act in ligand binding. F329A, like R316A, showed greater impairment of transport than binding, intimating mechanistic involvement during ligand internalization. Furthermore, immunochimical studies localized F329 in the FepA ligand binding site. The mutagenesis results suggested the existence of dual ligand binding sites in the FepA vestibule, and measurements of the rate of ferric enterobactin adsorption to fluoresceinated FepA mutant proteins confirmed this conclusion. The initial, outermost site contains aromatic residues and probably functions through hydrophobic interactions, whereas the secondary site exists deeper in the vestibule, contains both charged and aromatic residues and probably acts through hydrophobic and electrostatic bonds.

Introduction

Gram-negative bacteria acquire iron with transport systems that usually involve the elaboration of siderophores and the uptake of their ferric complexes. For example, enterobactin, the native siderophore of *Escherichia coli*, forms a hexaco-ordinate complex with Fe$^{3+}$, which enters prokaryotic cells through a multicomponent cell envelope transport system. An outer membrane (OM) ligand-gated porin (LGP), FepA, initiates ferric enterobactin (FeEnt) uptake into the cell; the periplasmic-binding protein (FepB) and an inner membrane complex of proteins (FepCDG) finish the transport process into the cytoplasm.

Multiple functional domains exist within ferric siderophore receptors such as FepA. Besides an external ligand binding site, they contain a transmembrane channel (Rutz *et al*., 1992; Liu *et al*., 1993) and a globular N-terminal domain that lodges within the pore (Buchanan *et al*., 1999). FepA is ligand gated (Rutz *et al*., 1992; Killmann *et al*., 1993), in that the binding of FeEnt on its exterior triggers a series of events that ultimately open the pore, resulting in internalization (Jiang *et al*., 1997). The N-terminal domain was proposed to regulate this transport reaction (Buchanan *et al*., 1999); the fact that it physically blocks the channel requires that it undergoes structural rearrangements during ligand uptake.

The chemistry of FeEnt may explain its interaction with proteins. The catecholate iron complex carries a net charge of $-3$, portending the involvement of aromatic and basic residues in its recognition by FepA. Mutagenesis verified the importance of basic residues (R286, R316) in FeEnt binding (Newton *et al*., 1997) and, in this report, we consider the function of aromatic amino acids in adsorption. FepA and another siderophore receptor, FhuA, contain aromatic residues in their surface loops that encircle the external entrances to their underlying pores (Fig. 1). These amino acids may function in the recognition and discrimination of the many ferric siderophores of the microbial world, or they may create a non-specific, non-polar surface that absorbs these hydrophobic molecules from the environment. We used site-directed mutagenesis and fluorescence spectroscopy to study the participation of aromatic amino acids in the recognition of FeEnt by FepA. Among seven conserved aromatic residues that we substituted with alanine, changes at three, Y260, Y272 and F329, affected the affinity of the FeEnt–FepA interaction. These residues define two sites, one on the
surface of the vestibule and another deep within it. The identification of these separate binding domains, and measurements of the rate of FeEnt binding to FepA mutants with alterations in them, support the two-stage model of siderophore binding to TonB-dependent OM receptor proteins (Payne et al., 1997).

Results

Target residues for mutagenesis

Several previous experiments suggested that the central portion of FepA contains residues that interact with ligands (Murphy et al., 1990; Rutz et al., 1992; Newton et al., 1997). Within this region of the Escherichia coli receptor, we identified numerous residues that we considered potentially important in FeEnt uptake, because they are mostly conserved in the FeEnt transport proteins of Gram-negative bacteria, but not in the other LGP of E. coli. The comparison (Fig. 1) highlighted seven aromatic residues, including five tyrosines (260, 272, 285, 289 and 309), one tryptophan (297) and one phenylalnine (329). Among these, Y260 was distinct, in that it was also conserved in BtuB, FhuA and FhuE.

The multideterminant nature of a ligand–receptor interaction may prevent single substitutions, even for amino acids involved in binding, from producing significant phenotypical effects (Newton et al., 1997). However, double mutagenesis, the pairwise alteration of candidate residues, yields more recognizable defects that may identify mechanistically relevant amino acids. Our initial approach to evaluating aromaticity in FeEnt recognition involved single and double Ala substitutions for the seven aromatic residues of interest, yielding 28 site-directed mutants. The results focused attention on three mutants (Y260A, Y272A and F329A), which we combined with another mutation (R316A) previously found to be critical to the binding reaction (Newton et al., 1997), yielding three more mutants. In addition, we made single and double Ala substitutions for five negatively charged amino acids in the same general region of E. coli FepA, yielding 15 more mutants. Finally, we combined two double mutants with the chemically modifiable mutation E280C to facilitate their biophysical analysis. In total, we engineered and analysed about 50 mutant FepA proteins.

The substitution mutations did not affect the expression of FepA: Western blots showed all the mutant proteins at wild-type levels. The immunoblots also revealed that the chemically modifiable mutation E280C to facilitate their biophysical analysis. In total, we engineered and analysed about 50 mutant FepA proteins.

Affinity screening

The nature of binding and transport assays that involve tripclicate measurements of FeEnt adsorption/uptake at 12 concentrations (Newton et al., 1999) made thorough analysis of so many constructs problematical. Therefore, and because we anticipated only a percentage of the total mutations to affect ligand–receptor interactions (&lt; 20%; Newton et al., 1997, this study), we sought methods of rapidly distinguishing constructs with aberrant phenotypes from those that function at wild-type levels. The use of affinity screening was one step towards this end. The ratio of $^{59}$FeEnt binding or transport at the wild-type $K_d$ (or $K_m$) and at a much higher concentration (e.g. 1000 $K_d$ or 1000 $K_m$) identified impaired FepA proteins (Fig. 3) for subsequent thermodynamic and kinetic analysis. This approach singled out Y260, Y272 and F329. The substitution Y260A strongly decreased affinity for FeEnt, whereas the effect of F329A was most apparent and that of Y272A was only seen in double combinations (Fig. 3). Double substitutions joining Y260A, Y272A or F329A with R316A confirmed the importance of the aromatics in FeEnt binding: each of the double mutants showed much lower affinity than R316A alone. Alanine substitutions for the other residues that we tested had no significant effects on FeEnt adsorption.

Binding equilibria

Recent methodological improvements (Newton et al., 1999) enhanced the sensitivity of $^{59}$FeEnt binding and uptake tests, which enabled us to accurately assess the impact of the site-directed mutations on the receptor protein. Among all the single aromatic mutants observed by affinity screening, and also including R316A, only Y260A showed an obvious (about 100-fold) reduction in adsorption affinity (Fig. 4): the $K_d$ of the FeEnt–FepA binding reaction increased from the wild-type level of 0.1 nM to 10 nM. Y272A did not change affinity at all, whereas F329A and R316A created only nominal alterations in $K_d$, to 0.2 nM and 0.4 nM respectively. Double combinations of these three mutations created predictable
and roughly additive reductions in affinity, from 100- to 2500-fold, which confirmed the relative order of importance seen in the comparisons of single mutants: Y260 > R316 > F329 > Y272 (Table 1). As their affinity decreased, the double mutants manifested a corresponding decrease in FeEnt binding capacity (Table 1, Fig. 4). This reduction probably derived from the inability of the experiment to accurately measure the capacity of low-affinity binding reactions, because of the rapid dissociation of the ferric siderophore from such defective receptors, relative to the time required for the 59FeEnt binding determination. Finally, FeEnt adsorbed more slowly to Y260A and its combinations than to the wild-type receptor protein or other mutants not including Y260A. In vivo, binding to FepA reached equilibrium in less than 5 s at 0°C, whereas for Y260A in the same conditions, even a 1 min incubation was insufficient to reach equilibrium (data not shown).

**Binding kinetics**

The effects of mutagenesis suggested the existence of two distinct stages (Payne et al., 1997), and it was of interest to study the rates of siderophore binding to the mutants with lower affinity. To do so, we combined the covalently modifiable mutation E280C with the double mutants Y272A/F329A and Y260A/R316A. E280 exists in L3 on the external surface of FepA, removed from the mouth of the vestibule (Fig. 1); its substitution with Cys and subsequent reaction with either fluorescent or paramagnetic (Jiang et al., 1997) labels does not affect FeEnt binding and transport. We purified the two triple mutant proteins, modified them with fluorescein and spectrally characterized their FeEnt adsorption rates. The mutants showed distinctly different kinetics from wild-type FepA and from each other (Fig. 5; Table 2). Although both mutant proteins bound the siderophore, neither underwent the complete association reaction of wild-type FepA, as shown by the lower amplitude of fluorescence quenching engendered by FeEnt binding to them. The Y260A/R316A mutation dramatically curtailed the second stage of the adsorption reaction, without impairment of the initial phase. Conversely, Y272A/F329A decreased the amplitude of both binding phases. These results strongly support the hypothesis of an initial binding site affected by the mutation Y272A/F329A and a second, subsequent site influenced by Y260A/R316A (see Discussion). Neither double mutation significantly changed the rate of initial adsorption (k1 in Table 2), as expected for a process that is presumably diffusion limited. However, the high concentration of FeEnt (10 μM) required for saturation of the mutants accelerated the association reaction so much that we could not measure precisely the first binding phase. Both double mutations increased the velocity of the second phase (k3 in Table 2), but the final determination
of these rates awaits more accurate measurements by other methods.

**Transport**

We performed siderophore nutrition tests as a first indicator of transport phenotype. In this assay, FepA mutants may show a variety of effects that differentiate them from wild type (Newton *et al.*, 1999). Constructs involving substitutions for Y260, Y272 or F329 produced three principal results, larger, fainter and fuzzy halos (Fig. 6, Table 1). These mutants, which showed transport deficiencies (see below), grew more slowly on plates than strains expressing wild-type FepA so, at any given time, their siderophore nutrition tests were fainter. Halos derived from wild-type strains developed in 6 h, whereas those from the mutant strains were sparse at that time and required as much as 24 h to reach a similar density. In some cases (Fig. 6), the mutants produced two growth rings: one nearly the same size as the wild type; and a second larger ring.

Uptake screening (Fig. 3) again underscored the inadequacies of Y260A and F329A, but, in this assay, the two mutants were equally, and completely, impaired. None of the other single mutants were deficient in transport screening assays, nor were they defective when combined with each other and tested at a variety of other concentrations. Transport assays over a full range of concentrations clarified the effects of the single Ala substitutions on FeEnt uptake. The maximum velocity of transport was comparatively intransigent. Among the mutations that we generated, only R316A reproducibly decreased the transport rate, but its impact was minor. The $V_{\text{max}}$ of FeEnt uptake by FepAR316A (117 pmol · min$^{-1}$ · 10$^{-9}$ cells) was 65% of the wild-type rate (181 pmol min$^{-1}$ · 10$^{-9}$ cells), and its combinations

![Fig. 3. Affinity screening of FepA mutants.](image)

A. $K_d$ screening of aromatic and basic substitution mutants. $^{59}$FeEnt binding assays were performed at 0.2 nM and 70 nM. The bars depict the ratio of FeEnt bound at these two concentrations. The error bars represent the standard deviations from three experiments.

B. $K_d$ screening of acidic substitution mutants. $^{59}$FeEnt binding assays were performed at 0.2 nM and 70 nM.

C. $K_m$ screening of aromatic substitution mutants. Ferric enterobactin uptake assays were performed at 0.3 nM and 100 nM. The bars depict the ratio of FeEnt transported at these two concentrations. The error bars represent the standard deviations from three experiments.

showed similar decreases. Changes in $K_m$, however, revealed that the alanine substitutions did not always affect transport as they did binding (Table 1, Fig. 4). The relative importance of the four residues was again $Y_260$ and $R_316$. Ala substitutions for $R_316$ and $F_329$ created disproportionate decreases in overall transport affinity, relative to their effects on binding (Table 1). In most cases, binding defects created

| Table 1. Phenotypic properties of FepA aromatic/charge substitution mutants. |
|-------------------------|------------------|------------------|------------------|------------------|------------------|------------------|
| Ferric enterobactin | Binding | | Transport | | Colicin B | | Colicin D |
| FepA mutant | Binding ratio | $K_d$ | Capacity | $K_m$ | $V_{max}$ | Nutrition | Killing | Killing |
| + + | 0.65 | 0.086 | 112 | 0.27 | 181 | 19 | 100 | 100 |
| Y260A | 0 | 9.7 | 88 | 0.33 | 150 | 19 | 100 | 100 |
| Y272A | 0.63 | 0.085 | 103 | 16 | 117 | 22 | 2.5 | 8 |
| R316A | 0.29 | 0.40 | 82 | 16 | 117 | 22 | 2.5 | 8 |
| E319A | 0.27 | 0.59 | 136 | 9.2 | 184 | 22 | 100 | 100 |
| F329A | 0.45 | 0.18 | 100 | 5.5 | 135 | 19 | 50 | 100 |
| Y260A/Y272A | 0 | 33 | 60 | 128 | 126 | 25 | 50 | 100 |
| Y260A/R316A | 0 | 222 | 18 | 793 | 93 | 23 | 2.5 | 8 |
| Y260A/F329A | 0 | 126 | 24 | 367 | 161 | 25 | 50 | 100 |
| Y272A/R316A | 0 | 17 | 80 | 133 | 129 | 24 | 2.5 | 8 |
| Y272A/F329A | 0 | 7.8 | 76 | 23 | 128 | 23 | 2.5 | 8 |
| R316A/F329A | 0 | 83 | 55 | 486 | 105 | 24 | 2.5 | 8 |

a. $K_d$ (nM) and capacity [pmol bound (10^9 cells)^{-1}] were determined from the concentration dependence of Ent binding by analysing the mean values from three independent experiments with GRAFIT 4 (Erithacus), using the ‘bound versus total’ equation. The mean standard errors for $K_d$ and capacity were 14% and 3% respectively.

b. $K_m$ (nM) and $V_{max}$ [pmol min^{-1} (10^9 cells)^{-1}] of uptake were determined from the concentration dependence of Ent transport by analysing the mean values from three independent experiments with GRAFIT 4 (Erithacus), using the ‘enzyme kinetics’ equation. The mean standard errors for $K_m$ and $V_{max}$ were 19% and 5% respectively. Nutrition test results for Ent present the diameters of growth halos in millimetres.

c. Colicin killing was determined by measuring the susceptibility of KDF541 expressing the mutant FepA proteins to limiting dilutions of colicins B and D, and is expressed as a percentage of the killing observed for KDF541/pITS449 (fepA).
congruent loss of transport efficiency, whereas R316A and F329A were more detrimental (about 10-fold) to $K_m$ than to $K_d$. As in the binding determinations, double mutagenesis showed the impact of Y272 on transport: its combinations with Y260A, F329A and R316A further decreased the overall uptake affinity, even though the transport properties of FepA were not impaired by Y272A alone.

Alanine substitutions for acidic residues

The triple-negative charge of FeEnt makes it unlikely that negatively charged residues directly participate in its recognition and binding. Nevertheless, the possibility of a complex network of electrostatic interactions in the binding site led us to evaluate the importance of acidic amino acids in the same region of FepA primary structure, using the same regimen of single and double Ala substitutions. Five acidic residues occur in the region bounded by residues 255 and 355, which localize to either the external vestibule (E280, E310, E319 and E326) or the periplasmic rim of the β-barrel (D298). Analysis of the 15 single and double combinations that we engineered for these acidic amino acids showed that only one created functional defects: E319A was deficient in both FeEnt binding and transport (Figs 3 and 6; Table 1), with $K_d \approx 0.6$ nM and $K_m \approx 9$ nM. Within error, all the other acidic substitution mutants functioned as well as FepA.

Effects of mutagenesis on colicin killing

Colicin B and D killed bacteria expressing the single and double mutants at, or very near, wild-type levels (Table 1), suggesting that unlike the conserved basic residues R286 and R316 the conserved aromatic residues that we studied are not essential to colicin penetration. For the aromatic replacement mutants, the
The amplitudes of the exponential decays are expressed as relative values that were derived from the decrease in F/F₀ previously (Payne et al., 1998; Locher et al., 1999). Those of FhuA: they differ in composition and in length through which ferric siderophores enter. The individual loops of FepA and FhuA, forming a large external vestibule through which ferric siderophores enter. The individual loops of FepA are not homologous or even similar to those of FhuA: they differ in composition and in length (Ferguson et al., 1998; Locher et al., 1998; Buchanan et al., 1999). In both proteins, however, aromatic amino acids that exist in different positions in different loops of the two OM proteins surround the vestibule mouth (Fig. 1). The first conclusion of our experiments is that these exteriormost residues, which are predominantly tyrosines, create an initial binding site (called here B1) for ferric siderophores. The effects of Ala substitution at Y272, one of the residues that form the aromatic ‘lips’ of the vestibule, indicate that changes in B1 impair the binding and, consequently, also the transport of FeEnt. The more dramatic effects of substitution for F329 show that it plays a more important role in the initial recognition site. Furthermore, although the position of this residue was not observed in the crystal structure because of its high flexibility, the discovery of F329 in the epitope of an antibody that competitively inhibits FeEnt binding localizes the residue in a ligand recognition site, exactly as suggested by the mutagenesis results. Thus, the data support both the existence of the outer B1 site, which primarily functions by hydrophobic bonds, and the importance of aromatic residues F329 and Y272 within it. Although the catecholate nature of FeEnt led us to postulate the involvement of aromatic amino acids in its recognition by FepA, in fact, other OM proteins (FhuA, LamB) that interact with non-aromatic ligands also contain aromatic residues in their binding pockets, illustrating the generality of this adsorption mechanism.

Deeper within the vestibule mouth, FepA and FhuA diverge. The former protein, which transports a trianionic catecholate siderophore, contains an abundance of basic residues grouped in a cluster at the top of its N-terminal domain. The latter protein, which transports an uncharged, hydrophobic, hydroxamate peptide siderophore, contains a concentration of aromatic residues in the same relative position. The properties of these potential secondary sites (B2) match the chemical attributes of the compounds that they recognize, perhaps conferring specificity to the binding reaction. The FhuA crystal structures contained ferrichrome bound in the B2 region. Although the FepA crystal structure did not resolve the position of FeEnt in the loops (it showed partial occupancy in their outer extremities; Buchanan et al., 1999), the co-ordinates depict R316 and Y260 within B2, appropriately located to interact with the ferric siderophore as it approaches and transits the channel (Fig. 1). The functional importance of Y260 constitutes a second conclusion of

### Table 2. Kinetic constants of FeEnt association with mutant FepA proteins.

<table>
<thead>
<tr>
<th>Protein</th>
<th>kₐ</th>
<th>A₁</th>
<th>k₃</th>
<th>A₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>E280C</td>
<td>0.68 ± 0.03</td>
<td>0.30 ± 0.005</td>
<td>0.30 ± 0.00001</td>
<td>0.24 ± 0.0006</td>
</tr>
<tr>
<td>E280C–Y272A–F329A</td>
<td>0.87 ± 0.06</td>
<td>0.26 ± 0.004</td>
<td>0.008 ± 0.0001</td>
<td>0.12 ± 0.001</td>
</tr>
<tr>
<td>E280C–Y260A–R316A</td>
<td>0.97 ± 0.07</td>
<td>0.33 ± 0.008</td>
<td>0.04 ± 0.002</td>
<td>0.09 ± 0.004</td>
</tr>
</tbody>
</table>

a. The proteins were purified, labelled with IAF, suspended in buffer at 25 nM, FeEnt was added to 10 μM, and the rates of association were measured by fluorescence spectroscopy.

b. Association rate constants, which were calculated based on a double exponential decay process, using GRAFIT 4, are denoted as described previously (Payne et al., 1997).

c. The amplitudes of the exponential decays are expressed as relative values that were derived from the decrease in F/F₀.

**Discussion**

Any experimental study involving mutagenesis as a diagnostic tool of protein function must consider potential changes in global conformation as a possible artifact. Three independent lines of evidence argue against the idea that the reduced siderophore binding and transport abilities of the mutant proteins arose from unanticipated but debilitating conformational changes as a result of mutagenesis. All the single mutants produced relatively small decreases in affinity, which, when combined, were additive. All the mutants, including double mutants that were severely impaired in binding and transport of FeEnt, showed virtually wild-type levels of colicin susceptibility, indicating that both the surface structures and the general functionality of the mutant proteins remained intact. Finally, the overall results from the series of residues that we considered in the experiments, 13 aromatic, acidic and basic amino acids in surface loops, β-strands and on the rim of the β-barrel, in the same general region of FepA, pertain to the question of structural distortions from mutagenesis. With the exception of Y260, Y272, F329 and E319, which all reside in appropriate locations for contact with the siderophore, no deficiencies were found in any of the other mutant proteins. These data rebut the idea that the site-directed mutations caused significant conformational changes in the region of interest.

Eleven surface loops encircle the membrane channels of FepA and FhuA, forming a large external vestibule through which ferric siderophores enter. The individual loops of FepA are not homologous or even similar to those of FhuA: they differ in composition and in length (Ferguson et al., 1998; Locher et al., 1998; Buchanan et al., 1999). In both proteins, however, aromatic amino acids that exist in different positions in different loops of the two OM proteins surround the vestibule mouth

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our experiments: the 100-fold reduction in affinity from Y260A is the largest effect of a single mutation seen in FepA so far, suggesting that the high affinity for FeEnt derives from interactions within the B2 site. The 2500-fold reduction in binding and transport seen in the Y260A/R316A double mutant leaves no doubt about the importance of B2 in siderophore acquisition. This region contains charged and aromatic amino acids, suggesting that both hydrophobic and electrostatic bonds contribute to its functionality. The proximity of the E319 carboxylate moiety (7 Å from the guanidino group of R316) underscores the inference of ionic interactions within the B2 site.

The hypothesis of two FeEnt binding sites in FepA originated from in vivo fluorescence measurements that showed kinetically discernible stages of FeEnt adsorption (Payne et al., 1997), and our rate measurements of binding by site-directed mutants corroborate and extend this theory. The adsorption of FeEnt quenches the emission spectrum of fluoresceinated FepA by about 60%, and the drop in fluorescence is consistent with the sum of two sequential exponential decay processes. The high-affinity, saturation effects of FeEnt on the fluorescent reporter group (Payne et al., 1997) imply that the quenching does not originate from direct, non-specific collisions between the siderophore and fluorescein; the distance of E280C from the mouth of the FepA vestibule (Fig. 1; Buchanan et al., 1999) strengthens this conclusion. Thus, the time-dependent decrease in fluorescence intensity derives from a permutation of FepA structure that changes the environment of the fluorescent label. In the mutants, the magnitude of quenching by FeEnt was less than in wild-type FepA, indicating that they underwent different conformational changes. Double mutagenesis of Y260 and R316 increased the amplitude of the first stage, and virtually eliminated the second stage. A straightforward interpretation of these results is that FeEnt binding causes one conformational change when it associates with the outer site, and another change when it moves to the inner site. In view of this, the Y260A/R316A mutation seriously compromised the second site, without measurable effects on the first and, as a result, the first binding phase predominated. This explanation also rationalizes the tendency of Y260A to retard progress towards binding equilibrium. Ala substitution for Y272 and F329, on the other hand, affected both stages of the association reaction. This result is consistent with the idea that decreased efficiency of the initial binding will also reduce the extent of ligand movement to the second site. Consequently, the amplitudes of both stages will lessen. It is noteworthy that more complex, alternative explanations are also plausible. Presteady-state experiments are needed to clarify the respective roles of B1 and B2 in the overall uptake process. In spite of this caveat, the spectroscopic results verify the existence of the B1 and B2 sites, which are kinetically, conformationally and functionally linked in the FeEnt association reaction. Thus, another conclusion of the experiments is that the first binding stage abstracts the ligand from solution into B1, whereas the second step involves inward movement of FeEnt to B2, where it sits poised above the transmembrane channel before transport.

The characteristic catechol groups of ferric enterobactin provide several potential modes of interaction between FeEnt and FepA. Aromatic rings may act as hydrogen bond acceptors (Levitt and Perutz, 1988; Mitchell et al., 1994), and they may form planar stacking interactions with other ring systems and with arginines (Flocco and Mowbray, 1994). Van der Waals’ forces and hydrophobic effects may also determine ligand interaction behaviour with proteins (Kuntz et al., 1999). In low-affinity protein recognition systems, hydrophobic interactions often provide affinity for ligand adsorption, whereas charge–charge interactions are responsible for recognition specificity (Davis et al., 1998). In this case, electrostatic forces usually contribute by creating a correct conformation or alignment that leads to complementarity between the ligand and its receptor (Jones and Thornton, 1996). On the other hand, in high-affinity recognition systems, charge–charge interactions may provide not only elements of specificity, but also enhanced ligand affinity (Chong et al., 1999). Our experiments suggest that the FeEnt–FepA binding equilibrium exhibits both kinds of interactions: the B1 site functions by hydrophobic bonds, whereas in B2, hydrophobic and electrostatic forces combine to create an extremely precise (Thulasiraman et al., 1998), avid (Newton et al., 1999) recognition phenomenon. The adsorption reaction is sensitive to alterations in these chemical characteristics of FepA, and double mutagenesis confirmed the multideterminant nature of the interaction: when two apparent contact residues were removed from FepA, whether two aromatic, two basic, or one of each, a substantial decrease in the receptor function (>1000-fold) occurred.

The independence of colicin killing from the aromatic residues that are important in FeEnt utilization was unexpected. F329 and Y272, which exist in the peripheral interface between the protein and the aqueous environment, were dispensable to colicin penetration. Furthermore, two positively charged amino acids in the interior of FepA, R286 and R316, were essential to colicin B and D susceptibility, but Y260, which was required for optimal FeEnt binding and transport, was also dispensable to colicin penetration. The close proximity of Y260 to R316 suggests that the positive charge of the latter amino acid, rather than its location, is most relevant to colicin reception.

The experiments described here and other studies
(Rutz et al., 1992; Newton et al., 1997) used mutagenesis to investigate the relationship between structure and function in FepA. An initial perception of the research, upon which some of our mutagenesis targets were based, was that one or a few surface loops in the central part of FepA sequence (L3, L4, L5) are pre-eminent in the recognition and binding of FeEnt. The primary argument against this conclusion is that FepA functions, albeit at much reduced efficiency, without these single loops (Newton et al., 1999). The current experiments provide insight into this result. The OM protein contains two binding sites for the ferric siderophore, one formed by residues in the loop extremities, and the other deep in the vestibule, primarily composed of side-chains from the interior barrel wall, and perhaps also residues in the N-terminal domain (although our data do not address this region). Single loop deletions did not abrogate FeEnt recognition and binding because (i) multiple loops participate in the initial adsorption reaction (evidenced by the identification of Y272 in L3 and F329 in L4); and (ii) they did not eliminate the secondary, underlying binding site (involving at least Y260 and R316). It is likely that other loops contribute contact residues to the binding reaction (Fig. 1), so we are left with the perception of two multideterminant binding domains formed by the coalescence of residues distributed throughout FepA primary structure. As the precision of the mutagenic approach escalated, it revealed this final conclusion with increasing certainty.

Experimental procedures

Bacterial strains and media

Escherichia coli K-12 strains were grown in Luria–Bertani broth (Miller, 1972), Tris media (Klebba et al., 1982) or MOPS media (Neidhardt et al., 1974).

Sequence comparisons

The sequences of six FeEnt transporters from E. coli (Lundrigan and Kadner, 1986), Salmonella typhimurium (Tumuru et al., 1990), Pseudomonas aeruginosa (Dean and Poole, 1993), Salmonella enterica (Baumler et al., 1998), Bordetella pertussis (Beall and Sanden, 1995) and five other E. coli LGPs, FecA, BtuB, IutA, FhuA and FhuE were compared using the PILEUP program (Genetics Computer Group, Madison, WI, USA). The alignment was manually adjusted on the basis of the E. coli FepA crystal structure (Buchanan et al., 1999).

Site-directed mutagenesis

We previously used pITS449 (Armstrong et al., 1990) as the starting point for site-directed mutagenesis. In the present work, we created vector pT944, a pITS449 derivative that incorporates restriction sites without changing the fepA coding sequence. PT944 introduced a Smal site at position 126 (GCCCGCGAT to GCCCGGGAT), removed the methylation site for Clal at position 1023 (ATCGATCTT to ATCGATTCT) and introduced a BamHI site at position 1701 (GGATCG to GGATCC). These modifications facilitate manipulation of the vector when combining independent mutations. Using PstI and SacI sites, the fepA structural gene from pT944 was inserted into the multiple cloning site of M13 mp19. To create individual fepA mutants, we performed site-directed mutagenesis with the structural gene in M13 (Kunkel, 1985). The mutation in fepA was then cloned back into the pT944 fepA structural gene using appropriate restriction sites and enzymes. The entire mutant gene was sequenced to rule out any random mutagenesis. The resultant plasmids were finally transformed into KDF541 (fepA; Rutz et al., 1992).

Expression

KDF541 strains harbouring pITS449, pT944 or their mutant derivatives were grown in LB broth with ampicillin and streptomycin (both at 100 μg ml⁻¹) for 16 h at 37°C before subculture at 1% into MOPS media with ampicillin (10 μg ml⁻¹). After growth for 5.5 h at 37°C with vigorous aeration, 5 × 10⁷ bacteria cells were collected by centrifugation and lysed by boiling with SDS–PAGE sample buffer. After electrophoresis on 10% slabs, the separated proteins were transferred to nitrocellulose paper, which was analysed by immunoblotting with anti-FepA mAb 41 or mAb 45 and [¹²⁵I]-protein A (Newton et al., 1997).

FeEnt binding

We intended to use alanine scanning mutagenesis, in conjunction with thermodynamic and kinetic analyses, to elucidate the FeEnt–FepA interaction, but the necessity of double substitutions expanded the scope of the experiments so much that screening methods were required to find constructs of interest for further study. Affinity screening was the improvement that expedited the experiments. The finding that prior binding and transport methodologies underestimated the affinities of ferric siderophore–receptor systems as much as 50 000-fold (Newton et al., 1999) focused our attention in the subnanomolar range, and the ratio of wild-type Kd or Km (≈ 0.1 nM) to a second, much higher concentration readily identified mutants with defects. The methodological corrections also facilitated accurate characterization of both single and double mutants. It did not, however, alleviate the need for double mutagenesis; Y272A again illustrated the concept that substitution for a functionally relevant residue may not compromise a multivalent ligand–receptor interaction (Newton et al., 1997).

To quickly assess siderophore binding, we compared ⁵⁹FeEnt adsorption (Newton et al., 1999) at the wild-type Kd and a much higher concentration. Mutants that were impaired in the binding screen were further analysed by determination of their Kd and capacity for ⁵⁹FeEnt (Newton et al., 1999). For FepA and mildly defective mutants, 2 × 10⁷ cells were tested with ⁵⁹FeEnt at approximately 200 c.p.m. pmol⁻¹⁻¹; for more impaired mutants, 10 times more cells were assayed, and the ⁵⁹FeEnt was diluted with non-radioactive FeEnt to reduce the
specific activity to about 20 c.p.m. pmol\(^{-1}\). For even more severely defective constructs, \(5 \times 10^8\) cells were used, and \(^{59}\)FeEnt was prepared at about 10 c.p.m. pmol\(^{-1}\). Both screening assays and \(K_d\) determinations were corrected with measurement of \(^{59}\)FeEnt binding to the negative control KDF541; the data were analysed by GRAFIT 4.

**FeEnt uptake**

To quickly assess siderophore transport, we performed siderophore nutrition tests (Newton et al., 1999) and compared \(^{59}\)FeEnt uptake (Newton et al., 1999) by screening at the wild-type \(K_m\), and a much higher concentration. Mutants of interest were further characterized by determination of transport \(K_m\), and \(V_{\text{max}}\). As with the binding experiments, uptake assays on mutants with decreased capabilities required an increase in the number of cells and a decrease in the specific activity of \(^{59}\)FeEnt. In addition, the incubation times of the transport experiments were increased (from 15 s for wild-type FepA to up to 2 min) to accommodate the lower uptake rates of the mutants. GRAFIT 4 was used to determine the thermodynamic and kinetic parameters.

**Fluorescence measurements**

Purified FepA proteins containing the E280C mutation (final concentration 0.15 mg ml\(^{-1}\)) in TTE buffer (50 mM Tris, pH 7.2, 2% Triton X-100, 5 mM EDTA) were incubated with 10 \(\mu\)M 5-iodoacetamidofluorescein (IAF; Molecular Probes) at room temperature in the dark with shaking for 45 min, ethanol precipitated three times and resuspended in 1 mM n-dodecyl \(\beta\)-d-maltoside (DM; Sigma), 50 mM MOPS, pH 7.0, 60 mM NaCl buffer. The reaction was then transferred to dialysis tubing (MWCO 12 000–14 000) and dialysed against the above buffer at 4 \(^\circ\)C with three changes of buffer for 48 h to remove unreacted probes. The concentration of the labelled proteins was determined before storage at –80 \(^\circ\)C.

In the FeEnt binding time course measurement, E280C labelled with IAF was diluted into 1 mM DM, 50 mM MOPS, pH 7.0, 60 mM NaCl buffer to a final concentration of 25 \(\mu\)M and incubated at 25 \(^\circ\)C for 1 h to ensure the stabilization of E280C–IAF. The time course was then started on the SLM8100 fluorimeter at an excitation and an emission wavelength of 490 nm and 520 nm, respectively, with the slits all set at 4 nm and an integration time of 1 s. After about 5 min of prerun, ferric enterobactin in 50 mM MOPS, pH 7.0, 60 mM NaCl was added to a final concentration of 10 \(\mu\)M. The same fluorescent labelling and time course measurement procedures were performed on Y272A–F329A–E280C and Y260A–R316A–E280C mutants. Wild-type protein FepA was used as a control.

**Colicin killing**

To examine the colicin sensitivity of each mutant FepA, a series of twofold dilution of colicins B and D were made in LB broth in a 96-well microplate. Using a Clonemaster (Immusine), 5 \(\mu\)l of diluted ColB or ColD was then transferred from the master plate to LB agar plated with KDF541 harbouring pTS449 or its mutant derivatives. After incubation at 37 \(^\circ\)C overnight, sensitivity was expressed as the maximum colicin dilution that resulted in clearing of the bacterial lawn to the agar.

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**References**


