The Ferric Enterobactin Transporter Fep Is Required for Persistent 
Salmonella enterica Serovar Typhimurium Infection

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Most bacterial pathogens require iron to grow and colonize host tissues. The Gram-negative bacterium Salmonella enterica serovar Typhimurium causes a natural systemic infection of mice that models acute and chronic human typhoid fever. S. Typhimurium resides in tissues within cells of the monocyte lineage, which limit pathogen access to iron, a mechanism of nutritional immunity. The primary ferric iron import system encoded by Salmonella is the siderophore ABC transporter FepBDGC. The Fep system has a known role in acute infection, but it is unclear whether ferric iron uptake or the ferric iron binding siderophores enterobactin and salmochelin are required for persistent infection. We defined the role of the Fep iron transporter and siderophores in the replication of Salmonella in macrophages and in mice that develop acute followed by persistent infections. Replication of wild-type and iron transporter mutant Salmonella strains was quantified in cultured macrophages, fecal pellets, and host tissues in mixed- and single-infection experiments. We show that deletion of fepB attenuated Salmonella replication and colonization within macrophages and mice. Additionally, the genes required to produce and transport enterobactin and salmochelin across the outer membrane receptors, fepA and iroN, are needed for colonization of all tissues examined. However, salmochelin appears to be more important than enterobactin in the colonization of the spleen and liver, both sites of dissemination. Thus, the FepBDGC ferric iron transporter and the siderophores enterobactin and salmochelin are required by Salmonella to evade nutritional immunity in macrophages and cause persistent infection in mice.

In humans, Salmonella enterica serovars Typhi and Paratyphi A, B, and C cause typhoid fever, a systemic infection with mortality rates exceeding 15% if left untreated (1, 2). S. enterica serovar Typhimurium causes murine typhoid in mice, a disease that models both the acute and chronic stages of human typhoidal infection. Upon ingestion, Salmonella bacteria pass through the stomach and traverse the intestinal epithelial barrier to colonize Peyer’s patches, mesenteric lymph nodes, spleen, and liver (3, 4). Salmonella bacteria reside within cells of the monocyte lineage, typically macrophages that contain the bacteria within a specialized membrane-bound compartment called a Salmonella-containing vesicle (5, 6).

Iron is an essential nutrient for animals. However, iron must be tightly regulated, as it can both accept and donate electrons, leading to the generation of damaging free radicals (7). Dietary ferric iron (Fe³⁺) is reduced to ferrous iron (Fe²⁺) by a ferrireductase (DcytB) on the luminal side of intestinal epithelial cells (8, 9). Ferrous iron is then taken up by enterocytes via DMT1/Nramp2/Slc11A2, exported into the blood via ferroportin 1, and oxidized to ferric iron by hematoxin on the basolateral membrane (10). Blood ferric iron is captured by transferrin, a high-affinity iron binding protein ($K_d$ [dissociation constant] = $10^{22}$ M⁻¹) (11). In tissues, cells endocytose transferrin, and ferric iron is released upon endosomol acidification. The released ferric iron is then reduced (12) and transported to the cytosol (13). Cytosolic iron is sequestered by ferritin, which accounts for about 16% of iron stores in the body (14).

A major host defense against infection is nutritional immunity, the sequestration of essential molecules, including metals, to prevent pathogen outgrowth (15). Sequestration of iron is an effective antimicrobial defense because iron is a cofactor required for crucial processes, including energy production and DNA replication. For instance, during acute or chronic immune activation, ferritin-bound iron accumulates in macrophages, a process that withholds iron from extracellular pathogens and is characteristic of the anemia of chronic disease (14, 16). In contrast, macrophages infected with Salmonella increase the export of iron, and the spleen and liver of Salmonella-infected mice do not accumulate iron (17–19). Decreased iron concentrations in macrophages limit Salmonella replication (20, 21), highlighting the importance of delineating how Salmonella acquires iron under such starved conditions.

Salmonella acquires ferric iron by secreting the catecholate siderophores enterobactin and salmochelin. Once bound to iron, enterobactin and salmochelin transit the bacterial outer membrane receptors FepA and IroN, respectively, in a TonB-dependent manner (22). Enterobactin and salmochelin then bind to FepB in the periplasm for import through FepDGC (23). During infection, macrophages respond to gamma interferon (IFN-γ) by increasing the secretion of lipocalin-2, a siderophore-capturing protein (20, 24–27). Lipocalin-2 binds enterobactin but not salmochelin, a glycosylated derivative of enterobactin that gives Salmonella a selective growth advantage in the intestine (26).

Whether Salmonella requires access to ferric iron has been examined in mice that lack functional Nramp1/Slc11a1, a ferrous
iron transporter in the membrane of the phagosome. Nramp1-deficient mice have dysregulated iron metabolism and are exquisitely sensitive to microbial pathogens that reside within vesicles, including Salmonella, Mycobacterium tuberculosis, and Leishmania species. Salmonella strains lacking fepA and iroN have no apparent phenotype in Nramp1−/− mice in single and mixed infections (23, 28) or in survival assays upon intragastric or intraperitoneal inoculation compared with the wild type (29). Together, these studies indicate that enterobactin and salmochelin uptake via FepA and/or IroN is not essential for Salmonella infection of Nramp1−/− mice.

Nramp1−/− mice, such as the Sv129S6 strain, survive acute infection and develop chronic infection. In these animals, the synthesis and secretion of salmochelin are required for Salmonella virulence upon intraperitoneal inoculation, as measured by survival assays (26, 30). Intraperitoneal inoculation allows bacteria to bypass the gastrointestinal tract and directly access the spleen (31). However, Pearson et al. demonstrated that upon intranasal inoculation of piglets, there was no colonization defect of a triple mutant lacking fepA, iroN, and cirA compared to the wild-type strain (32). CirA is a third outer membrane receptor for siderophore-mediated iron uptake via FepBDGC (27, 28, 33). However, CirA does not directly bind enterobactin or salmochelin but rather captures catecholate breakdown products containing ferric iron. In the work presented here, we establish the requirement of Salmonella for FepB and the enterobactin and salmochelin binding proteins FepA and IroN for gastric and deep-tissue colonization of Nramp1−/− mice upon orogastric inoculation.

Materials and Methods

Bacterial strains and growth conditions. Salmonella enterica serovar Typhimurium wild-type strain SL1344 (34) and mutant derivatives were grown overnight at 37°C with aeration in Luria-Bertani (LB) medium prior to infections. Antibiotics were used at the following concentrations: streptomycin at 30 μg/ml and kanamycin at 30 μg/ml.

Deletion mutants in genes involved in iron uptake and metabolism were constructed by using lambda Red methodology (35, 36), ΔiroN, ΔfepA, and ΔentC mutants of background strain 14028 were generated previously by using a similar methodology (36), and the ΔfepB mutant was generated in this study by using primers fepB fwd (5′-GCGCTAACC TAAGAGTAAAACGTCGCTCTGTCAACTGTGTAGGCTGGAGCTGC-3′) and fepB rev (5′-AATCGGTCTGGTCAGTCGGATAAGACTCG-3′) and P22 phage lysates were prepared from each mutant strain and insertion of the cassette by using site-specific and internal kanamycin primers. P22 phage lysates were prepared from each mutant strain and used to transduce each mutation into wild-type strain SL1344.

Restoration of fepB was achieved by P22 phage transduction of a wild-type copy into the fepB-deleted strain. To enrich for colonies in which the ΔfepB mutant had 100- to 1000-fold lower CFUs, the transduced strain was plated on LB agar and then serially diluted for plating to enumerate CFUs. The infectious dose was verified by plating for CFUs on selective LBS agar. The period in which poor growing may be observed, mice were monitored twice daily. Cages containing “scruffy” mice were supplied with food and water in small dishes on the floor of the cage to alleviate suffering from dehydration and malnourishment. Two weeks after inoculation, infected animals were euthanized by CO2 asphyxiation, followed by cervical dislocation. Spleen, liver, mesenteric lymph nodes, Peyer’s patches, and cecum were collected; homogenized in 1 ml PBS; and then serially diluted for plating to enumerate CFU. Competitive indexes (CIs) were calculated as follows: (CFUwild type/CFUmutant) input/output (CFUwild type/CFUmutant) output. Gentamicin protection assays. RAW264.7 cells stably transfected with the pHBA-1-neo expression plasmid containing the full-length Slc11a1 cDNA (25) were seeded at 1.5 × 104 cells per well in poly-L-lysine-coated 24-well tissue culture plates. Cells were activated with 20 ng/ml lipopolysaccharide (LPS) (S. enterica serovar Typhimurium LPS; Sigma-Aldrich) and 20 U/ml IFN-γ (Peprotech) for 18 h. Bacteria opsonized with normal mouse serum (sigma) were added to macrophages at a multiplicity of infection of 10:1 (bacteria to macrophage). After 30 min, free and loosely adherent bacteria were removed by washing with PBS twice, and cells were incubated for a further 1.5 h at 37°C in fresh medium supplemented with gentamicin (100 μg/ml) to kill extracellular bacteria. Medium was then exchanged for medium supplemented with 10 μg/ml gentamicin to inhibit extracellular bacterial growth. At 2, 18, and 24 h, wells were washed twice with prewarmed PBS, incubated with 1% Triton X-100 for 5 min, and lysed, and serial dilutions were plated to enumerate CFU.

Statistics. P values were calculated with GraphPad Prism 5 (GraphPad Software Inc.) and considered significant if the P value was <0.05. For nonparametric data, Wilcoxon signed-rank or Mann-Whitney tests were used. Otherwise, one-sample t tests or Student’s t tests were used.

Results

Salmonella fepB ferric iron transporter mutants colonize mice poorly in competitive infections. We constructed a mutant Salmonella Typhimurium SL1344 strain with a deletion of fepB, encoding a protein crucial for ferric iron uptake. To establish whether FepB is needed for Salmonella survival and growth in tissue, mixed-infection experiments were performed in Sv129S6 (Nramp1−/−) mice. These mice become chronically infected with Salmonella and can be used to monitor infection long term (38). We evaluated the number of wild-type organisms and the number of ΔfepB mutant bacteria were recovered from fecal pellets, indicating that the ΔfepB mutant colonizes the intestine well initially (Fig. 1A). However, by 3 days postinfection, the ΔfepB mutant had 100- to 10,000-fold-decreased colonization relative to the wild-type organism (Fig. 1A).

Nagy et al.
At 2 weeks postinfection, mice were euthanized, and tissues were collected to enumerate CFU. In gut-associated tissues, including the cecum, Peyer’s patches, and mesenteric lymph nodes, the numbers of \( \Delta fepB \) strain bacteria were 100-fold reduced relative to the numbers of wild-type organisms (Fig. 1B). In the spleen and liver, the phenotype was more severe, as indicated by the competitive index, which was 1,000 times lower for \( \Delta fepB \) than for wild-type strains (Fig. 1B). These data suggest that ferric iron transport by the Fep system is required for tissue colonization in the context of coinfection with wild-type bacteria.

Salmonella requires the FepB ferric iron transporter for persistent infection of mice. We used single-infection experiments in Sv129S6 (Nramp1\(^{+/+}\)) mice to establish whether the ferric iron transporter mutant colonizes tissues normally in the absence of the wild-type organism. At 1 day postinfection, the \( \Delta fepB \) mutant was recovered at slightly but significantly lower numbers from fecal pellets of mice infected with the \( \Delta fepB \) mutant than from mice infected with the wild-type organism (Fig. 1C). Within 3 days postinfection, the difference in recovery from fecal pellets was >10,000-fold, with mice infected with the \( \Delta fepB \) mutant exhibiting a strong colonization defect. The large difference in relative strain recovery at 1 and 3 days suggests that the \( \Delta fepB \) mutant initially colonizes the intestine well. At 2 weeks postinfection, there were \( 10^3 \) to \( 10^6 \) CFU per gram of wild-type Salmonella bacteria in each tissue examined, while tissues of mice infected with the \( \Delta fepB \) mutant lacked detectable colonization (Fig. 1D). These results, in combination with the mixed-infection experiments, clearly demonstrate that Salmonella infection of Nramp1\(^{+/+}\) mice requires the FepB ferric iron transporter in all tissues examined.

Salmonella requires the FepB ferric iron transporter for replication in macrophages. During persistent infection, S. Typhimurium resides within macrophages (5, 38). To establish whether FepB is required for Salmonella replication in macrophages, macrophage-like Nramp1\(^{+/+}\) tissue culture cells were individually infected with equivalent numbers of wild-type or \( \Delta fepB \) mutant Salmonella bacteria. Infected cells were lysed, and CFU were enumerated over a time course (Fig. 2A). The wild-type strain replicated during the first 18 to 24 h, while CFU recovered for the \( \Delta fepB \) mutant strain remained similar throughout the experiment. Expression of fepB on a plasmid is toxic to Escherichia coli (40) and Salmonella (data not shown). Therefore, to confirm that fepB is needed for replication in macrophages, wild-type fepB was restored in the fepB-deleted strain by phage transduction. Replication of the restored strain in macrophages was comparable to that of the wild-type strain. Macrophages were also treated with 50 \( \mu \)M deferasirox, a ferric iron-specific chelator, over a time course of infection. At each time point examined, replication of the wild type in the presence of deferasirox was significantly reduced compared to the replication of bacteria in the presence of the vehicle alone (Fig. 2B), similar to results obtained by Nairz et al. (41). These results demonstrate that ferric iron transport via FepB is required for replication of Salmonella in macrophages.

Salmonella can utilize ferric iron via FepB as its sole iron source. Next, we determined whether ferric iron is sufficient to
FepA and IroN in mice. 

Plated to determine numbers of CFU/ml. Error bars indicate standard errors of the means (n = 3 experiments). *P < 0.05 versus the wild type.

Supporting Information. 

FIG 2 Salmonella requires ferric iron (fepB) for growth in macrophages. (A) Cell culture macrophages (Nramp1+/+) were infected with the wild-type, ΔfepB, or fepB-restored (fepB+) strain at a multiplicity of infection of 10. At 2, 18, and 24 h postinfection, macrophages were lysed, serially diluted, and plated to determine numbers of CFU/ml. Error bars indicate standard errors of the means (n = 3 experiments). *P < 0.05 versus the wild type. (B) Cell culture macrophages (Nramp1−) were infected with the wild type in the absence or presence of 50 μM deferasirox at a multiplicity of infection of 10. At 2, 18, and 24 h postinfection, macrophages were lysed, serially diluted, and plated to determine numbers of CFU/ml. Error bars indicate standard errors of the means (n = 3 experiments). *P < 0.05 versus the wild type.

FIG 3 Salmonella can grow with ferric iron as the sole iron source. (A) The indicated strains were grown in LB medium supplemented with 200 μM iron chelator 2,2'-dipyridyl (dipi). The optical density at 600 nm was monitored for 16 h. Error bars indicate standard deviations (n = 3 experiments). **P < 0.05 versus the wild type; *P < 0.05 versus the ΔfepB mutant. (B) The indicated strains were grown in M9 minimal medium supplemented with ferric chloride (Fe3+) and/or 200 μM dipiridyl. The optical density at 600 nm was monitored for 16 h. Error bars indicate standard deviations (n = 3 experiments). *P < 0.05 versus the wild type; **P < 0.05 versus the ΔfepB mutant.

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poor tissue colonization in vivo. To determine whether the in vivo growth defects observed for the ΔiroN, ΔfepA, or ΔiroN ΔfepA strains reflected enhanced siderophore production, a strain unable to import or produce enterobactin and salmochelin (ΔiroN ΔfepA ΔentC) was generated. Mice orogastrically inoculated with individual wild-type or ΔiroN ΔfepA ΔentC bacteria were sacrificed at 2 weeks postinfection, and tissue colonization was determined (Fig. 6). Tissue colonization by the triple mutant strain was significantly reduced compared to that of the wild-type strain in the cecum, Peyer’s patches, mesenteric lymph nodes, and spleen, although the difference in liver colonization was not significant. These observations indicate that the reduced tissue colonization observed for ΔfepA and ΔiroN mutants is not dependent upon siderophore overproduction.

FIG 4 Salmonella requires enterobactin and salmochelin during persistent infection of mice. In separate experiments (left and right), mice were orogastrically inoculated with the wild-type, ΔiroN, ΔfepA, or ΔiroN ΔfepA strain. After 2 weeks, mice were sacrificed, and the cecum (A), Peyer’s patches (B), mesenteric lymph nodes (C), spleen (D), and liver (E) were immediately homogenized and plated to determine bacterial loads. Each symbol represents one mouse (n = 5 to 10). *, P < 0.05 versus the wild type; **, P < 0.005 versus the wild type.

FIG 5 Salmonella requires either fepA or iroN for growth in macrophages. Cell culture macrophages (Nramp1−/−) were infected with the wild-type, ΔiroN, ΔfepA, or ΔiroN ΔfepA strain at a multiplicity of infection of 10. At 2, 18, and 24 h postinfection, macrophages were lysed, serially diluted, and plated to determine numbers of CFU/ml. In panel B, the CFU index is defined as the number of CFU at 18 h relative to the number of CFU at 2 h (18 h) and the number of CFU at 24 h relative to the number of CFU at 18 h (24 h). Error bars indicate standard deviations (n = 3 experiments). *, P < 0.05 versus the wild type.
DISCUSSION

Salmonella encodes transporters for both ferric and ferrous iron. Ferrous iron is taken up by FeoB (43) and by divalent cation transporters (MntH and SitA) (44, 45). FeoB and SitA have known roles in BALB/c Nramp1−/− mice during acute infection (43, 44). Ferric iron is acquired upon the import of iron-bound siderophores through FepA and IroN and siderophore capture in the periplasm by FepB (23). Previous work established that fepA and iroN are dispensable for infection of BALB/c Nramp1−/− mice (23, 28, 29). We demonstrate that siderophore capture and import via fepB are required for Salmonella survival and replication within macrophages as well as for colonization of mice upon orogastric inoculation. In addition, the ability to produce and/or take up either the enterobactin or salmochelin siderophores is needed for Salmonella colonization of both gastric and deep tissues in Sv129S6 Nramp1−/− mice. Our observations are consistent with those of Gorbacheva et al., who found that an S. Typhi mutant strain lacking the ability to synthesize enterobactin (and salmochelin) failed to replicate in human macrophages (46). The growth defect observed in strains lacking fepB was more severe than that in strains unable to produce or import either siderophore. This may reflect that FepBDGC also takes up derivatives of enterobactin or salmochelin (27, 28, 33). Overall, the data suggest that S. Typhimurium requires ferric iron during persistent infection of mice. Moreover, the results from infection of mice with single mutant strains indicate that enterobactin acquisition through FepA is important for colonization of the cecum and Peyers patches and that enterobactin or salmochelin import via IroN is important for colonization of all tissues examined. Our studies thus provide an experimental foundation to make hypotheses about the temporal-spatial requirements of siderophores during different stages of infection.

Mice that lack hemochromatosis gene alleles (Hfe) have increased resistance to Salmonella infection. This is due to decreased levels of macrophage iron via an enhanced production of ferroportin (47) as well as increased iron sequestration via augmented lipocollin-2 production, which scavenges enterobactin (25). These results are in concordance with our current studies indicating a preferential uptake of ferric iron via IroN and FepA during infection. Because hemochromatosis studies have been performed in Nramp1−/− mice, it will be of interest to determine if similar mechanisms occur in Nramp1−/+ mice, which tightly regulate iron stores and mimic the acute-to-chronic progression of human typhoid fever.

Patients with β-thalassemia and thus ferric iron overload, including within macrophages, have increased susceptibility to bacterial infection. Mice with experimentally induced iron overload have significantly enhanced Salmonella growth rates in all tissues examined (48, 49). Dysregulation of hepcidin, a peptide that normally limits excess iron absorption, leads to iron overload and increased susceptibility to Salmonella infection (50). The mechanism by which iron overload increases susceptibility to Salmonella infection remains to be fully resolved, but the results of previous studies and the current work indicate that iron overload may provide abundant stores of ferric iron for bacterial utilization.

There are at least two pathways by which Salmonella may access ferric iron in macrophages. First, ferric iron is transported to different tissues by the host protein transferrin, which binds cognate receptors on cells, including macrophages (51). Host transferrin receptor expression is increased in response to many intracellular pathogens, such as Francisella tularensis, Mycobacterium tuberculosis, and Ehrlichia (52–54). Upon acidification of transferrin-containing endosomes, ferric iron is released from transferrin and may be accessed by these pathogens. In Nramp1−/− macrophages, Salmonella does not require transferrin expression for successful intracellular survival (52). However, Nramp1+ macrophages limit Salmonella access to iron (20, 25). Therefore, Salmonella may, like other pathogens that live within vesicles, intercept transferrin-containing vesicles to obtain ferric iron. Ferric iron released from transferrin can alternatively be transported to the cytosol and stored within ferritin. Some cytosolic microbes, such as Neisseria meningitidis and Listeria monocytogenes, utilize ferritin as an iron source through mechanisms that directly degrade ferritin pools in the cytosol (55, 56). Since Salmonella is typically an intravascular pathogen in macrophages, it may not be able to directly access cytosolic ferritin. However, iron starvation and cellular stress induce autophagy of ferritin, followed by ferritin degradation in vesicles (57–59). Salmonella has been demonstrated to induce autophagy in macrophages (60, 61) and can also direct vesicular trafficking events (62). Thus, one hypothesis is that Salmonella facilitates fusion of the vesicle where it resides with ferritin-containing vesicles and then uses enterobactin or salmochelin to remove Fe3+ from the cytosolic storage protein ferritin. The addition of exogenous iron-loaded ferritin to Nramp1−/− macrophage-like cells increases Salmonella survival, consistent with this hypothesis (63).

Together, our studies combined with others highlight the particular importance of ferric iron in both nutritional immunity and Salmonella survival in the host. Salmonella has successfully evolved ferric iron-specific siderophores, enterobactin and salmochelin, that may be especially suited to take advantage of intracellular ferric iron storage pools. The current work also indicates that Salmonella may utilize siderophores differentially based upon tissue localization. Modulation of host iron regulators or the preferentially utilized Salmonella iron uptake systems has the potential to become pivotal in the development of new pharmacological approaches to control and/or prevent disease.

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REFERENCES


