Fructose-Asparagine Is a Primary Nutrient during Growth of Salmonella in the Inflamed Intestine

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Abstract

Salmonella enterica serovar Typhimurium (Salmonella) is one of the most significant food-borne pathogens affecting both humans and agriculture. We have determined that Salmonella encodes an uptake and utilization pathway specific for a novel nutrient, fructose-asparagine (F-Asn), which is essential for Salmonella fitness in the inflamed intestine (modeled using germ-free, streptomycin-treated, ex-germ-free with human microbiota, and IL10−/− mice). The locus encoding F-Asn utilization, fra, provides an advantage only if Salmonella can initiate inflammation and use tetrathionate as a terminal electron acceptor for anaerobic respiration (the fra phenotype is lost in Salmonella SPI1− SPI2− or trrA mutants, respectively). The severe fitness defect of a Salmonella fra mutant suggests that F-Asn is the primary nutrient utilized by Salmonella in the inflamed intestine and that this system provides a valuable target for novel therapies.

Introduction

Salmonella is a foodborne pathogen that causes significant morbidity and mortality in both developing and developed countries [1,2]. It is widely believed that there are no undiscovered drug targets in Salmonella enterica, largely due to the high number of nutrients available during infection and redundancy in metabolic pathways [3,4]. To acquire nutrients in the intestine, Salmonella initiates inflammation, which disrupts the microbiota and causes an oxidative burst that leads to the formation of tetrathionate [1–3,5–7]. Tetrathionate is used as a terminal electron acceptor for the anaerobic respiration of carbon compounds that otherwise would not be metabolized [8]. One of these carbon sources is ethanolamine, which is derived from host phospholipids. Ethanolamine can be respired by Salmonella, but not fermented [8]. Salmonella actively initiates inflammation using two Type 3 Secretion Systems (T3SS), each encoded within a distinct, horizontally acquired pathogenicity island. SPI1 (Salmonella Pathogenicity Island 1) contributes to invasion of host cells and elicitation of inflammation in the host. SPI2 is required for survival within macrophages and contributes to intestinal inflammation. Salmonella strains lacking SPI1 and SPI2 cause very little intestinal inflammation [5,6,8,9]. Here, we have identified fructose-asparagine (F-Asn) as another carbon source that is consumed by Salmonella using tetrathionate respiration during the host inflammatory response. The phenotypes of mutants lacking this utilization system are quite severe, suggesting that this is the primary nutrient utilized during Salmonella-mediated gastroenteritis. No other organism is known to synthesize or utilize F-Asn.

Results

The fructose-asparagine (F-Asn) utilization system was discovered during a genetic screen designed to identify novel microbial interactions between Salmonella and the normal microbiota. Transposon site hybridization (TraSH) was used to measure and compare the relative fitness of Salmonella transposon insertion mutants after oral inoculation and recovery from the cecum of two types of gnotobiotic mice, differing from each other by a single intestinal microbial species [10–15]. The two types of mice were germ-free and ex-germ-free colonized by a single member of the normal microbiota, Enterobacter cloacae. E. cloacae was chosen because it is a commensal isolate from our laboratory mice, easily cultured, genetically tractable, and it protects mice against
**Author Summary**

It has long been thought that the nutrient utilization systems of *Salmonella* would not make effective drug targets because there are simply too many nutrients available to *Salmonella* in the intestine. Surprisingly, we have discovered that *Salmonella* relies heavily on a single nutrient during growth in the inflamed intestine, fructose-asparagine (F-Asn). A mutant of *Salmonella* that cannot obtain F-Asn is severely attenuated, suggesting that F-Asn is the primary nutrient utilized by *Salmonella* during inflammation. No other organism has been reported to synthesize or utilize this novel biological compound. The novelty of this nutrient and the apparent lack of utilization systems in mammals and most other bacteria suggest that the F-Asn utilization system represents a specific and potent therapeutic target for *Salmonella*.

*Salmonella* infection (Figure 1). In total, five genes conferred a greater fitness defect in the mice containing *Enterobacter* than in the germ-free mice (Table 1).

Two of these genes, barA and sirA (uvrY), encode a two component response regulator pair that is conserved throughout the γ-proteobacteria [16–18]. BarA/SirA control the activity of the CorA protein (carbon storage regulator) which coordinates metabolism and virulence by binding to and regulating the translation and/or stability of mRNAs for numerous metabolic and virulence genes including SPI1, SPI2, and glgCAP (glycogen biosynthesis) [17,19,20]. To confirm the fitness phenotype of the BarA/SirA regulatory system, we performed competition experiments in which wild-type *Salmonella* was mixed in a 1:1 ratio with an isogenic *sirA* mutant and inoculated orally into germ-free mice and ex-germ-free mice colonized by *Enterobacter*. The results of TraSH analysis suggested that the *sirA* mutant would be at a greater growth disadvantage in *Enterobacter* mono-associated mice than in germ-free mice (Table 1). Results of the competition experiment confirmed this prediction (Figure 2).

The other three genes identified by TraSH analysis had not been characterized previously, and are located together in a putative operon. Genome annotation suggested that they encode a C4 dicarboxylate transporter, a sugar kinase, and a phosphosugar isomerase (Figure 3). A putative asparaginase lies at the end of the operon, and a separate gene upstream of the operon encodes a putative transcriptional regulator of the GntR family. These genes are not present in *E. coli* and appear to represent a horizontal acquisition inserted between the *fur* and *treF* genes at 77.7 centisomes of the *Salmonella* 14028 genome (ORFs STM14_4328 to STM14_4332). We have named these genes *fraBDAE* and *fraR* for reasons to be described below. A *fraB1:kan* mutation was constructed and tested for fitness in germ-free and *Enterobacter* mono-colonized mice using 1:1 competition assays against the wild-type *Salmonella*. The TraSH results suggested that this locus would exhibit a differential fitness phenotype in germ-free mice and *Enterobacter* mono-associated mice. Indeed, disruption of the *fra* locus caused a severe fitness defect in germ-free mice and a more severe defect in *Enterobacter*-colonized mice (Figure 4A, B).

**The fra locus confers a fitness advantage during inflammation and anaerobic respiration**

Competition experiments between wild-type and the *fraB1:kan* mutant were performed as described above using conventional mice (with normal microbiota) and mice treated orally with streptomycin (strep-treated) one day earlier to disrupt the microbiota (Figure 4C, D, E). Conventional mice do not become inflamed from *Salmonella*, while strep-treated mice (or germ-free) do become inflamed [5,6,8,21–24]. Disruption of the *fra* locus caused no fitness defect in conventional mice, but caused a severe defect in the strep-treated mice at one and four days post-infection (Figure 4C, D, E). The phenotype in strep-treated mice was confirmed by complementation (Figure 4F). It is expected that the *fraB1:kan* mutation is polar on the remainder of the *fraBDAE* operon. Therefore, the *fraB1:kan* mutation was complemented with a low copy number plasmid encoding the entire *fra* island (Figure 4F). The phenotype was confirmed again using a separately constructed mutation, *fraBE:kan*, and complementation (Figure 4G, H, I). In both instances, greater than 99% of the phenotype was restored (Figure 4F, I).

The observation of a phenotype in germ-free and strep-treated mice, but not conventional mice, suggested that *Salmonella* might require inflammation in order to acquire or utilize the *fra*-dependent nutrient source. It is known that inflammation causes the accumulation of tetrathionate in the lumen, a terminal electron acceptor that allows *Salmonella* to respire anaerobically [6]. Histopathology results confirmed that infection with *Salmonella* caused inflammation in the germ-free and strep-treated mice, but not in the conventional mice (Figure 5A, D, E). To test the hypothesis that *Salmonella* must induce inflammation for *fra* to affect the phenotype, we repeated the competition experiments in a *Salmonella* genetic background lacking SPI1 and SPI2, so that both the wild-type and the *fra* mutant would be defective for induction of inflammation. The severe fitness phenotype of the *fra* mutant was not observed in these strains (Figure 4J–L) and histopathology results confirmed that inflammation was indeed low during these experiments (Figure 5B, F).

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![Graph](image-url)

**Figure 1. Protection of mice against *Salmonella* serovar Typhimurium strain 14028 by *Enterobacter cloacae* strain JLD400.** Germ-free C57BL/6 mice were divided into two groups. One group was colonized with $10^7$ cfu of *Enterobacter cloacae* via the intragastric route (i.g.) and one group was not. One day later both groups were challenged i.g. with $10^7$ cfu of *Salmonella*. After 24 hours, the cecum and spleen were homogenized and plated to enumerate *Salmonella*. Each point represents the CFU/g recovered from one mouse with the geometric mean shown by a horizontal line. Statistical significance between select groups was determined by using an unpaired two-tailed Student t test. ** = P value < 0.01, *** = P value < 0.001.

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To test the hypothesis that tetrathionate respiration was required for use of the fra-dependent nutrient source, the competition experiments were repeated in a ttrA mutant background. TtrA is part of a tetrathionate reductase, which is required for the utilization of tetrathionate as a terminal electron acceptor during anaerobic respiration [6,25]. As in the SPI1 SPI2 background, there was no phenotype of a fra mutant in a ttrA mutant background indicating that Salmonella must be able to respire using tetrathionate to gain advantage from the fra locus (Figure 4M–O). Histopathology results confirmed the presence of moderate inflammation during these experiments (Figure 5C, G).

To determine if the fra locus is required during the systemic phase of disease, we performed competition experiments between the wild-type and fra mutant after intraperitoneal inoculation of conventional or strep-treated mice, with bacterial recovery from the spleen. The fra mutant had no fitness defect during systemic infection (Figure 4P, Q).

So far, we have seen the fra phenotype in C57BL/6 mice, which are mutated at the Nramp1 locus, and this required that the mice be either germ-free or strep-treated so that Salmonella could induce inflammation. Ideally, we would like to determine the significance of the fra locus in a model that is not mutated and does not require strep-treatment or a germ-free status. It is known that humans with a complete microbiota are quickly inflamed by Salmonella infection while conventional mice are not, and more recently it was discovered that germ-free mice colonized with human fecal microbiota (“humanized” mice) become inflamed from Salmonella infection without disturbance of the gut microbiota by streptomycin [26]. Therefore, we “humanized” germ-free Swiss Webster mice, which are Nramp1+/+, with human feces obtained from a healthy adult donor from the Ohio State University fecal transplant center. Competition experiments were then performed between wild-type and fra mutant Salmonella in these mice. Histopathology results confirmed the presence of mild inflammation during these experiments and the fra locus had a greater than 10,000-fold fitness phenotype (Figure 6).

IL10 knockout mice were used as another method to facilitate Salmonella-induced inflammation without using streptomycin [5]. Histopathology results indicated that, unexpectedly, there was not very much inflammation in these mice by day 3 post-infection although the fra locus still had a modest fitness phenotype (greater than 100-fold) (Figure 6). The phenotypes of the fra locus in IL10 knockout mice and in the humanized Swiss Webster mice demonstrate that the fra phenotype is not limited to germ-free or streptomycin-treated mice.

Finally, to test for the possibility that these severe fra mutant phenotypes were the result of interaction between the wild-type and fra mutant during infection, we performed experiments in which strep-treated C57BL/6 Nramp1+/− heterozygous mice were infected separately with the wild-type, the fra mutant, or the complemented fra mutant. The strains were quantitated in the

| Locus tag   | Symbol | Description                | Germ-free miceᵇ | Enterobacter monoassociated miceᵇ | Differenceᵇ |
|------------|--------|----------------------------|-----------------|----------------------------------|-------------|
| STM14_2365 | sirA   | response regulator         | 1.88            | -0.27                            | -2.15       |
| STM14_3566 | basA   | hybrid sensory histidine kinase | 1.09           | -0.55                            | -1.64       |
| STM14_4330 | fraD   | putative sugar kinase      | -0.07           | -1.29                            | -1.22       |
| STM14_4331 | fraB   | putative phosphosugar isomerase | 0.05           | -1.12                            | -1.18       |
| STM14_4329 | fraA   | putative transporter       | -0.06           | -1.23                            | -1.17       |

*a* The locus tag is from the Salmonella serovar Typhimurium strain 14028s genome (accession number NC_016856.1).
*b* The log2 hybridization intensity of this locus after recovery of the Salmonella library from germ-free mice.
*c* The log2 hybridization intensity of this locus after recovery of the Salmonella library from germ-free mice that had been previously monoassociated with Enterobacter cloacae.
*d* The difference in log2 hybridization intensity of this locus between Enterobacter monoassociated mice and germ-free mice.

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**Table 1.** Genes that are differentially required in germ-free mice and ex-germ-free mice monoassociated with *Enterobacter cloacae.*
Figure 3. Map of the fra locus of Salmonella enterica. The five genes of the fra locus are shown as grey arrows. The gor and tref genes are shown as black arrows and are conserved throughout the Enterobacteriaceae while the fra locus is not, suggesting that the fra locus was horizontally acquired. The proposed functions and names of each gene are shown below and above the arrows, respectively. The names are based upon the fraB1 homolog, while the fra locus does not have an asparaginase. Therefore, the name fraC was not used and the asparaginase was named fraE. The locus tags using the Salmonella nomenclature for strains 14028 (STM14 numbers) and LT2 (STM numbers) are shown above the gene names.

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Figure 4. Fitness defect of a fraB1::kan mutant as measured by competitive index (CI) in various genetic backgrounds and mouse models. A) 10^7 wild-type MA43 and fraB1::kan mutant MA59 in germ-free (GF) C57BL/6 mice, via the intragastric route (i.g.) and recovered from the cecum after 24 hours. B) 10^7 wild-type MA43 and fraB1::kan mutant MA59 in germ-free C57BL/6 mice mono-associated with Enterobacter cloacae, via the i.g. route and recovered from the cecum after 24 hours. C) 10^9 wild-type MA43 and fraB1::kan mutant MA59 in C57BL/6 conventional mice, via the i.g. route and recovered from the cecum after 24 hours. D) 10^7 wild-type IR715 and fraB1::kan mutant MA59 in streptomycin-treated (ST) C57BL/6 mice, via the i.g. route and recovered from the cecum after 4 days. E) 10^7 wild-type IR715 and fraB1::kan mutant MA59 in streptomycin-treated C57BL/6 mice, via the i.g. route and recovered from the cecum after 4 days. F) Complementation of the fraB1::kan mutation with a plasmid encoding the entire fra island, pASD5006. 10^7 ASD6090 and ASD6000 in streptomycin-treated C57BL/6 mice, via the i.g. route and recovered from the cecum after 4 days. G) 10^7 wild-type IR715 and fraB4::kan mutant MA5901 in streptomycin-treated C57BL/6 mice, via the i.g. route and recovered from the cecum after 24 hours. H) 10^7 wild-type IR715 and fraB4::kan mutant MA5901 in streptomycin-treated C57BL/6 mice, via the i.g. route and recovered from the cecum after 4 days. I) Complementation of the fraB4::kan mutation with a plasmid encoding the entire fra island, pASD5006. 10^7 wild-type ASD6090 and fraB4::kan mutant MA5901, both strains are in a SPI2 background, in streptomycin-treated C57BL/6 mice, via the i.g. route and recovered from the cecum after 4 days. J) 10^7 fraB1::kan mutant MA4310 and fraB1::kan mutant MA5901, both strains are in a SPI1 background, in streptomycin-treated C57BL/6 mice, via the i.g. route and recovered from the cecum after 4 days. K) 10^7 fraB1::kan mutant MA4310 and fraB1::kan mutant MA5901, both strains are in a SPI1 background, in streptomycin-treated C57BL/6 mice, via the i.g. route and recovered from the cecum after 4 days. L) 10^7 fraB1::kan mutant MA4310 vs fraB1::kan mutant MA5901, both strains are in a SPI1 background, in streptomycin-treated C57BL/6 mice, via the i.g. route and recovered from the cecum after 4 days. M) 10^7 fraB1::kan mutant MA4310 vs fraB1::kan mutant MA5910, both strains are in a SPI1 background, in streptomycin-treated C57BL/6 mice, via the i.g. route and recovered from the cecum after 24 hours. N) 10^7 fraB1::kan mutant MA4310 vs fraB1::kan mutant MA5910, both strains are in a SPI1 background, in streptomycin-treated C57BL/6 mice, via the i.g. route and recovered from the cecum after 4 days. O) 10^7 wild-type MA43 and fraB1::kan mutant MA59 in conventional C57BL/6 mice, via the intraperitoneal route (i.p.) and recovered from the spleen after 24 hours. P) 10^7 wild-type MA43 and fraB1::kan mutant MA59 in streptomycin-treated C57BL/6 mice, via the i.p. route and recovered from the spleen after 24 hours. Q) 10^7 wild-type MA43 and fraB1::kan mutant MA59 in streptomycin-treated C57BL/6 mice, via the i.p. route and recovered from the spleen after 24 hours. Each data point represents the CI from one mouse with the median shown by a horizontal line. Statistical significance of each group being different than 1 was determined by using a one sample Student’s t test. Statistical significance between select groups was determined using a Mann-Whitney test. * = P value < 0.05, ** = P value < 0.01, *** = P value < 0.001.

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feces each day post-infection for four days at which point the mice were sacrificed and the strains were quantitated in the cecum. The fraB1 mutant was recovered in 30-fold lower numbers than wild-type on the fourth day in the feces and 98-fold lower in the cecum (Figure 7). This defect was restored by complementation with the fra locus on a plasmid in the cecum, while in the feces the restoration did not reach statistical significance (Figure 7).

The fra locus is required for growth on fructose-asparagine (F-Asn)

FraA is homologous to the Dcu family of dicarboxylate transporters. However, authentic dicarboxylate acquisition loci do not encode a sugar kinase or phosphosugar isomerase. Furthermore, none of the dicarboxylates that we tested (malate, fumarate or succinate) provided a growth advantage to the wild-type strain vs. a fraB1::kan mutant, suggesting that they are not substrates of the Fra pathway. BLAST searches using the entire operon revealed that the closest homolog is the \( \text{fil} \) operon of \( E. \text{coli} \), although the \( \text{fil} \) operon is at a different location within the genome and does not encode an asparaginase (and the \( S. \text{salmonella} \) fra locus does not encode a \( \text{fil} \) homolog). The products of the \( E. \text{coli} \text{fil} \) operon transport and degrade the Amadori product fructose-lysine (F-Lys) [27,28]. Amadori products most often result from a spontaneous reaction between a carbonyl group (often of glucose, although numerous other compounds can also react) and an amino group of an amino acid in vivo, and are then referred to as non-enzymatic glycation products [29,30]. With F-Lys and fructose-arginine (F-Arg) this can happen with the free amino acid, or the side groups of the lysine and arginine residues of a protein. In contrast, fructose-asparagine (F-Asn) can only result from reaction of glucose with the alpha amino group of free asparagine or the N-terminal asparagine of a protein. We synthesized three different Amadori products, F-Lys, F-Arg, and F-Asn and used them as sole carbon sources during growth experiments. The preparations were free of glucose but contained some free amino acid. However, control experiments demonstrated that \( S. \text{salmonella} \) was unable to grow on any of the three amino acids alone, so these contaminants are inconsequential (Figure 8D). \( S. \text{salmonella} \) was unable to grow on F-Arg, and grew slowly and with low yield on F-Lys (Figure 8B, C). The growth on F-Lys was independent of the fra locus. In contrast, \( S. \text{salmonella} \) grew as well on F-Asn as on glucose, and growth on F-Asn was dependent upon the fra locus (hence the name fra, for fructose-asparagine utilization) (Figure 8A). A commercial source of F-Asn was obtained and it also allowed \( S. \text{salmonella} \) to grow in a fra-dependent manner (structure shown in Figure 8F). Complementation of the \( \text{fraB1}:\text{kan} \) mutant with a plasmid encoding the fra island restored the ability of the mutant to grow on F-Asn (Figure 8E). In addition to serving as a sole carbon source, F-Asn, also served as sole nitrogen source (Figure 9).

Growth with F-Asn was tested under aerobic and anaerobic conditions in the presence or absence of the terminal electron acceptor tetraionate (Figure 10). The F-Asn was utilized under all conditions, but respiratory conditions were superior with a doubling time of 1.6+/−0.1 hours aerobically with tetraionate, 2.0+/−0.3 hours aerobically without tetraionate, 1.9+/−0.1 hours anaerobically with tetraionate, and 2.9+/−0.4 hours anaerobically without tetraionate. Competition experiments in...
Figure 7. Quantitation of *Salmonella* in feces on days 1 through 4, and cecum on day 4, post-infection. Groups of five C57BL/6 mice heterozygous for *Nramp1* were orally inoculated with 10⁷ CFU of IR715 (wild-type), MA59 (fraB1::kan mutant), or ASD6000 (fraB1::kan mutant with complementation plasmid pASD5006). The geometric mean+SE is shown. Statistical significance between select groups was determined by using an unpaired two-tailed Student t test. * = P value < 0.05, ** = P value < 0.01.

Figure 8. Growth of wild-type and fraB1::kan mutant *Salmonella* on Amadori products. Growth of wild-type MA43 and fraB1::kan mutant MA59 on F-Asn (A), F-Arg (B), F-Lys (C), asparagine, arginine, lysine, or glucose (D). Bacteria were grown overnight in LB at 37°C shaking, centrifuged, resuspended in water, and subcultured 1:1000 into NCE medium containing the indicated carbon source at 5 mM. The optical density at 600 nm was then read at time points during growth at 37°C with shaking. Controls included NCE with no carbon source, and NCE with glucose that was not inoculated, as a sterility control (D). E) Complementation of a fraB1::kan mutation with plasmid pASD5006 encoding the fra island (ASD6000) or the vector control, pWSK29 (ASD6010). Each point in (A)–(E) represents the mean of three cultures with error bars indicating standard deviation. F) The structure of F-Asn (CAS # 34393-27-6).
which the wild-type and fraB1::kan mutant were grown in the same culture were performed in minimal medium containing F-Asn. As expected, the mutant was severely attenuated during aerobic and anaerobic growth, and in the presence or absence of tetrathionate (Figure 11). The attenuation was most severe during anaerobic growth in the presence of tetrathionate.

Discussion

The mechanisms by which microbes interact with each other in the gastrointestinal tract are largely unknown. Screening large libraries of bacterial mutants for fitness defects in animals with defined microbiota can be used to identify those genes that are only required in the presence of specific members of the microbiota [15]. In this report, we took a highly reductionist approach and screened for genes that were differentially required in germ-free mice versus ex-germ-free mice colonized with a single commensal Enterobacter cloacae isolate. Only five genes were differentially required, a two component response regulatory pair, barA/sirA, and three genes within the fra locus (Table 1). Individual sirA and fraB mutants were used to confirm the findings. The sirA gene was required for fitness in the presence of E. cloacae but not in its absence (Figure 2). The fra locus was required for fitness in both situations, but the phenotype was more severe in the presence of E. cloacae (Figure 4A, B). Thus, the differential screening strategy was successful in identifying genes that are more important in the presence of other bacteria within the gastrointestinal tract. The reason(s) that sirA is required in the presence, but not the absence, of E. cloacae is not known. It is thought that BarA detects short chain fatty acids produced by the normal microbiota and then phosphorylates SirA [31–34]. SirA then activates the transcription of two small RNAs, csrB and csrC, which antagonize the activity of the CsrA protein [20,35–39]. The CsrA protein is an RNA-binding protein that regulates the stability and translation of hundreds of mRNAs involved with metabolism and virulence [17,19,40]. One possible reason that sirA differentially affects fitness in the two mouse models may be that the Enterobacter-colonized mouse offers an environment richer in carboxylic acids that act as stimuli for BarA-SirA signaling with resulting effects on metabolism and growth [31–34]. The fitness effects could also be due to the regulation of genes involved in the induction of inflammation and/or anaerobic metabolism including SPI1, SPI2, ethanolamine utilization, and vitamin B12 biosynthesis by CsrA [19,20,41–44]. Finally, SirA or CsrA may regulate the fra locus itself.

The fra locus was annotated as a C4 dicarboxylate uptake system. However, we found that the fra locus played no role in the utilization of C4 dicarboxylates. BLAST searches revealed that the operon is similar to the fil locus of E. coli which is required for the utilization of fructose-lysine (F-Lys). The fil locus of E. coli has a different genomic context than the fra locus of Salmonella, and is only distantly related. We determined that the fra locus of Salmonella plays no role in the utilization of F-Lys (Figure 8C). However, the presence of an asparaginase in the fra locus (fraE), but not the fil locus, led us to hypothesize that F-Asn may be the correct nutrient, and indeed, this was the case. Wild-type Salmonella is able to grow as well on F-Asn as on glucose, and this ability is

![Figure 9. Growth of Salmonella on F-Asn as sole nitrogen source.](image)

Growth of wild-type MA43 and fraB1::kan mutant MA59 on F-Asn. Bacteria were grown overnight in LB at 37°C shaking, centrifuged, resuspended in water, and subcultured 1:1000 into NCE medium lacking a nitrogen source (NCE-N) but containing the indicated carbon source at 5 mM. The optical density at 600 nm was then read at time points during growth at 37°C with shaking. Controls included NCE-N with no carbon source, NCE-N with 5 mM glucose, and NCE-N with glucose that was not inoculated, as a sterility control. Each point represents the mean of four cultures and error bars represent standard deviation.

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dependent upon the \textit{fim} locus (Figures 8, 10). While the individual members of the \textit{fim} operon have not been characterized, we hypothesize as to their functions in Figure 12. F-Asn differs from ethanolamine in that it can be fermented (Figure 10B), which would be consistent with the proposed release of glucose-6-P by \textit{FraB} (Figure 12). Although F-Asn can be fermented, it only provides a fitness advantage \textit{in vivo} when it can be respired, i.e., when tetrahionate reductase is functional (Figure 4M–O), possibly because of the much greater energy yield from respiration versus fermentation. \textit{E. cloacae} grows very poorly on F-Asn and does not encode the \textit{fra} locus. Therefore, \textit{E. cloacae} likely exacerbated the \textit{fra} phenotype of \textit{Salmonella} by competing for other nutrients.

F-Asn is an Amadori compound (also known as a glycation product) formed by reaction between glucose in its open chain form with the alpha amino group of asparagine followed by a rearrangement that gives the fructose derivative. Until this report, no organism had been shown to synthesize or utilize F-Asn. However, in the early 2000s it was discovered that acrylamide is present in many foods, especially French fries and potato chips. F-Asn is a precursor to acrylamide. After the acrylamide discovery, numerous papers measured acrylamide concentration, and the precursor molecules, glucose and asparagine, in foods [45–52]. However, to the best of our knowledge, only two reports have measured the concentration of F-Asn in a few fruits and vegetables [53,54]. The concentrations are surprisingly high, ranging between 0.1% (carrot) and 1.4% dry weight (asparagus) [54]. Factors that influence these concentrations are time, temperature, pressure, and perhaps less obviously, moisture content [55]. Any reducing sugar and any amino acid (or other amines) can form compounds analogous to F-Asn. It is important to note that these Amadori compounds are not the ultimate products since with further time and heating they decompose to a large variety of

\textbf{Figure 10. Growth of \textit{Salmonella} on F-Asn in the presence or absence of tetrahionate or oxygen.} Growth of wild-type MA43 and \textit{fraB}::kan mutant MA59 on 5 mM F-Asn or 5 mM glucose anaerobically (A and B) or aerobically (C and D) in the presence (A and C) or absence (B and D) of 40 mM tetrahionate (S\textsubscript{4}O\textsubscript{6}\textsuperscript{2-}). Bacteria were grown overnight in LB at 37 °C shaking, centrifuged, resuspended in water, and subcultured 1:1000 into NCE medium containing the indicated carbon source. The optical density at 600 nm was then read at time points during growth at 37 °C with shaking. Each point represents the mean of four cultures with error bars indicating standard deviation. doi:10.1371/journal.ppat.1004209.g010
other products, some of which are responsible for a variety of flavors, and the brown color, in cooked foods [35–37]. In fact, glycation products form spontaneously in the human body and flavors, and the brown color, in cooked foods [55–57]. In fact, glycation products form spontaneously in the human body and provide an indication of glucose concentration over time [30,58–60]. A common diabetes test measures the glycation of the N-terminal valine of hemoglobin [30].

The severity of the fra fitness phenotype suggests that F-Asn is the primary nutrient used by Salmonella during growth in the inflamed intestine. For perspective, in strep-treated mice the fitness defect of a fra mutant is 1000-fold, while mutants unable to utilize ethanolamine or sialic acid are attenuated 10-fold and 2-fold, respectively [8,61]. The fra operon was previously identified by transcription profiling as up-regulated by Fur under anaerobic conditions [62]. Other genes activated under the same conditions included ethanolamine utilization (eux), and hilA, a regulator of SPI1 expression. Both of these loci are associated with induction of inflammation or growth during inflammation [8,62]. The fra locus is present among most Salmonella serovars, but is disrupted in serovars Typhi and Paratyphi A, consistent with the marked degradation of numerous loci involved with anaerobic respiration among these extra-intestinal serovars [63]. Interestingly, a putative fra locus is present in Citrobacter rodentium and Citrobacter freundii, but not in numerous other non-pathogenic Citrobacter species. The fil locus, encoding the ability to utilize F-Lys is present in E. coli, Shigella, and Cronobacter. It will be interesting to determine which, if any, members of the normal microbiota can compete with E. coli and Salmonella for Amadori products.

The apparent species-specificity of the F-Asn utilization system, and the severity of the fitness defect associated with mutants that cannot metabolize F-Asn, indicate that the Fra system represents a specific and valuable therapeutic target. Further studies are needed to determine the role of each gene in the fra locus with regard to F-Asn metabolism. Similarly, further studies are needed to determine the mechanism by which the proposed transcription factor, FraR, regulates F-Asn metabolism. These structure-function studies will facilitate small molecule drug screens targeting F-Asn utilization. It will also be interesting to determine the concentration of F-Asn and other Amadori products in a wide variety of foods, to determine if these products can affect disease susceptibility, and to explore the possibility of preventing salmonellosis or other infections by removing Amadori products from specific food products or from the diet in general. The utilization systems for many more Amadori products are likely awaiting discovery within bacterial genomes and these may play interesting roles in microbial ecology and human health.

Materials and Methods

Bacterial strains and media

Bacteria were grown in Luria-Bertani (LB) broth or on LB agar plates (EM Science) unless otherwise noted. The minimal medium used was NCE (no carbon E) [64] containing trace metals [25]. Chloramphenicol (cam), streptomycin (strepl), or kanamycin (kan) were added at 30, 200, or 60 μg/ml, respectively, when appropriate. Fructose-asparagine was either synthesized or purchased from Toronto Research Chemicals, catalog #F792525.

Anaerobic growth was performed in a Bactron 1 anaerobic chamber containing 90% N2, 5% CO2, and 5% H2 (Shel Lab). Strains used are described in Table 2. Enterobacter cloacae strain JLD400 was isolated in our laboratory by plating fecal samples from a conventional BALB/c mouse onto LB agar plates. This particular isolate was chosen because it is easy to culture and genetically manipulable (the strain can be electroporated, maintains ColE1-based plasmids, and can act as a recipient in RP4-mediated mobilization of a suicide vector used to deliver mTn5-lux(CDABE, not shown). The species identification was performed using a Dade Microscan Walkaway 96si at the Ohio State University medical center. Additionally, genomic DNA sequences have been obtained that flank mTn5-luxCDABE insertions in JLD400 and these DNA sequences match the draft genome sequence of E. cloacae NCTC 9394.

Salmonella mutant library

A transposon mutant library was constructed in S. enterica serovar Typhiimmurium strain 14028. EZ-Tn5 <T7/kan> transposomes from Epicentre Technologies were delivered to Salmonella by electroporation. This transposon encodes kanamycin resistance and has a T7 RNA Polymerase promoter at the edge of the transposon pointed outward. The resulting library contains between 190,000 and 200,000 independent transposon insertions and is referred to as the JLD200k library. The insertion points of this library have been determined previously by next-generation sequencing [65]. It is estimated that approximately 4400 of the 4800 genes in the Salmonella genome are non-essential with regard to growth on LB agar plates [65]. Therefore, the JLD200k library...
is saturated with each gene having an average of 43 independent transposon insertions.

**Construction of mutations**

A FRT-kan-FRT or FRT-cam-FRT cassette, generated using PCR with the primers listed in Table 3 and pKD3 or pKD4 as template, was inserted into each gene of interest (replacing all but the first ten and last ten codons) using lambda Red mutagenesis of strain 14028+ followed by growth at 37°C to remove the plasmid [66]. A temperature sensitive plasmid encoding FLP recombinase, pCP20, was then added to each strain to remove the antibiotic resistance marker [66]. The pCP20 plasmid was cured by growth at 37°C. A fraB4::kan mutation was constructed using primers BA2552 and BA2553 (Table 3). A FRT-cam-FRT was placed in an intergenic region downstream of pagC using primers BA1561 and BA1562 (deleting and inserting between nucleotides 1342878 and 1343056 of the 14028 genome sequence (accession number NC_016856.1) (Table 3).

**Animals**

Germ-free C57BL/6 mice were obtained from Balfour Sartor of the NIH gnotobiotic resource facility at the University of North Carolina and from Kate Eaton at the University of Michigan. Germ-free Swiss Webster mice were obtained from Taconic Farms. The mice were bred and maintained under germ-free conditions in sterile isolators (Park Bioservices). Periodic Gram-staining, 16s PCR, and pathology tests performed by the Ohio State University lab animal resources department and our own laboratory were used to confirm that the mice contained no detectable microorganisms. Conventional C57BL/6 mice were obtained from Taconic Farms. C57BL/6 mice that were heterozygous for the Nramp1 gene were generated by breeding the standard Nramp1+/+ mice from Greg Barton [67]. IL10 knockout mice (B6.129P2-IL10tm1Cgn/J) were obtained from Jackson Laboratory. Germ-free Swiss Webster mice were “humanized” by intragastric inoculation of 200 μl of human feces obtained from an anonymous healthy donor from the OSU fecal transplant center.

**Transposon Site Hybridization (TraSH)**

The JLD200k transposon mutant library was grown in germ-free C57BL/6 mice in the presence or absence of E. cloacae strain JLD400. Four mice were inoculated intragastrically (i.g.) with 10⁷ cfu of Enterobacter cloacae strain JLD400 that had been grown overnight in LB shaking at 37°C. After 24 hours these mice, and an additional four germ-free mice, were inoculated with 10⁷ cfu of the JLD200k library that had been grown overnight in shaking LB kan at 37°C. Prior to inoculation of the mice, the library was spiked with an additional mutant, JLD1214, at a 1:10:000 ratio. This mutant contains a chloramphenicol resistance (cam) gene at a neutral location in the chromosome in the intergenic region downstream of pagC [68]. After inoculation of mice with the spiked library, the inoculum was dilution plated to quantitate the kanamycin resistant (kan) Salmonella library members and the cam spike strain. The remainder of the inoculum was pelleted and saved as the “input” for hybridization to microarrays.
24 hours of infection with the JLD200k library, the mice were euthanized and organs were harvested (small intestine, cecum, large intestine, and spleen). One germ-free mouse died prior to organ harvest and was not used. All samples were homogenized and diluted plated to determine Salmonella counts. The remainder of the homogenate was added to 25 ml LB kan and grown overnight with shaking at 37°C to recover the library members. Each culture was then pelleted and frozen as a potential “output” sample for microarray analysis. The kan’ and cam’ colony counts recovered from each organ indicated that the spike ratio of 1:10,000 was maintained in the intestinal samples but not in the spleen samples. This indicates that the library underwent a population bottleneck on the way to the spleen so microarray analysis of spleen samples would not be informative. The cecum samples were chosen for microarray analysis. There was one “input” sample for all arrays. There were seven separate “output” samples for the arrays; four from the cecums of Enterobacter-associated mice and three from germ-free mice. The output from each mouse was compared to the input on a single array. We also did a single “in vitro” array experiment in which the JLD200k library was grown in the presence of Enterobacter in liquid LB broth at 37°C. Genomic DNA was isolated from the input and output bacterial pellets. The purity and concentration of the DNA samples was assessed using a Nanodrop spectrophotometer and the quality of the DNA was assessed via agarose gel electrophoresis. All seven samples had high quality intact genomic DNA. The DNA was digested using a restriction endonuclease (RsaI). Labeled RNA transcripts were obtained from the T7 promoter by in vitro transcription. A two-color hybridization strategy was employed. RNA transcripts from the output samples were fluorescently labeled with Cyanine-3 (Cy3, green) and the input sample was labeled with Cyanine-5 (Cy5, red), while the input sample was labeled with Cyanine-3 (Cy3, green). Equal molar concentrations of the output and input sample were combined and hybridized to genome-wide tiling microarrays printed commercially by Agilent Technologies. Agilent’s SurePrint technology employs phosphoramidite chemistry in combination with high performance Hewlett Packard inkjet technology for in situ synthesis of 60-mer oligos. Using Agilent eArray, an easy-to-use web-based application, we were able to synthesize the arrays used by Chaudhuri et al. that completely tiled both the sense and anti-sense strands of the Salmonella SL1344 genome (AMADID 015511) [10]. Each slide contained 2 arrays, each array with 105,000 features, densely tiling the entire genome. The strain of Salmonella used in our experiments was 14028 and its genome sequence was only recently published (GenBank Nucleotide Accession CP001363 (complete genome) and CP001362 (plasmid)). As such, each of the 60-mer probes used by Chaudhuri et al. [10] were mapped to the 14028 genome using

| Strain or plasmid | Genotype or description | Source or reference |
|-------------------|------------------------|---------------------|
| 14028             | wild-type Salmonella enterica serovar Typhimurium | American Type Culture Collection |
| ASD6000           | MA59 fraB1::kan+pASD5006 amp’| This study |
| ASD6010           | MA59 fraB1::kan+pWSK29 (amp’)| This study |
| ASD6040           | MA59 fraB1::kan+pASD5006 (amp’)| This study |
| ASD6090           | MA59 fraB1::kan+pASD5006 (amp’)| This study |
| IR715             | 14028 nai’| This study |
| JLD400            | wild-type Enterobacter cloacae isolated from a laboratory mouse | This study |
| JLD1214           | 14028 IGPagC-STM14_1502::cam | lambda red mutation downstream of pagC created using PCR primers BA1561 and BA1562, then transduced into 14028 |
| MA43              | IR715 phoN1::saudA | phoN1::str mutation from Helene Andrews-Polymenis collection transduced into IR715 |
| MA45              | IR715 sirA2::kan | IR715 transduced with P22 grown on BA736 [44,72] |
| MA59              | IR715 fraB1::kan | fraB1::kan mutation from Helene Andrews-Polymenis collection transduced into IR715 |
| CS1032            | IR715 fraB4::kan | lambda red mutation of fraB, created using PCR primers BA2552 and BA2553, then transduced into IR715. |
| MA4301            | 14028 (avrA-invH1) ssaK::kan | ssaK::kan from Micah Worley strain MJW1836 transduced into YD039 [46,81] |
| MA4310            | MA43 ttrA1::cam | ttrA1::cam mutation from Helene Andrews-Polymenis collection transduced into MA43 |
| MA5900            | 14028 (avrA-invH1) ssaK::kan fraB1::cam | fraB1::cam mutation from Helene Andrews-Polymenis collection transduced into MA5901 |
| MA5910            | IR715 fraB1::kan ttrA1::cam | ttrA1::cam mutation from Helene Andrews-Polymenis collection transduced into MA59 |
| pASD5006          | pWSK29 fraRBDAE::amp’| This study |
| pWSK29            | pSC101 cloning vector amp’| [82] |
| pCP20             | cI857 P_r-f-frp pSC101 oriTS amp’ cam’| [66] |
| pKD3              | FRT-cam-FRT oriR6K amp’| [66] |
| pKD4              | FRT-kan-FRT oriR6K amp’| [66] |

Table 2. Bacterial strains and plasmids.
blast, and then annotated with any open reading frames (ORFs) that the probe spanned. A total of 96,749 probes mapped to the 14028 genome, with a median gap between each probe of 35 nucleotides on both strands.

After purification, the labeled samples were denatured and hybridized to the array overnight. Microarray slides were then washed and scanned with an Agilent G2505C Microarray Scanner, at 2 μm resolution. Images were analyzed with Feature Extraction 10.5 (Agilent Technologies, CA). Median foreground intensities were obtained for each spot and imported into the mathematical software package “R”, which was used for all data input, diagnostic plots, normalization and quality checking steps of the analysis process using scripts developed specifically for this analysis. In outline, the intensities were not background corrected as this has been shown to only introduce noise. The dataset was filtered to remove positive control elements and any elements that had been flagged as bad, or not present in the 14028 genome. Using the negative controls on the arrays, the background threshold was determined and all values less than this value were flagged. Finally, the Log2 ratio of output Cy5/input Cy3 (red/green) was determined for each replicate, and the data was normalized by the loess method using the LIMMA (Linear models for microarray data) package in “R” as described [69,70]. Complete statistical analysis was then performed in “R”. Insertion mutants where the ORF is essential for survival will be selected against, and thus a negative ratio of Cy5/Cy3 (red/green) will be observed in the probes adjacent to the insertion point, resulting from higher Cy3 (green) signal from the input. Conversely, insertion mutants that were advantageous to growth in the output samples would have a positive ratio, resulting from the higher Cy5 (red) signal in the output. Mutants having no effect on growth would have equal ratios in both the output and input samples (yellow). A spreadsheet of these data is available in Dataset S1.

Synthesis of Amadori products

We carried out the syntheses of three fructosyl amino acids with asparagine, lysine, and arginine. Hodge and Fisher’s review of Amadori products was consulted as an essential starting point for synthesis [71] and the recent review by Mossine and Mawhinney of all aspects of fructose-amines was a treasure house of information [55]. We found the method of Wang et al. [72] to be the most satisfactory, however reaction times cannot be standardized and excess glucose must be removed. The reaction with asparagine is slow because asparagine is sparingly soluble in methanol. By contrast, the reaction with 2-Boc-lysine is fast. Arginine is an intermediate case. Previous syntheses of F-Asn

| Table 3. Oligonucleotides used. |
|----------------------------------|
| **Gene targeted** | **Primer name** | **Description** | **Sequence** |
| pagC | BA1561 | Used for lambda red mutagenesis in which the cat (cam') gene was placed downstream of pagC in a neutral site using pKD3 as PCR template. | CTTCTTTACCCAGTACACGGGTATCGCTGTTTCTTCTTTGTAGGCTGGAGCTGGCTTCG |
| pagC | BA1562 | Used for lambda red mutagenesis in which the cat (cam') gene was placed downstream of pagC in a neutral site using pKD3 as PCR template. | CGAAGGCGGTCAAAATCTTTGATGACATTGGATTAACATATAGAATATCCTTCCTTAG |
| fra island | BA2228 | Used for amplifying the fra island and cloning it into a complementation vector, resulting in pASD5006. | GCAGAATCTATCCGTCGACACGAAC |
| fra island | BA2229 | Used for amplifying the fra island and cloning it into a complementation vector, resulting in pASD5006. | GCAGGTTAAGGCTCTCCGTTAAGGCAATC |
| fraB | BA2552 | Used for lambda red mutagenesis in which the aph (kan') gene was placed within the fraB gene using pKD4 as PCR template. | CCTGATGTAATATATCCTTCCACATATAGCGCCATATGAATATCCTTCCTTAG |
| fraB | BA2553 | Used for lambda red mutagenesis in which the aph (kan') gene was placed within the fraB gene using pKD4 as PCR template. | AGAGGAAAGCAGTATGGGTATGAAGAGACAGTTGCAATGTAAGGCTGGAGCCTTC |

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include those of Stadler et al. [46], Wang et al. [72], and Miura et al. [73]. The procedure of Stadler et al. [46] uses alkaline conditions which we thought could bring about isomerization of the sugar and racemization of the amino acid. We chose to develop the synthesis of Wang et al. [72] after trying a number of different protocols described for other amino acids [74–77]. Wang et al. [72], however, describe only a general method and asparagine presents some particular problems, the most important of which is the poor solubility of asparagine in methanol. We added bisulphite to the reaction mixture to reduce the formation of colored by-products [57] and finally removed excess glucose by use of a cation-exchange column according to the method of Mossine et al. [78]. Using methanol alone as solvent gives the product after refluxing for 24 hr. in approximately 10–15% yield together with recovery of about 90% of the asparagine. Although the yield is low, the starting materials are inexpensive, and the insolubility of asparagine has the advantage that F-Asn, which is quite soluble in methanol, emerges from the ion exchange column almost free of asparagine. This gave a free-flowing off-white non-hygroscopic solid. The $^1$H-NMR spectrum is complex due to the equilibrating mixture of alpha- and beta- pyranose and furanose forms [55], but integration of the upfield resonances due to asparagine and the downfield resonances due to the sugar are in the proper ratio. The material was also characterized by its specific rotation and infrared (IR) spectrum: $[\alpha]_D^23+48$° ($c$ = 0.1, water) (reference [73] $-40$°, $c$ = 1, water); IR (Nujol): 3350, 3155, 1668, 1633, 1455, 1408, 1080 cm$^{-1}$. Compare our preparations to results in [71,73].

Competition assays

Competition assays were performed in which a mutant strain was mixed in a 1:1 ratio with an isogenic wild-type and inoculated by the intragastric (i.g.) or intraperitoneal (i.p.) route to mice. Fecal samples, intestinal sections, spleen and liver were recovered at specific times post-infection, homogenized and plated on selective plates. The wild-type and mutant strains were differentiated by antibiotic resistance. The competitive index was calculated as $CI = (\text{cfu of mutant recovered}/\text{cfu w.t. recovered})/(\text{cfu mutant input}/\text{cfu w.t. input})$. If the mutant is defective compared to the wild-type it will have a CI of less than 1.

Complementation assays

The fra island was PCR amplified from purified 14028 genomic DNA with primers BA2228 and BA2229 using Phusion polymerase (New England Biolabs). The PCR product was cloned into pPCR-Blunt II-TOPO (Invitrogen). The resulting clones were digested with EcoRI (New England Biolabs), run on an agarose gel and the 8.6 kbp fra fragment was gel purified (Qiagen). This purified DNA fragment was ligated into pWSK29 digested with EcoRI (NEB) using T4 DNA ligase (New England Biolabs) overnight at 4°C. The ligation reaction was transformed into DH5α and plated on LB containing ampicillin at 37°C. The resulting plasmid, pASD5006, or the vector control pWSK29, were electroporated into the appropriate strains.

Ethics statement

All animal work was performed in accordance with the protocols approved by our Institutional Animal Care and Use Committee (OSU 2009A0035). The IACUC ensures compliance of this protocol with the U.S Animal Welfare Act, Guide for Care and Use of Laboratory Animals and Public Health Service Policy on Humane Care and Use of Laboratory Animals. Human fecal material was obtained from an anonymous healthy donor at the Ohio State University fecal transplant center in accordance with the protocol approved by our Institutional Review Board (OSU 2012H0367).

Supporting Information

Dataset S1  Transposon Site Hybridization data from germ-free mice and germ-free mice monoassociated with Enterobacter Cloacae. As explained more fully in the Materials and Methods, a normalized Log2 ratio of output/input hybridization intensity was determined for each replicate. Insertion mutants where the ORF is essential for survival were selected against, and thus yielded a negative ratio in the probes adjacent to the insertion point. Conversely, insertion mutants that were advantageous to growth in the output samples yielded a positive ratio. The average ratio for all probes and all replicates for each locus are shown in the spreadsheet for germ-free mice and germ-free mice mono-associated with Enterobacter cloacae. The difference column shows the difference of the ratios for that locus between the two mouse groups to facilitate the identification of differentially required genes. The spreadsheet has two tabs, one sorted by locus tag and one sorted by difference. (XLSX)

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Author Contributions

Conceived and designed the experiments: MMA DLN JFB PW BMMA. Performed the experiments: MMA DLN JFB ASD CS BS JD JNS. Analyzed the data: MMA DLN JFG ASD CS BMMA. Contributed to the writing of the manuscript: MMA DLN JFG ASD BS TR EJB PW BMMA.

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