A putative P-type ATPase required for virulence and resistance to haem toxicity in *Listeria monocytogenes*

Heather P. McLaughlin¹†, Qiaobin Xiao³†, Rosemarie B. Rea¹,⁴, Hualiang Pi³, Pat G. Casey¹,
Trevor Darby¹, Alain Charbit⁵,⁶, Roy D. Sleator⁴, Susan A. Joyce¹, Richard E. Cowart⁷, Colin
Hill¹*, Phillip E Klebba³‡ and Cormac G.M. Gahan¹,²

¹Alimentary Pharmabiotic Centre, Department of Microbiology, University College Cork, Cork,
Ireland

²School of Pharmacy, University College Cork, Cork, Ireland

³Department of Chemistry and Biochemistry, University of Oklahoma, Norman, Oklahoma
73019, USA

⁴Department of Biological Sciences, Cork Institute of Technology, Cork, Ireland

⁵Université Paris Descartes, Faculté de Médecine Necker-Enfants Malades, Paris, France

⁶INSERM, U1002, Unité de Pathogénie des Infections Systémiques, Paris, France

⁷Division of Biological Science, Department of Natural and Applied Sciences, University of
Dubuque, Dubuque, Iowa  52001, USA

¹Heather P Mc Laughlin and Qiaobin Xiao contributed equally to this work

¹Current address: David Geffen School of Medicine at UCLA, 6222 MacDonald Research
Laboratories, 675 Charles E. Young Drive South, Los Angeles, CA 90095

*Corresponding Author:* Prof. Colin Hill, Department of Microbiology, University College Cork,
Cork, Ireland.

Email: chill@ucc.ie  Telephone: ++353-21-4901373  Fax: ++353-21-4903101

Running title: Fur-regulated virulence factor in *L. monocytogenes*
Abstract

Regulation of iron homeostasis in many pathogens is principally mediated by the ferric uptake regulator, Fur. Since acquisition of iron from the host is essential for the intracellular pathogen *Listeria monocytogenes*, we predicted the existence of Fur-regulated systems that support infection. We examined the contribution of nine Fur-regulated loci to the pathogenicity of *L. monocytogenes* in a murine model of infection. While mutating the majority of the genes failed to affect virulence, three mutants exhibited a significantly compromised virulence potential. Most striking was the role of the membrane protein we designate FrvA (*Fur* regulated virulence factor A; encoded by *frvA* [lmo0641]), which is absolutely required for the systemic phase of infection in mice and also for virulence in an alternative infection model, the Wax Moth *Galleria mellonella*. Further analysis of the ΔfrvA mutant revealed poor growth in iron deficient media and inhibition of growth by micromolar concentrations of haem or haemoglobin, a phenotype which may contribute to the attenuated growth of this mutant during infection. Uptake studies indicated that the ΔfrvA mutant is unaffected in the uptake of ferric citrate but demonstrates a significant increase in uptake of haem and haemin. The data suggest a potential role for FrvA as a haem exporter that functions, at least in part, to protect the cell against the potential toxicity of free haem.
Introduction

Iron is indispensable for the growth of most bacteria, serving as a cofactor for enzymes involved in essential metabolic pathways such as glycolysis, DNA synthesis, energy generation, and detoxification of oxygen radicals [1,2]. The correlation between iron acquisition and bacterial virulence has been well documented [3,4,5] and the absolute requirement for this metal for both host metabolism and bacterial growth results in significant competition for iron in the host [6]. Following bacterial infection host responses are evoked which sequester iron, making it relatively unavailable for bacterial metabolism [7].

In the Gram positive intracellular pathogen Listeria monocytogenes, iron deficient environments have been shown to upregulate the expression of the principal virulence regulator PrfA and significantly increase the production of the haemolysin Listerialysin O promoting phagosomal escape, and the actin polymerisation protein ActA which plays a role in cell-to-cell spread [8,9,10]. It has been hypothesized that the requirement for iron has played a part in driving the evolution of an intracellular life-cycle for L. monocytogenes as the bacterium can utilize the iron-saturated protein ferritin stored in the cytosol of host cells (as reviewed by McLaughlin et al. [11]).

As iron-limiting conditions can be encountered in both the natural environment and during host infection, free-living pathogenic bacteria such as L. monocytogenes have evolved mechanisms to acquire iron from a variety of sources. Iron acquisition is mediated by a number of distinct systems that have been characterized in L. monocytogenes: a citrate inducible receptor for the uptake of ferric citrate, utilization of exogenous siderophores, catechol siderophore-like molecules, and catecholamine complexes, and iron acquisition via a cell-surface transferrin-binding protein [12]. A comprehensive analysis of the iron acquisition systems in L. monocytogenes identified a variety of iron sources which can be used for growth, including eukaryotic iron-binding proteins (haemoglobin, ferritin, transferrin and lactoferrin), ferric siderophores (enterobactin and corynebactin) and iron complexes of hydroximates (ferrichrome, ferrichrome A, and ferrioxamine B) [2]. In addition, the same study also identified two genetic loci responsible for the uptake of ferric hydroximates and haemin/haemoglobin.
Deletions in fluD or fluC strongly reduced ferrichrome uptake and a deletion in hupC eliminated uptake of haem and haemoglobin and resulted in decreased virulence potential [2]. However, it is clear that many other loci putatively involved in iron homeostasis in L. monocytogenes remain to be characterized by functional genetics approaches [13,14].

Maintaining a balanced acquisition of iron from the external environment is essential for bacterial survival. Whilst pathogens must compete for iron during infection, excess intracellular iron can lead to the generation of toxic hydroxyl radicals via the Fenton reaction. Iron homeostasis in most bacteria, including L. monocytogenes, is controlled by the regulatory protein Fur (ferric uptake regulator) or a functional equivalent [15]. In the presence of sufficient levels of iron, Fur acts as a repressor whereby an iron-Fur complex prevents gene transcription by binding to specific Fur-box sequences upstream of the start codon of target genes [16].

Recently Ledala et al. [17] used DNA microarray analysis to examine gene expression in a Fur mutant and identified Fur-regulated genes in L. monocytogenes, including genes encoding iron transporters and proteins involved in iron storage. In this study, we undertook an independent genome-wide search to identify putative Fur-box consensus sequences in the genome of L. monocytogenes. This approach identified a number of candidate Fur-regulated loci, including some (such as lmo0641) that were not identified previously through microarray analysis [17]. We undertook a systematic functional analysis of selected Fur-regulated loci by creating plasmid-insertion mutants and subsequently testing these for virulence potential in the murine model. This led to the identification of Fur-regulated virulence factor A, FrvA (encoded by frvA/lmo0641), a novel Fur regulated virulence factor which is absolutely required for growth of L. monocytogenes under restricted iron conditions and for systemic infection. We carried out iron uptake studies on the frvA mutant and determined that the mutant demonstrates a significant increase in uptake of haem and is also sensitive to elevated haem concentrations. Sensitivity to haem toxicity may account for the significant attenuation of virulence during the systemic phase of infection in the murine infection model.
Results and Discussion

In silico identification of putative Fur regulated genes

Fur has been identified as a major regulator of iron homeostasis in numerous Gram-positive and Gram-negative bacteria [16,18,19]. Regulation of iron uptake is particularly important during infection as pathogens must scavenge iron from sources in the host organism. Indeed, deregulation of iron uptake through elimination of Fur has been shown to significantly impact upon virulence potential in a number of pathogenic bacteria, including *L. monocytogenes* [20,21]. Surprisingly, recent approaches to identify novel *in vivo*-induced genes in *L. monocytogenes* (such as microarray and IVET approaches) have failed to identify the key inducible systems for iron-uptake during infection [22,23,24]. In addition, signature tagged mutagenesis approaches have also failed to identify the mechanisms of intracellular iron uptake in this pathogen [25]. We therefore employed a systematic functional genetic analysis of selected Fur-regulated genes and identified a locus (*lmo0641*, now designated *frvA*) that is absolutely required for the systemic phase of *L. monocytogenes* infection.

Ledala and coworkers have recently utilised microarray analysis to identify members of the Fur regulon in *L. monocytogenes* [17]. We concurrently used the classical 19 bp Fur-binding motif defined in *B. subtilis* [26] (Figure 1A) to mine the *L. monocytogenes* EGDe genome for similar motif sequences. We used two primary criteria to limit the number of sequences identified. Firstly, the identified sequence should be within 350bp of an annotated start codon and secondly, a match at 16 or more of the 19 positions was required. Anything less than 16/19 was not considered unless the annotated ORF was deemed to have a likely role in iron acquisition based on bioinformatic analysis. This approach identified a subset of the Fur-regulated loci determined through microarray analysis [17]. However, we also identified Fur-regulated loci at *lmo2431* (previously identified as a potential Fur-regulated locus by Jin *et al.* [2] and *lmo0641* (the subject of this study) which were not detected using the cut-off criteria employed by Ledala *et al.* [17]. Another locus (*lmo0484*) was identified here which is adjacent to a gene (*lmo0485*) identified using microarrays and therefore may form part of an operon.
The loci identified as containing Fur-binding motifs are represented in Figure 1. In each case, where the Fur box was upstream of a putative operon, RT-PCR confirmed co-transcription of all the genes in the operon (data not shown). The Fur boxes were aligned and a graphical display of the results was generated using ‘sequence logo’ which generates a consensus for Listeria that is identical to that in Bacillus (Figure 1B) [27]. Fur regulation was confirmed through RT-PCR analysis of representative genes in both wild-type L. monocytogenes and a Δfur mutant. The results validated the microarray data described previously [17] and also confirmed that lmo2431 and lmo0641 are regulated by Fur.

Virulence analysis of plasmid insertion mutants.

We created mutants using the pORI19 integration strategy as this method is relatively rapid, results in stable mutations and lends itself to analysis of a large number of loci in a reasonable timeframe [20,28]. Two of the identified loci (lmo1007 and lmo0484) consisted of a single small (<500nt) gene and were considered too small for plasmid disruption and were not analysed here. Mutation of fri has been described elsewhere [29,30,31]. Where the Fur box was upstream of an operon we chose the first open-reading frame for plasmid disruption as this would increase the likelihood of causing pleiotropic effects on co-transcribed downstream genes. Plasmid disruptions at the correct locations were confirmed by PCR, using a primer based on the EDGe chromosome and one based on the plasmid. The absence of the repA gene in mutant strains selects against excision and extrachromosomal maintenance of the integration plasmid, ensuring stable integrants for subsequent analysis (see experimental procedures for details). mRNA was extracted from each of the mutants and RT-PCR analysis confirmed that plasmid disruption of the target gene was associated with the complete elimination of expression from each locus with the exception of the lmo2431 mutant (a locus previously analysed by Jin et al. [2]) in which gene expression was greatly reduced (data not shown).
In this initial screen, three of the mutants in Fur-regulated loci exhibited a significant reduction in virulence potential relative to the wild type (P<0.05) (Figure 1E). The most significantly affected mutant in this screen was pORI19::frvA.

**The Fur-regulated virulence (frvA) locus is required for effective infection.** To confirm an essential role for frvA in the virulence of *L. monocytogenes* two precise in-frame deletion mutants were created (see experimental procedures). An initial mutant was created through the deletion of the central region of the *lmo0641* gene, from residues 85-416 inclusive (mutant designated ΔfrvA[85-416]). As toxicity has previously been associated with the generation of truncated membrane proteins through partial deletion [31] we also created a precise deletion mutant in which the entire open reading frame was deleted. This mutant was designated ΔfrvA. Both mutants were complemented using the pPL2 plasmid to re-introduce a single copy of frvA (designated ΔfrvA::pPL2frvA and ΔfrvA[85-416]::pPL2frvA). Although growth of ΔfrvA was unaffected in nutrient-rich media (BHI), this mutant was recovered at significantly lower levels (three-log reduction) from the spleens of infected mice on day three post-infection when compared to the wild-type. Numbers recovered from the liver at three days post inoculation indicated a lesser, but still significant reduction in levels as compared to the wild-type. The reintroduction of frvA fully restored the virulence potential (Figure 2A). These data definitively establish a critical role for frvA in *L. monocytogenes* virulence potential and pathogenesis. Notably, ΔfrvA[85-416] also demonstrated a dramatic reduction in virulence potential in the murine model (Supplemental Fig S1).

Larvae of the Wax Moth (*Galleria mellonella*) have recently been utilized as an alternative pathogenicity model for *L. monocytogenes* [32,33]. Here we also analysed the virulence potential of ΔfrvA using this model system (Figure 2B). While no deaths were observed over time for the insects that received PBS, a significant difference was seen between insects receiving the wild-type and the ΔfrvA mutant. The LT-50 for insects infected with the wild-type strain was 26 hours while over 50% of the ΔfrvA-infected insects were still alive after 43 hours. The significance of iron acquisition to the virulence of bacterial pathogens has previously been investigated in this insect model. Work by
Daou et al. [34] demonstrates a role for IlsA, a surface protein in Bacillus cereus that binds haemoglobin and ferritin, to pathogenesis in the G. mellonella host. In order to determine the possible influence of the downstream gene lmo0642 in murine virulence of L. monocytogenes we created an in-frame mutation in this locus. Interestingly, this gene is apparently not required for pathogenesis. Deletion of lmo0642 failed to affect the ability of L. monocytogenes to reach either the liver or spleen in mice infected by the intraperitoneal (i.p.) route (Figure 2C) or to grow intracellularly (data not shown).

ΔfrvA was compared to the wild-type and complemented strains for their ability to replicate within J774 macrophage cells (Supplemental Figure S2). After one hour ΔfrvA displayed no significant difference in invasion of J774 cells when compared to the wild-type or complement strains. Subsequent readings taken after three, five and seven hours represent intracellular survival and multiplication of these strains within the cell line. Similarly, no significant difference was observed in the ability of ΔfrvA to survive inside J774 macrophages over time, as all three strains displayed growth of approximately one log after 7 hours.

**Bioinformatic analysis of frvA.** FrvA is a putative transmembrane protein consisting of six predicted transmembrane regions (SOSUI) and is annotated as being similar to a heavy metal-transporting ATPase (http://genolist.pasteur.fr/ListiList/). The closest non-listerial homologues reside in Bacillus spp. A predicted heavy metal-transporting ATPase in B. megaterium was found to share 56% identity and 72% similarity (over a query coverage of 621/626 amino acids) with FrvA. A predicted cadmium-transporting ATPase in B. halodurans C-125 also shared close homology with 55% identity and 72% positives (over 618/626 amino acids) (NCBI Blast). Three conserved domains were identified in FrvA using the Conserved Domain Search from NCBI including a P-type ATPase superfamily, a haloacid dehalogenase-like (HAD) hydrolase, and a cation transport ATPase. In addition, FrvA was found to contain several classic P-type ATPase motifs such as the phosphorylation motif D321KTGTLT and the hinge region motif G518DGIND. Similar to other type I heavy metal-transporting ATPases such as YkvW in Bacillus subtilis and CtpA, a P-type ATPase involved in copper homeostasis in L.
monocytogenes, FrvA also possesses both an M4 motif S^{277}PC and an HP motif, S^{358}LHPLA, respectively [35].

Lmo0642, the product of the downstream gene, is also predicted to be localized to the bacterial membrane (PSORT) and also has 6 transmembrane regions (SOSUI). No conserved domains were identified (NCBI) and its closest homolog is a hypothetical protein (EF0716) from Enterococcus faecalis V583 (NCBI Blast).

Regulation of frvA by Fur. qRT-PCR analysis of the wild-type L. monocytogenes EGDe strain and a Δfur mutant confirmed that lmo0641 is under the negative regulation of Fur. Using the 2-ΔΔCt method to calculate the relative changes in gene expression, lmo0641 was shown to be up-regulated 93-fold in Δfur compared to the wild-type. Transcription of frvA was also found to be under the control of PerR, a Fur homolog which functions as an Fe(II)-dependent peroxide stress sensor and which regulates putative metal transport and storage functions [36]. In addition to the classical Fur box a putative PerR binding region was identified upstream of the annotated start codon of frvA. De-repression of frvA was seen in the absence of either regulator. However, no further impact was observed in a ΔfurΔperR double mutant (data not shown). The significance of this dual regulation by Fur and PerR is unclear, but highlights some degree of interplay between these two regulators. It is interesting to note that frvA (lmo0641) was also previously determined to be regulated by PrfA, the master regulator of virulence gene expression in L. monocytogenes [37]. The locus is not preceded by a detectable PrfA binding motif but the authors noted the presence of a binding site recognized by Sigma B, the general stress response regulator. Taken together, the evidence suggests that the locus is a member of multiple regulatory networks, perhaps reflecting the importance of FrvA in L. monocytogenes niche adaptation.

ΔfrvA displays increased haemin uptake and elevated sensitivity to haem toxicity. In an attempt to understand the virulence defect displayed by ΔfrvA we carried out extensive physiological analysis of the mutant strain. A [^{59}Fe]-citrate uptake assay indicated that
the ability of ΔfrvA to acquire ferric citrate was not impaired when compared to the wild-type *L. monocytogenes* strain (Fig. 3A). Both strains transport ferric citrate with similar affinity (K_m) and velocity (V_max). Although Adams *et al.* [38] have reported that a citrate inducible iron uptake system exists in *L. monocytogenes* we demonstrate here that the FrvA system is not involved in the direct uptake of ferric citrate. The existence of an iron reductase has previously been suggested in *L. monocytogenes* based upon physiological data [39,40] although this remains the subject of some debate [2]. We performed iron reductase assays but could find no significant difference between wild-type and mutant cells in ability to reduce iron in these assays, suggesting that this locus does not encode an iron reductase (see Supplementary Table T1).

During infection free iron is not available to bacterial cells whereas haem (Hb) and haemin (Hn) represent a potentially abundant source of iron [41]. However haem can be relatively toxic to cells at elevated concentrations [42,43]. We investigated the rates of haemin uptake in ΔfrvA and observed significant differences between the wild-type and mutant strains in the acquisition of [59Fe]-Hn (Fig. 3B). The rate of haemin transport by ΔfrvA (V_max = 30.6 pMol per 1x10^9 cells per minute) was nearly twice that of the wild-type strain (V_max = 18.8 pMol per 1x10^9 cells per min). Subsequent analysis of the mutant in iron-limiting MOPS-L media supplemented with haemoglobin and haemin revealed that *L. monocytogenes* ΔfrvA displayed growth behavior distinct from that of the wild-type and complement strains (Fig. 4). Growth of the wild-type and complement was restored upon addition of increasing concentrations of Hb and Hn (0.2 and 2.0µM) to iron-limiting media (Fig. 4A, 4C, 4F). In contrast, growth of ΔfrvA required addition of 0.2µM Hn and Hb, whereas a higher concentration of 2.0µM was shown to reduce growth suggestive of toxicity (Fig. 4B and 4E). Nutrition tests were performed to assess the capability of the strains to utilize iron from several different sources. ΔfrvA displayed no impairment in ability to utilize ferric siderophores, Hb or Hn when compared to the wild-type and complement strains (Fig. 5).

As FrvA displays homology to bacterial heavy-metal transporting ATPases and with the knowledge that cation-transporting ATPases function in maintaining cation homeostasis [35], we investigated the sensitivity of ΔfrvA to toxic levels of heavy metal
sulfates such as copper, cobalt, cadmium, and zinc as well as iron. Exposure to a disk that
contained 1M FeSO$_4$ resulted in a larger zone of clearance in $\Delta$frvA when compared to
the wild-type, indicative of elevated toxicity. However sensitivity to other heavy metals
such as CdSO$_4$, CoSO$_4$, CuSO$_4$ and ZnSO$_4$ was comparable in both the wild-type and
mutant (Supplemental Figure S3). The data suggest that deletion of frvA does not affect
the sensitivity of cells to heavy metals such as cadmium, cobalt, copper and zinc but
confirms the contribution of this locus to iron homeostasis.

Global disruption of iron homeostasis in the $\Delta$frvA mutant. As physiological analysis
of $\Delta$frvA revealed iron-related phenotypes, we investigated the possibility that deletion of
this locus could lead to altered expression of other genes in the L. monocytogenes
genome involved in iron homeostasis. qRT-PCR was used to evaluate the differential
expression of three iron-related genes in the wild-type and mutant strains (Figure 6). We
chose two Fur-regulated genes; lmo2186 which encodes a homologue of SaulsdC and
bears homology to a haemin binding protein IsdC in S. aureus [44], and lmo1959,
designated as fhuD encoding the L. monocytogenes ferrichrome binding protein [2,44].
In addition, lmo2431 (hupD) was also analyzed as this gene is part of the hupDGC
operon encoding an ABC transporter involved in uptake of haemin and haemoglobin
[2,44]. qRT-PCR analysis revealed a strong induction of both lmo2431 and lmo1959 in
$\Delta$frvA compared to the wild-type strain. lmo2431 was shown to be up-regulated 210-fold
and lmo1959 up-regulated 164-fold in the mutant strain. lmo2186 also displayed a an
induction in $\Delta$frvA, with an almost 5-fold difference observed between the wild-type and
mutant. As Fur is generally considered a repressor of transcription [16], the induction of
two Fur-regulated genes in $\Delta$frvA is supported by our finding that the fur gene was shown
to be down-regulated almost 6-fold in $\Delta$frvA.

Conclusions

Using a functional genetics approach we identified a novel Fur-regulated locus
(frvA) in L. monocytogenes that is essential for virulence and for resistance to haem and
haemin-mediated toxicity. It is known that *L. monocytogenes* has the capacity to utilise iron-loaded haemoglobin and haemin as sources of iron [31]. Furthermore, elimination of haemoglobin and haemin uptake through mutation of the HupC transport system significantly impairs virulence potential, indicating that iron acquisition from haem is essential for pathogenesis [2]. However, haem and haemin are known to be toxic for bacteria and many bacteria express specific mechanisms for detoxification of haem [43].

FrvA possesses P-type ATPase and hydrolase conserved domains and is homologous to other heavy-metal transporting ATPases in *Staphylococcus* and *Bacillus*. Previous work by Francis and Thomas [35] identified another P-type ATPase, encoded by *ctpA*, which is involved in copper homeostasis in *L. monocytogenes*. Significantly, mutagenesis of the *ctpA* locus resulted in a strain that was unaffected in intracellular growth in the J774 macrophage cell line, but was impaired in ability to cause infection in the murine model [45]. Although P-type ATPases are known to mediate the transport of various heavy metals in bacteria, iron transport is most often associated with the structurally unique ATPases of the ABC transporter family [46]. However, Mta72, a P-type ATPase in *M. tuberculosis*, has been shown to transport iron transferred from the siderophore carboxymycobactin and is another rare example of a P-type ATPase involved in iron homeostasis [6].

It is interesting to note that the HrtA system in *S. aureus* also functions as a haem exporter and deletion of *hrtA* in that organism causes dysregulation of Fur expression resulting in pleiotrophic effects [47]. Whilst HrtA is an ABC transporter rather than a P-type ATPase we note homologies between FrvA and HrtA (21% identity over 221 amino acids). Certainly the results presented here suggest functional similarities between FrvA and HrtA though further experimental work will be necessary to directly compare both systems.

We did not demonstrate a role for FrvA in transport of ferric citrate or in iron reduction by *L. monocytogenes* and the mutant was not impaired in intracellular growth in *vitro*. Rather the predominant phenotype of ∆*frvA* is an increased uptake of haemin and significantly increased sensitivity to both haemin and haemoglobin toxicity and reduced virulence during systemic infection. However we acknowledge that further work is
necessary to determine the precise biochemical mechanisms underpinning FrvA activity. The profound dysregulation of iron homeostasis in ∆frvA results in the de-repression of other Fur-regulated loci which complicates interpretation of the analysis of the mutant and which may necessitate the future use of isolated liposomal protein models to delineate its precise function.

Materials and Methods

Ethics statement

All animal procedures were approved by the University Research Ethics Board (UREB) in University College Cork (approval ID 2008/32) and were carried out in a specialized facility. Work was carried out under license from the Irish Department of Health.

Bacterial strains, plasmids and culture conditions. Listeria monocytogenes strains were grown in Brain Heart Infusion (BHI) (Oxoid) broth at 37°C and Escherichia coli strains were grown in Luria-Bertani (LB) broth at 37°C. Strains and plasmids used in this study are listed in Table 1. For solid media, agar (1.5%) was added. Antibiotics, obtained from Sigma Chemical Company, were added in the following concentrations; 50µg/ml ampicillin for pKSV7 in E. coli and 10 µg/ml chloramphenicol for pKSV7 in L. monocytogenes. For pPL2 in E. coli and L. monocytogenes, concentrations of 15 and 7.5 µg/ml chloramphenicol were used, respectively. Where indicated L. monocytogenes strains were sub-cultured at 1% into iron-deficient MOPS minimal salts medium [Neidhardt, 1974 #1524], with appropriate supplements (MOPS-L; [Xiao, #7571] to stationary phase (OD600 of approximately 1.2), and then subcultured again (1%) into MOPSL and grown to mid-log phase. Ferrichrome (Fc) and ferrichrome A (FcA) were purified from cultures of Ustilago sphaerogena [Emery, 1971 #2185]. Ferrioxamine B (FxB) was a gift from J. B. Neilands. We purchased purified hemin (Hn) and bovine hemoglobin (Hb) from Sigma-Aldrich (St. Louis, Mo).
DNA manipulations. Gel extraction was performed using the Qiagen Gel Extraction Kit (Qiagen). Plasmid DNA isolation was carried out utilizing the Qiagen QIAprep Spin Miniprep Kit (Qiagen). PCR reagents and T4 DNA ligase, supplied by Roche Diagnostics GmbH (Mannheim, Germany), and restriction enzymes (New England Biolabs) were all used according to the manufacturer’s instructions. Oligonucleotide primers were synthesized by MWG and are listed in Table 2. PCR reactions were completed using a PTC-200 (MJ Research) PCR system. Colony PCR was performed following lysis of cells with IGPAL CA-630 (Sigma). Genomic DNA was isolated from L. monocytogenes using a chromosomal kit (Sigma) according to the manufacturer’s instructions.

Creation of plasmid insertion mutants. A central portion of the gene of interest was amplified by PCR and cloned into the multiple cloning site of pORI19 (RepA−) [20,28]. Following plasmid isolation, electrotransformation of L. monocytogenes EGDe containing pVE6007 (RepA+/Temperature sensitive) was performed according to the protocols outlined by Park and Stewart, (1990). Loss of pVE6007 was achieved by transferring 10μl of a 30°C overnight culture to BHI broth prewarmed to 42°C with subsequent growth for 16hrs at 42°C and isolation on prewarmed BHI-Em (5µg/ml) agar plates at 42°C. Loss of pVE6007 (Cm+) was confirmed by replica plating onto BHI-Em (5µg/ml) and BHI-Cm (10µg/ml) plates with overnight incubation at 30°C. Integration results in the formation of a stable Em+r mutant and was confirmed by PCR using a primer outside the region of integration and a primer specific to the plasmid.

Construction of deletion mutants. As described by Horton et al. [48] the Splicing by Overlap Extension (SOE) procedure was utilized to create a complete gene deletion mutant. This is an in-frame, non-polar deletion of a gene in the L. monocytogenes EGDe chromosome. Two pairs of primers were designed, SOEA/SOEB and SOEC/SOED, to amplify two fragments of approximately equal size on either side of the gene to be deleted using the proofreading enzyme Vent polymerase (New England Biolabs). These AB and CD products were then gel extracted to ensure purity, mixed in a 1:1 ratio, and
were spliced together using SOEA/SOED primers. This AD product was digested and cloned into pKSV7, a temperature sensitive plasmid. The resulting construct was electroporated into competent *E. coli* DH5α cells and transformants were selected on Luria-Berani plates with ampicillin. The plasmid was isolated using the Qiagen QIAprep Spin Miniprep kit. The presence of the correct insert was verified by sequencing (Lark Technologies Inc., Essex, UK) using the pKSV7 MCS primers M13F and M13Rmut. The isolated plasmid was electroporated into competent *L. monocytogenes* EGDe cells. Transformant selection took place on Brain-Heart Infusion agar containing chloramphenicol. Clones in which chromosomal integration of the plasmid had occurred are selected by serial passaging at 42°C and are streaked onto pre-warmed BHI-Cm agar plates. Continuous passaging at 30°C in BHI broth followed by replica plating onto BHI and BHI-Cm plates ensures plasmid excision. Chloramphenicol sensitive colonies were tested for gene deletion using primers SOEX, located upstream, and SOEY, located downstream of the gene of interest.

**Complementation of deletion mutants.** A site-specific phage integration vector, pPL2, was used for the complementation of SOEing deletion mutants. This vector integrates within the tRNA^Arg^ gene on the chromosome. Vent polymerase (New England Biolabs), a proof reading enzyme, was used to amplify the entire deleted gene and flanking regions, including the upstream gene promoter. Primers CompA and CompB included restriction sites corresponding to those on the MCS site of pPL2. The PCR product was gel extracted to ensure purity, and was digested and cloned into pPL2. The resulting construct was electroporated into competent *E. coli DH5α* cells and transformants were selected on Luria-Berani plates with chloramphenicol. The plasmid was isolated using the Qiagen QIAprep Spin Miniprep kit. The presence of the correct insert was verified by sequencing (Lark Technologies Inc., Essex, UK) using the pPL2 MCS primers T3F and T7R. The isolated plasmid was electroporated into competent SOEing mutant cells. Transformant selection took place on Brain-Heart Infusion agar containing chloramphenicol. The presence of the gene was authenticated using a forward running check primer that anneals to the middle of the gene and SOE D, located on the cloned
insert. Integration of pPL2 to the correct site was confirmed using primers PL102, located upstream of the integration site, and the SOE D primer.


RNA extraction. Total RNA was extracted using both the Macaloid Clay method, outlined by Raya et al. [49], and the Qiagen RNeasy Mini Kit. Cultures were grown overnight shaking at 37°C. A 1% inoculum was added to 30mLs BHI broth and cultures were grown at 37°C until an OD<sub>600nm</sub> of 0.3 was reached. 30mLs of culture were pelleted by centrifugation at 4,000g for 7 minutes. The supernatant was removed, the pellet was washed with 1mL cold TE buffer (10mM Tris, 1mM EDTA: pH 8.0), and centrifuged again for 13,000g for 1 minute. Again the supernatant was removed, and the pellet was resuspended in 20µL lysozyme (50mg/mL), 400µL cold TE buffer, and left at room temperature for 3 minutes. Subsequently, the cell suspension was added to a 1.5mL screw-cap plastic tube containing 50µL 10% sodium dodecyl sulphate, 500µL phenol-chloroform (5:1), 175µL Macaloid Clay and 0.5g 425-600µm glass beads (Sigma). Cell disruption was achieved using a bead beater (Mini-beadbeater 8TM cell disrupter, Biospec products.) Cells were beaten for 1 minute, placed on ice for 1 minute, beaten
again for 1 minute, and then centrifuged for at 13,000g for 3 minutes. The organic layer was removed and precipitated with 1:10 volume sodium acetate, and 2.5 volume 96% ethanol at -80°C for 20 minutes. Following this step, samples were put through the Qiagen RNeasy Mini Kit and then eluted in 50µL TE buffer. RNA samples were treated with RNase-free DNase I set (Qiagen) and DNA-free (Ambion) was used to remove any DNA. The concentration of RNA was quantified utilizing a NanoDrop (ND-1000 spectrophotometer). A PCR, carried out with 16S rRNA primers; L142 and U141, was used to ensure the absence of DNA in the samples. The reverse transcriptase PCR was run using 4µL random primer p(dN)$_6$, 2µL RNA, and 2µL DEPC water (Sigma) at 65°C for 10 min, and put directly on ice. To these samples, 32µL of a mastermix was added containing: 1µL Expand Reverse Transcriptase (Roche), 8µL 5x Buffer (Roche), 4µL 100mM dTT (Roche), 1µL dNTP mix (dATP, dCTP, dGTP, dTTP; 10mM) and 18µL DEPC water. This reaction was carried out at 30°C for 10 min, 42°C for 3 hours, and held at 4°C. cDNA was confirmed through PCR using L142 and U141 primers and the wild-type L. monocytogenes extracted DNA as a positive control.

Quantitative real-time PCR. The Universal Probe Library Assay Design Center (https://www.roche-applied-science.com/sis/rtpcr/upl/adc.jsp) was used to design PCR primers which correspond to a specific probe in the library. Primer sequences and corresponding probes are listed in Table 2. The 16S rRNA gene was used as a housekeeping gene to compensate for any variability in the initial amount of starting total RNA. Amplification reactions consisted of 2.5µL of cDNA, 6.4µL of 2x FastStart TaqMan Probe Master (Roche), primers (900nM) and probe mix (250nM). RNase-free water was added to bring the total volume of the reaction to 10µL. Reactions were performed in duplicate on 384-well plates using the LightCycler 480 System (Roche). Negative control reactions, without cDNA, were also included on the plate. Thermal cycling conditions were carried out according to manufacturer’s instructions (Roche) and the $2^{-ΔΔCt}$ method [50] was used to calculate the relative changes in gene expression from the qRT-PCR experiments.
Growth curves. Growth of *Listeria monocytogenes* in MOPS-L media. EGD-e and its mutant derivatives were grown in BHI overnight, and then subcultured at 1% into BHI broth or MOPS-L media. In the latter case, the bacteria were grown to stationary phase, and for growth rate determinations they were subcultured again at 1% into MOPS-L containing Hn or Hb at varying concentrations. The cultures were shaken at 37 °C and OD$_{600}$ nm was monitored at indicated time points up to 26 hours.

Metal disk assay. Cultures were grown overnight shaking at 37°C. A 2% inoculum was added to 10mL of fresh BHI and cultures were grown to logarithmic phase (0.3OD) at 37°C. 400µL of log phase cell cultures were added to 4mL of cooled, molten soft agar (0.75%) and poured on top of a petri dish containing 20mL BHI agar. After solidifying, a sterilized 13mm disk (Whatman) was placed on top of the overlay. Metals used were made up in 1M stocks in which 35µL of each metal were dispensed onto the center of the disk. The plate was then incubated overnight at 37°C, and the zone of clearance surrounding the disk was measured.

$^{59}$Fe binding and uptake experiments. For binding and transport determinations, we prepared $^{59}$Fe complexes of citrate (specific activity 150 to 1,000 cpm/pMol) and haemin [44]. For $^{59}$Fe-citrate, we provided the organic ligand in 50-fold molar excess. We conducted adsorption and transport experiments [2,44] over a range of concentrations, by adding appropriate amounts of $^{59}$Fe complexes to two aliquots of 2 x 10$^7$ cells of EGD-e or its mutants, and incubating the aliquots for 15 s and 75 s, respectively, before collecting and washing the cells on 0.2 micron filters. The 15 s aliquot measured the amount initially bound to the cells, which when subtracted from the second time-point, gave the amount transported during a 1 min period. At each concentration, data were collected in triplicate and averaged. The $K_d$ and capacity of $^{59}$Fe-siderophore binding were determined by using the “Bound-versus-Total” equation of Grafit 5.09 (Erithacus, Ltd., Middlesex, UK), and $K_m$ and $V_{max}$ of transport were calculated using the “Enzyme Kinetics” equation.
Macrophage assay. This intracellular survival assay was carried out using J774 mouse macrophage cells (originally obtained from the American Type Culture Collection, Manassas, VA). 24-well tissue culture plates were seeded with $1 \times 10^5$ live cells per well in DMEM (Gibco) containing 10% fetal calf serum and incubated in 5% CO$_2$ at 37°C for 40 hours. For infection, bacteria were prepared by centrifuging 1 mL of an overnight culture which was then washed once in PBS, and resuspended in 1mL DMEM. Bacteria were diluted in DMEM and $1 \times 10^7$ CFU was added to each well containing washed macrophage cells. To increase contact between macrophages and bacteria, the 24-well plates were centrifuged at 1500rpm for 10min and incubated for 1 hour in 5% CO$_2$ at 37°C. To kill extracellular bacteria, 1mL of 100μg/mL gentamycin (Sigma) was added to each well and incubated for an additional 30 min. Bacteria surviving intracellularly were enumerated at time points taken after addition of gentamycin. Each well was washed twice in PBS, cells were lysed with 250μL ice cold water containing 0.02% Triton X (Sigma), and scraped in a similar manner using a pipette tip. Serial dilutions were carried out on the lysate and plated on BHI agar overnight at 37°C.

Murine virulence assay. Cultures were grown overnight with shaking at 37°C. Cultures were centrifuged, washed in PBS (Sigma), resuspended and diluted to $1 \times 10^6$ CFU/mL in PBS. BALB/c mice were inoculated with $4 \times 10^5$ CFU in 200 μL PBS intraperitoneally (i.p.). The mice were euthanized 3 days post-infection. Spleens and Livers were harvested and then homogenized in PBS. Bacteria were enumerated by plating the serial dilutions of organ homogenates on BHI agar left to incubate overnight at 37°C.

Galleria mellonella virulence assay. Cultures were grown overnight with shaking at 37°C. Cultures were centrifuged, washed, and resuspended in an equal volume of PBS (Sigma). Infection of Galleria mellonella was performed according to the protocol outlined by Joyce et al. [32]. Briefly, insects were obtained from Livefood, UK and were
stored in the dark at room temperature prior to use. 3 groups, containing 10 insects per
group, were injected with $1 \times 10^6$ CFU/10µL of the wild-type *L. monocytogenes* EGD-e
strain (group 1), $1 \times 10^6$ CFU/10µL EGDΔ0641 (group 2), or 10 µL PBS (group 3) to
serve as a negative control. Bacterial suspensions were injected using a sterile Hamilton
syringe and a 30-Gauge disposable needle into the first right pro-leg of the second set of
pro-legs. All ten insects per group were placed together in a Petri-dish lined with
Whatman paper and incubated in the dark at 37°C. Insects were examined over several
days and time of death was recorded.

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Institutes of Health (NIH) grant GM53836 and National Science Foundation (NSF) grant
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Author Contributions

Conceived and designed the experiments: AC, RC, CH, PK, CG. Performed the
experiments: HM, QX, RR, HP, PC, TD, RS, SJ, RC. Analyzed the data: HM, QX, RR,
HP, TD, AC, RS, SJ, RC, CH, PK, CG. Wrote the paper: HM, QX, PK, CG.
# References


Figure Legends

Figure 1. Identification and role in virulence of Fur-regulated gene systems. (A) The classical Fur box is represented as a 19 bp sequence. Recent studies have suggested that a more accurate representation of the Fur box is that of a 7-1-7 motif. The 19bp sequence was used to search the Listeria monocytogenes EGDe genome sequence (Listilist). (B) Identified sequences were aligned and a graphical display of the results was generated using the web based programme sequence logo (17). (C) Genetic organisation of 29 putative Fur regulated genes (black/gray) at 12 chromosomal loci. All genes are drawn approximately to scale using the L. monocyctogenes EGDe genome sequence data. Lmo numbers refer to the National Centre for Biotechnology Information annotation scheme. Fur boxes are represented by black circles. Gray genes indicate those disrupted in EGDe in the course of this study. Lollipops are used to illustrate putative stem loop terminator regions. (D) RT-PCR analysis was used to confirm Fur regulation of all identified genes and to give an indication of the increase in expression levels. Control primers were used to ensure that template cDNAs were of equal concentration. Samples were removed at various cycles of PCR (cycle number in brackets) and visualised on agarose gels. A repeat experiment demonstrated similar results. Results were also verified through real-time PCR analysis. (E) In vivo survival of disruption mutants in Fur-regulated loci in the murine infection model. Mice were injected i.p. with either the wild-type or mutants and the number of bacteria recovered from the spleen was determined three days post-inoculation. Error bars represent the standard deviations from the mean (n=4). * indicates means are significantly different to the wild-type (P<0.05). ND, not detected.

Figure 2. Pathogenesis of strains in murine and Wax Moth models of infection. (A) CFUs of the ΔfrvA and ΔfrvA complemented strain enumerated from livers and spleens three days post infection. Error bars represent standard error of the mean and asterisks represent P < 0.001 by the Student’s t-test when compared to the wild-type and complement strains. (B) Pathogenesis of strains in the Galleria mellonella model of infection. Dotted line and cross indicates LT-50 (time in which 50% of insects had perished). (C) Pathogenesis of the Δlmo0642 mutant in the murine model of infection.
Strains were inoculated into mice by the ip route and numbers were enumerated in the spleens at day three post-infection. Student t-test did not detect a significant difference between the wild-type and mutant strain.

**Figure 3.** $^{59}$Fe binding and uptake assays. Uptake affinity ($K_m$ in nM) and velocity ($V_{max}$ in pMol per $10^9$ cells per minute) by which the wild-type (open circles) and Δ0641 (ΔfrvA) (closed circles) strains transport $[^{59}\text{Fe}]$-citrate (A) and $[^{59}\text{Fe}]$-Hn (B) were assessed. Overall $K_m$ and $V_{max}$ of $[^{59}\text{Fe}]$ transport are listed in the tables on right-hand side. Data was plotted using the Enzyme Kinetics algorithm of Grafit 7 (Erithacus Ltd, West Sussex, UK) and represent the mean of independent experiments done in triplicate.

**Figure 4.** Bacterial growth. The rates and extent of bacterial growth (A: EGD-e; C: Δlmo0641; E:Δlmo0641/pPL2lmo0641) were determined in iron-restricted MOPS-L media supplemented with Hb (panels A - C; open, gray and black symbols represent addition of 0.0, 0.02 and 2 μM Hb, respectively) or Hn (D, E; open, gray and black symbols represent addition of 0.0, 0.2 and 2 μM Hn, respectively), and in BHI broth (F). The bacteria were cultured in BHI broth overnight. In A-E they were then subcultured in MOPS-L to stationary phase, and at t = 0 subcultured again at 1% into MOPS-L containing different concentrations of Hb or Hn. In F, at t = 0 they were subcultured into BHI broth. The flasks were shaken at 37 °C and absorbance at 600 nm (initially close to zero for all cultures) was monitored for 12 - 26 h (note different scales). Because of the slow growth of *L. monocytogenes* in iron-restricted minimal media, this graphic representation focuses on the comparison of the mutant strains at later times in the growth cycle.

**Figure 5.** Nutrition tests. Tests demonstrate the halo of growth surrounding a paper disc embedded with 10 μl aliquots of the test iron compound. Concentrations of compounds are indicated as μM. Fc (ferrichrome) and FcA (ferrichrome A), FxB (ferrioxamine B), Hb (haem/haemoglobin) and Hn (haemin) were tested on BHI agar containing 0.1 mM
BP. The experiment was repeated several times with similar results. No differences were seen between mutant strains and the wild-type in these iron nutrition assays.

Figure 6. Quantitative real-time PCR. Induction of lmo0641 (frvA) transcription in Δfur compared to the wild-type (black bar) and induction of gene transcriptions in ΔfrvA compared to the wild-type (gray bars) in BHI. Up-regulated genes are represented by bars above the x-axis and the down-regulated gene (fur) is represented by the bar below the axis. Asterisks represent Fur-regulated genes. Error bars represent the mean ± SD of the relative change in gene expression of independent duplicate samples.
### Table 1. Bacterial strains and plasmids used in this study

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Table 2. Oligonucleotide primers used in this study

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Restriction sites are in boldface and complementary overhangs are underlined
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756
757