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Published in:
eLife

DOI:
10.7554/eLife.57322

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Document Version
Final author's version (accepted by publisher, after peer review)

Publication date:
2020

Link to publication in University of Groningen/UMCG research database

Citation for published version (APA):

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An ECF-type transporter scavenges heme to overcome iron-limitation in *Staphylococcus lugdunensis*

Jochim A\(^1\), Adolf LA\(^1\), Belikova D\(^1\), Schilling NA\(^2\), Setyawati I\(^3\), Chin D\(^4\), Meyers S\(^5\), Verhamme P\(^5\), Heinrichs DE\(^4\), Slotboom DJ\(^3\) and Heilbronner S\(^1,6,7^{*}\)

1 - Interfaculty Institute of Microbiology and Infection Medicine, Department of Infection Biology, University of Tübingen, Tübingen, Germany

2 - Institute of Organic Chemistry, University of Tübingen, Tübingen, Germany

3 - Groningen Biomolecular Sciences and Biotechnology Institute, University of Groningen, Groningen, The Netherlands

4 - Department of Microbiology and Immunology, University of Western Ontario, London, Ontario, Canada.

5 - Center for Molecular and Vascular Biology, KU Leuven, Belgium.

6 - German Centre for Infection Research (DZIF), Partner Site Tübingen, Tübingen, Germany

7 - (DFG) Cluster of Excellence EXC 2124 Controlling Microbes to Fight Infections

* Corresponding author

Abstract:

Energy-coupling factor type (ECF-transporters) represent trace nutrient acquisition systems. Substrate binding components of ECF-transporters are membrane proteins with extraordinary affinity, allowing them to scavenge trace amounts of ligand. A number of molecules have been described as substrates of ECF-transporters, but an involvement in iron-acquisition is unknown. Host-induced iron limitation during infection represents an effective mechanism to limit bacterial proliferation. We identified the iron-regulated ECF-transporter Lha in the opportunistic bacterial pathogen *Staphylococcus lugdunensis* and show that the transporter is specific for heme. The recombinant substrate-specific subunit LhaS accepted heme from diverse host-derived hemoproteins. Using isogenic mutants and recombinant expression of Lha, we demonstrate that its function is independent
of the canonical heme acquisition system Isd and allows proliferation on human cells as sources of
nutrient iron. Our findings reveal a unique strategy of nutritional heme acquisition and provide the first
element of an ECF-transporter involved in overcoming host-induced nutritional limitation.

**Key words**
Iron, heme, ECF-transporter, Staphylococcus lugdunensis, nutritional immunity

**Background:**
Trace nutrients such as metal ions and vitamins are needed as prosthetic groups or cofactors in
anabolic and catabolic processes and are therefore crucial for maintaining an active metabolism. Metal
ions such as iron, manganese, copper zinc, nickel and cobalt must be acquired from the environment
by all living organisms. In contrast many prokaryotes are prototrophic for vitamins like riboflavin,
biotin and vitamin B$_{12}$. However, these biosynthetic pathways are energetically costly [1], and
prokaryotes have developed several strategies to acquire these nutrients from the environment. ABC
transporters of the Energy-coupling factor type (ECF-transporters) represent highly effective trace
nutrient acquisition systems [2, 3]. In contrast to the substrate-binding lipoproteins/periplasmic
proteins of conventional ABC transporters, the specificity subunits of ECF transporters (ECF-S) are
highly hydrophobic membrane proteins (6-7 membrane spanning helices) [2]. ECF-S subunits display
a remarkably high affinity for their cognate substrates in the picomolar to the low nanomolar range,
which allows scavenging of smallest traces of their substrates from the environment [4]. Whether ECF
type transporters can be used to acquire iron or iron-containing compounds is unknown.

The dependency of bacteria on trace nutrients is exploited by the immune system to limit
bacterial proliferation by actively depleting nutrients from body fluids and tissues. This strategy is
referred to as “nutritional immunity” [5, 6]. In this regard, depletion of nutritional iron (Fe$^{2+}$/Fe$^{3+}$) is
crucial as iron is engaged in several metabolic processes such as DNA replication, glycolysis, and
respiration [7, 8]. Extracellular iron ions are bound by high-affinity iron-chelating proteins such as
lactoferrin and transferrin found in lymph and mucosal secretions and in serum, respectively.
However, heme is a rich iron source in the human body and invasive pathogens can access this heme
pool by secreting hemolytic factors to release hemoglobin or other hemoproteins from erythrocytes or other host cells. Bacterial receptors then extract heme from the hemoproteins. This is followed by import and degradation of heme to release the nutritional iron. To date, several heme acquisition systems of different Gram-positive and Gram-negative pathogens have been characterized at the molecular level (see [9] for an excellent review).

Staphylococci are a major cause of healthcare-associated infections that can lead to morbidity and mortality. The coagulase-positive *Staphylococcus aureus* represents the best-studied and most invasive species. Coagulase-negative staphylococci (CoNS) are regarded as less pathogenic than *S. aureus* and infections caused by CoNS are normally subacute and less severe. In this regard, the CoNS *Staphylococcus lugdunensis* represents an exception. *S. lugdunensis* infections frequently show a fulminant and aggressive course of disease that resembles that of *S. aureus*. Strikingly, *S. lugdunensis* is associated with a series of cases of infectious endocarditis [10]. The reasons for the apparently high virulence potential of *S. lugdunensis* remains largely elusive and few virulence factors have been identified so far. In this respect, it is interesting to observe that *S. lugdunensis*, unlike other staphylococci but similar to *S. aureus*, encodes an iron-dependent surface determinant locus (Isd) system [11, 12]. Isd facilitates the acquisition of heme from hemoglobin and can be regarded as a hallmark of adaptation towards an invasive lifestyle. However, to ensure continuous iron acquisition within the host, many pathogens encode multiple systems to broaden the range of iron-containing molecules that can be acquired [13].

Here we report the identification of an iron-regulated ECF-type ABC transporter (named LhaSTA) in *S. lugdunensis*. We found LhaSTA to be specific for heme, thus representing a novel strategy to overcome nutritional iron limitation. Recombinant LhaS was able to take up heme from several host hemoproteins such as hemoglobin, myoglobin or hemopexin. Consistent with these data, LhaSTA expression allowed proliferation of *S. lugdunensis* in the presence of these iron sources as well as human erythrocytes or cardiac myelocytes as a sole source of nutrient iron. Our data indicate that LhaSTA function is independent of the presence of surface-displayed hemoprotein receptors suggesting Isd-independent acquisition of heme from host hemoproteins. Our work identifies LhaSTA...
as the first ECF transporter that facilitates iron acquisition, thus participating to overcome host immune defenses.

Results:

LhaSTA encodes an iron regulated ECF transporter

The *isd* locus of *S. lugdunensis* shows several characteristics that distinguish it from the locus of *S. aureus*. Amongst these is the presence of three genes that encode a putative ABC transporter and are located between *isdJ* and *isdB* (Fig. 1A) [11]. Analysis of the open reading frame using Pfam [14] revealed that the three adjacent genes encode components of a putative ECF-transporter, namely a specificity subunit (*lhaS* - SLUG_00900), a transmembrane subunit (*lhaT* - SLUG_00910) and an ATPase (*lhaA* - SLUG_00920), and they might be part of a polycistronic transcript. The location within the *isd* operon suggested a role of the transporter in iron acquisition. Bacteria sense iron limitation using the ferric uptake repressor (Fur) which forms DNA-binding dimers in the presence of iron ions [15]. Under iron limitation, Fe dissociates from Fur and the repressor loses affinity for its consensus sequence (*fur box*) allowing transcription. Interestingly, a *fur-box* was located upstream of *lhaS* (Fig. 1A). qPCR analysis in *S. lugdunensis* N920143 revealed that the expression of *lhaS* and *lhaA* increased ~21 and ~12 fold, respectively, in the presence of the Fe-specific chelator EDDHA (Fig. 1B). The effect of EDDHA could be prevented by addition of FeSO₄ (Fig. 1B). This confirmed iron-dependent regulation and suggested that LhaSTA is involved in iron acquisition.

LhaSTA allows bacterial proliferation on heme as a source of nutrient iron

LhaSTA is encoded within the *isd* operon and the Isd system facilitates the acquisition of heme from hemoglobin [12, 16]. Therefore, we speculated that LhaSTA might also be involved in the transport of heme. To test this, we used allelic replacement and created isogenic deletion mutants in *S. lugdunensis* N920143 lacking either *lhaSTA* or *isdEFL* the latter of which encodes the conventional lipoprotein-dependent heme transporter of the *isd* locus. Further we created a Δ*lhaSTA*Δ*isdEFL* double mutant. In the presence of 20 µM FeSO₄ all strains showed similar growth characteristics (Fig. 1C). The two single mutants had a slight growth defect compared to wild type when heme was the only iron
source. However, the ΔlhaSTAΔisdEFL mutant showed a significant growth defect under these conditions (Fig. 1D). These data strengthen the hypothesis that LhaSTA is a heme transporter.

**LhaS binds heme**

To confirm the specificity of LhaSTA, we heterologously produced the substrate-specific component LhaS in *E. coli* and purified the protein. We observed that the recombinant protein showed a distinct red color when purified from *E. coli* grown in rich LB medium, which contains heme due to the presence of crude yeast extract [17] (Fig. 2A). The absorption spectrum of the protein showed a Soret peak at 415 nm and Q-band maxima at 537 and 568 nm, suggesting histidine coordination of the heme group. Both the visible color and the spectral peaks were absent when LhaS was purified from *E. coli* grown in heme-deficient RPMI medium (Fig. 2A). We conducted MALDI TOF analysis of holo-LhaS purified from *E. coli* grown in LB and identified two peaks, one of which corresponds to full length recombinant LhaS (24074.437 Da expected weight), and the other to heme (616,1767 Da expected weight). Importantly, the heme peak was not detectable when LhaS was purified from RPMI (Fig 2B and 2C). Furthermore, ESI MS confirmed the presence of heme only in LhaS samples purified from LB (Figure 2 - figure supplement 1). Using the extinction coefficients of LhaS and heme we calculated a heme-LhaS binding stoichiometry of 1:0.6 for the complex isolated from heme containing medium.

**LhaSTA represents an iron acquisition system**

Next, we sought to investigate whether LhaSTA represents a functional and autonomous iron acquisition system. LhaSTA is located within the *isd* locus of *S. lugdunensis*, which, besides the conventional heme membrane transporter IsdEFL, also encodes the hemoglobin receptor IsdB and the cell wall-anchored proteins IsdJ and IsdC. Furthermore, the locus encodes the putative secreted/membrane-associated hemophore IsdK, whose role in heme binding or transport is currently unknown [16], and the autolysin *atl* remodeling the cell wall [18]. To study solely LhaSTA-dependent effects, we disabled all known heme import activity in *S. lugdunensis* by creation of a deletion mutant lacking the entire *isd* operon (from the *atl* gene to *isdB*, Fig. 1A). Then we expressed *lhaSTA* under the
control of its native promoter on a recombinant plasmid in the $\Delta$isd background. *S. lugdunensis* has been reported to degrade nutritional heme in an IsdG-independent fashion due to an unknown enzyme (OrfX) [19]. Therefore, we speculated that heme degradation in this strain might still be possible. LhaSTA deficient and proficient strains showed comparable growth in the presence of FeSO$_4$ (Figure 1 - figure supplement 1). However, only the *lhaSTA* expressing strain was able to grow in the presence of heme as sole source of nutrient iron (Fig. 3A). To further support a role for LhaSTA in iron import, we isolated the cytosolic fraction of the strains prior and after incubation with heme and measured iron levels using the ferrozine assay [20]. Consistently, we found that LhaSTA expression increased cytosolic iron levels post incubation with heme (Fig. 3B). These data suggest that LhaSTA represents a “bona fide” and functional autonomous iron acquisition system.

LhaSTA enables acquisition of heme from various host hemoproteins

We wondered how *S. lugdunensis* might benefit from a heme specific ECF-transporter when a heme acquisition system is already encoded by the canonical Isd system. Indeed, Isd represents a highly effective heme acquisition system. Interactions between the surface receptor IsdB and the proteinaceous part of hemoglobin are thought to enhance heme release to increase its availability [21, 22]. The downside of this mechanism is the specificity for hemoglobin because heme derived from other host hemoproteins such as myoglobin remains inaccessible. In contrast, the HasA hemophore produced by Gram negative pathogens is reported to bind heme with sufficient affinity to enable heme acquisition from a range of host hemoproteins without the need of protein-protein interactions to enhance heme release [23, 24]. As ECF-transporters are known to have high affinity towards their ligands, we speculated that LhaS might represent a membrane-located high affinity ‘hemophore’ allowing heme acquisition from hemoproteins other than hemoglobin. We explored this idea using the hemoprotein myoglobin which is abundant in muscle tissues. Myoglobin was previously reported to not interact with *S. lugdunensis* IsdB [16] and is therefore unlikely to be a substrate for the Isd system. We analyzed the growth of the *S. lugdunensis* wild type (WT) strain as well as of the isogenic *lhaSTA* deficient strain (Fig 4A) on human hemoglobin (hHb) or on equine myoglobin (eqMb) as sole sources of nutrient iron (Fig. 4B). Unlike the WT strain, the *lhaSTA* deficient strain displayed a mild
proliferation defect on hHb and a pronounced growth defect on equine eqMb (Fig. 4B). Interestingly, lhaSTA deficiency did not impact proliferation on hemoglobin-haptoglobin (Hb-Hap) complexes suggesting Isd-dependent acquisition of heme from Hb-Hap. These data strongly indicate that LhaSTA possesses a hemoprotein substrate range that differs from that of the Isd system. To further validate this, we used the above-described S. lugdunensis isd mutant expressing lhaSTA (Fig. 4A) and tested its ability to proliferate on a range of different hemoproteins. (Fig. 4C). We found that lhaSTA expression enabled growth on hemoglobin (human and murine origin) and myoglobin (human and equine origin) as well as with hemopexin (Hpx). Consistent with the above observations, Hb-Hap complexes did not enable growth of the lhaSTA proficient strain strengthening the notion that Hb-Hap acquisition is Isd-dependent. These data further indicate that LhaSTA allows extraction and usage of heme from a diverse set of host hemoproteins, thus expanding the range of hemoproteins accessible to S. lugdunensis.

To confirm the activity of LhaSTA at the biochemical level, we isolated E. coli-derived membrane vesicles that carried apo-LhaS. Following incubation of the vesicles with or without different host hemoproteins, LhaS was purified using affinity chromatography. Heme saturation of LhaS was assessed using SDS-PAGE and tetramethylbenzidine (TMBZ) staining, a reagent that turns green in the presence of hemin-generated peroxides [25] (Fig. 4D). In the absence of hemoproteins during incubation, apo-LhaS did not stain with TMBZ, but TMBZ staining was observed after incubation with all the hemoproteins tested except for Hb-Hap. These data nicely correlate with the ability of the lhaSTA proficient strain to grow on all hemoproteins but Hb-Hap complexes.

**LhaSTA allows usage of human host hemoproteins as an iron source**

Usage of host derived hemoproteins requires the combined action of hemolytic factors to damage host cells as well as hemoprotein acquisition systems to use the released hemoproteins. Since we realized that the S. lugdunensis N920143 strain is non-hemolytic on sheep blood agar, we reproduced the Δisd deletion as well as the plasmid-based expression of lhaSTA in the hemolytic S. lugdunensis N940135 strain. As for N920143, LhaSTA-dependent usage of hemoproteins was also observed in the N940135 background (Figure 4 - figure supplement 1).
We speculated that the expression of LhaSTA is beneficial to *S. lugdunensis* during invasive disease as it allows usage of a wide range of hemoproteins as iron sources. To test this, we attempted to establish septic disease models for *S. lugdunensis*. However, we found that *S. lugdunensis* N940135 was unable to establish systemic disease in mice. Even when infected with $3 \times 10^7$ CFU/animal, mice did not show signs of infection (weight loss / reduced movement). Three days post infection, the organs of infected animals showed low bacterial burdens frequently approaching sterility (Figure 5 - figure supplement 1) and the expression of LhaSTA did not increase the bacterial loads within the organs. We speculate that the presence of human-specific but lack of mouse-specific virulence factors might reduce *S. lugdunensis* pathogenesis in mice. Little is known about virulence factors encoded by *S. lugdunensis*, however, human specific toxins that lyse erythrocytes to release nutritional hemoglobin have previously been described for *S. aureus* [26]. To further assess this, we performed hemolysis assays using human as well as murine erythrocytes (Figure 5 - figure supplement 2). Hemolytic activity of *S. lugdunensis* culture filtrates was low compared to those of *S. aureus*. Nevertheless, we observed lysis of human erythrocytes while murine erythrocytes were not affected by *S. lugdunensis* culture filtrates. This suggests human specific factors mediating host cell damage (Figure 5 - figure supplement 2).

Therefore, we used an *ex-vivo* model to investigate whether LhaSTA facilitates the usage of human cells as a source of iron. First, we supplied freshly isolated human erythrocytes as a source of hemoglobin. Figure 5A shows, that the presence of human erythrocytes allowed significantly improved growth of the Isd deficient but LhaSTA-expressing strain. Secondly, we used a human cardiac myocyte cell line as a source of iron. *S. lugdunensis* is associated with infective endocarditis and myocytes are a source of myoglobin which can be acquired via LhaSTA. Indeed, we found that *lhaSTA* expression enhanced the growth of *S. lugdunensis* in the presence of cardiac myocytes.

In conclusion, our results suggest that LhaSTA represents a novel broad-range heme-acquisition system that expands the hemoprotein substrate range accessible to *S. lugdunensis* to overcome nutritional iron restriction (Fig. 6).

**Discussion**
Nutritional iron restriction represents an effective host strategy to prevent pathogen proliferation within sterile tissues. In turn, bacterial pathogens have developed a range of strategies to overcome nutritional iron limitation during infection. Amongst these is the production and acquisition of siderophores which scavenge the smallest traces of molecular iron to make it biologically available.

The highly virulent *S. aureus* species produces the siderophores staphyloferrin A (SF-A) and staphyloferrin B (SF-B) which are important during infection [27, 28]. *S. lugdunensis* is associated with a series of cases of infective endocarditis and the course of disease mimics that of *S. aureus* endocarditis. In contrast to *S. aureus*, *S. lugdunensis* does not produce endogenous siderophores [29], suggesting that the iron requirements during infection must be satisfied through alternative strategies.

Host hemoproteins can be used by pathogens to acquire iron-containing heme and a plethora of hemoproteins are available during infection. Hemoglobin or myoglobin becomes available if the intracellular pool of the host is tapped by secretion of hemolytic factors. Alternatively, host hemopexin or hemoglobin-haptoglobin complexes involved in heme/hemoglobin turnover are extracellularly available to pathogens. Host hemoproteins are characterized by a remarkable affinity towards heme: Both, globin and hemopexin bind heme with dissociation constants (Kds) smaller than 1 pM [30, 31].

The usage of heme by invasive pathogens is widely distributed, however, the molecular pathways and hemoprotein range availability differs dramatically (see [9] for an excellent review). Iron dependent surface determinant loci are used to acquire heme from hemoglobin by several Gram positive pathogens including *S. aureus* [32], *S. lugdunensis* [11, 12], *Bacillus anthracis* [33], *Streptococcus pyogenes* [34] and *Listeria monocytogenes* [35].

ABC transporters of the Energy-coupling factor type (ECF) are trace element acquisition systems [3, 4]. ECF-type transporters are characterized by high affinity towards their ligands and ECF systems specific for the vitamins riboflavin [36], folate [37], thiamine [38], biotin [39], cobalamine [40, 41], pantothenate [42, 43], niacin [44] and pyridoxamine [45] as well as for the trace metals nickel and cobalt [46, 47] have been described. However, ECF-transporters that allow iron acquisition have so far remained elusive.

Now we show that *S. lugdunensis* encodes the iron regulated ECF-transporter LhaSTA. LhaS binds heme and enables accumulation of iron within the cytoplasm. Therefore, the system represents a
new type of “bona fide” iron acquisition system. Recombinant LhaS acquired heme from human and
murine hemoglobin, from human and equine myoglobin as well as from human hemopexin. The
ability of LhaS to accept heme from several sources strongly suggest an affinity-driven mechanism
relying on passive diffusion of heme between proteins rather than on active extraction. Such a
mechanism has been suggested for HasA-type hemophores of Gram negative pathogens such as
*Serratia marcescens, Yersina pestis* and *Pseudomonas aeruginosa* [48, 49]. Similar to LhaS, HasA has
been shown to possess a broad hemoprotein substrate range and allows the usage of hemoglobin from
different species as well as myoglobin and hemopexin [48]. This ability of HasA was attributed to its
high affinity towards heme (Kd = 0.2 nM) [23]. ECF-specificity subunits frequently possess Kds
towards their ligands in the low nanomolar to picomolar range [4], supporting the idea that LhaS
might directly accept heme from hemoproteins. We attempted isothermal titration calorimetry to
determine the affinity of LhaS towards heme, but our efforts failed to deliver a precise Kd. However,
co-purification of heme with heterologously expressed LhaS suggests that the off-rates are low,
consistent with high-affinity binding. Therefore, the system might be superior to heme acquisition
systems, which depend on specific interactions between bacterial hemoprotein-receptors and host
hemoproteins to extract heme. The *S. aureus* Isd system is well-studied in this regard. The surface
located receptor IsdB binds hemoglobin through its N-terminal NEAT domain (IsdB-N1). This
binding is proposed to induce a steric strain that facilitates heme dissociation. Heme is then captured
by the C-terminal NEAT domain (isdB-N2) and transported across the cell wall and membrane [22,
28, 50-52]. Similarly, the secreted hemophores IsdX1 and IsdX2 of Bacillus anthracis possess NEAT
motifs and perform the same two-step process as IsdB of *S. aureus* to acquire heme [53]. This
mechanism harbours the disadvantage of facilitating heme acquisition only from a single hemoprotein.
IsdB allows acquisition from hemoglobin but does not interact with myoglobin or hemopexin [51] and
even hemoglobin from different species reduces the efficacy of the system [54, 55]. The same is true
for IsdB of *S. lugdunensis* [16]. *Haemophilus influenza* uses the specific interaction between the
surface exposed receptor HxuA and hemopexin to facilitate heme dissociation. Heme is subsequently
captured by HxuC [56, 57]. Again, the specificity for hemopexin prevents usage of other hemoproteins
by HxuA. Specific interactions between LhaS and multiple host hemoproteins seem unlikely,
suggesting that the superior affinity of LhaS towards the heme group bypasses the need for protein-
protein interactions and enables usage of different hemoproteins. However, additional experimental
evidence is required to strengthen this hypothesis of passive heme transfer.

The LhaSTA operon of *S. lugdunensis* is located within the *isd* operon which encodes the
hemoglobin receptor IsdB, the cell wall-anchored, heme-binding proteins IsdJ and IsdC as well as the
conventional heme membrane transporter IsdEFL. Deletion of *lhaSTA* in combination with *isdEFL* did
not completely abrogate acquisition of free heme. A similar effect has been observed in *S. aureus*
suggesting the presence of additional, low affinity heme transporters within these species [58].
Furthermore, a putative secreted/membrane associated hemophore (IsdK) is encoded within the operon
[16]. Interestingly, we show LhaSTA to be functionally independent of the Isd cluster because
LhaSTA-dependent usage of all host hemoproteins except for Hb-Hap was observed in the absence of
all Isd-associated proteins. This indicates that LhaSTA does not rely on Isd-dependent funneling of
heme across the cell wall, but also raises interesting questions about the spatial organization of heme
acquisition and donor proteins. For an efficient transfer of heme between host hemoproteins and LhaS
one would expect that spatial proximity between the proteins is required. Yet, LhaS is situated in the
bacterial membrane and host hemoproteins are too large (hemoglobin ~ 64-16 kd (tetramer-monomer),
myoglobin ~16 kDa, Hemopexin-heme ~70.6 kDa) to readily penetrate the peptidoglycan layer of
Gram-positive bacteria. However, it has been shown that staphylococcal peptidoglycan contains pores
that might allow access of proteins to the bacterial membrane [59, 60]. Along this line, it is tempting
to speculate that surface receptor-dependent acquisition of Hb-Hap might be needed as these
complexes exceed 100 kDa and might be unable to access the bacterial membrane. However, we also
observed that recombinant LhaS did not accept heme from Hb-Hap which might indicate that binding
of haptoglobin to hemoglobin increases the strength of heme binding to the protein complex. Such an
effect of haptoglobin is to our knowledge not known and might be interesting for further investigation.

We failed to establish a functional mouse model of systemic disease to study the *in vivo* role of
LhaSTA for the pathogenicity of *S. lugdunensis*. The reasons for this can be plentiful as little is known
about virulence factors of *S. lugdunensis*. Genome analysis showed that *S. lugdunensis* lacks the wide
variety of virulence and immune evasion molecules found in *S. aureus* [11]. This is the most likely
explanation for the apparent reduced virulence of *S. lugdunensis* in mice. Nevertheless, the co-existence of the Isd and LhaSTA heme-acquisition system in this species may represent a virulent trait and be required for invasive disease. In line with this, most *S. lugdunensis* strains are highly hemolytic on blood agar plates suggesting that the release of hemoproteins from host cells can be achieved by this species. The hemolytic SLUSH peptides [61] of *S. lugdunensis* resemble the β-PSMs of *S. aureus* [62]. Additionally, the sphingomyelinase C (β-toxin) is conserved in *S. lugdunensis* [11]. However, recent research suggested that *S. aureus* targets erythrocytes specifically using the bi-component toxins LukED and HlgAB recognising the DARC receptor [26]. This creates human specificity. Whether similar mechanisms are used by *S. lugdunensis* is unclear, but we found that, in contrast to human cells, *S. lugdunensis* failed to lyse murine erythrocytes. This suggests that host specific virulence factors are present in *S. lugdunensis*. Bi-component toxin genes are not located in the chromosome but genes encoding a streptolysin-like toxin were identified [11].

We found that LhaSTA facilitated growth of *S. lugdunensis* in the presence of human cells such as erythrocytes and myelocytes strongly suggesting that the system allows usage of these cells during invasive disease.

Altogether our experiments identify LhaSTA as an ECF-transporter able to acquire iron and place this important class of nutrient acquisition system in the context of bacterial pathogens and immune evasion strategies. During the revision of this manuscript Chatterjee and colleagues published the identification of a heme-specific ECF transporter in streptococci [63]. In addition, a preprint manuscript that reports the identification of a heme-specific ECF transporter in *Lactococcus sakei* is present in bioarchives [64]. Although these transporters seem functionally redundant to the one of *S. lugdunensis* described here, the specificity subunits of the systems show remarkably little amino-acid sequence similarity. This suggests that the genes encoding them might have developed independently in bacterial species. This was also suggested for the cobalamin-specific components BtuM and CbtT which bind the same ligand despite little sequence conservation [40].

Additional experiments are required to determine whether heme-specific ECF transporters are also present in other bacterial pathogens and the biochemical properties of heme-binding need to be
further characterized to better understand the role of these systems in overcoming nutritional iron limitation.

Methods:

Table 1: Bacterial strains, plasmids and chemicals used in this study

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### Chemicals

If not stated otherwise, reagents were purchased from Sigma.

### Bacterial strains and growth in iron limited media

All bacterial strains generated and/or used in this study are listed in Table 1. For growth in iron limited conditions, bacteria were grown overnight in Tryptic Soy Broth (TSB) TSB (Oxoid). Cells were harvested by centrifugation, washed with RPMI containing 10 µM EDDHA (LGC standards), adjusted to an OD$_{600}$ = 1 and 2.5 µl were used to inoculate 0.5 ml of RPMI+ 1 % casamino acids (BACTO) + 10 µM EDDHA in individual wells of a 48 well microtiter plate (NUNC). As sole iron source 200 nm porcine hemin (Sigma), 2.5 µg/ml human hemoglobin (own preparation), 10 µg/ml human myoglobin (Sigma) or equine myoglobin (Sigma), 117nM human haptoglobin-hemoglobin or 200 nM hemopexin-heme (Sigma) were added to the wells. Bacterial growth was monitored using an Epoch2 reader (300 rpm, 37°C). The OD$_{600}$ was measured every 15 minutes.

### Creation of markerless deletion mutants in *S. lugdunensis*

For targeted deletion of *lhaSTA* and *isdEFL*, 500 bp DNA fragments upstream and downstream of the genes to be deleted were amplified by PCR. A sequence overlap was integrated into the fragments to allow fusion and creating an ATG-TAA scar in the mutant allele. The 1 kb deletion fragments were created using spliced extension overlap PCR and cloned into pIMAY. All the oligonucleotides are summarized in Supplementary File 1. Targeted mutagenesis of *S. lugdunensis* was performed using allelic exchange described elsewhere [65]. The plasmids and the primers used are listed in Table 1 and Supplementary File 1, respectively.

### Heterologous expression of LhaS and membrane vesicle preparation

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<th>chemical compound, drug</th>
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<th>BIO RAD</th>
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LhaS was overexpressed with a N-terminal deca-His tag using pQE-30 in *E. coli* XL1 blue in either Lysogeny broth (LB) medium or RPMI+1% casamino acids. 100 ml overnight culture in LB with 100 μg ml⁻¹ ampicillin was harvested by centrifugation and washed once in PBS. Cells were resuspended in 5 ml PBS and used for inoculation of 2 L RPMI + 1% casamino acids or LB medium. Cells were allowed to grow at 37°C to an OD₆₀₀ = 0.6 - 0.8. Expression was induced by adding 0.3 mM IPTG for 4 – 5 h at 25°C. Cells were harvested, washed with 50 mM potassium phosphate buffer (KPi) pH 7.5, and lysed through 3 rounds of sonification (Branson Digital Sonifier; 2 min, 30% amplitude), in presence of 200 μM PMSF, 1 mM MgSO₄ and DNaseI. Cell debris were removed by centrifugation for 30 min at 7000 rpm and 4°C. The supernatant was centrifuged for 2 h at 35000 rpm and 4°C to collect membrane vesicles (MVs). The MV pellet was homogenized in 50 mM KPi pH 7.5 and flash frozen in liquid nitrogen, stored at -80°C and used for purification.

**Purification of LhaS**

His-tagged LhaS MVs were dissolved in solubilisation buffer (50 mM KPi pH 7.5, 200 mM KCl, 200 mM NaCl, 1% (w/v) n-dodecyl-b-D-maltopyranosid (DDM, Roth) for 1 h at 4°C on a rocking table. Non-soluble material was removed by centrifugation at 35000 rpm for 30 min and 4°C. The supernatant was decanted into a poly-prep column (BioRad) containing a 0.5 ml bed volume Ni²⁺-NTA sepharose slurry, equilibrated with 20 column volumes (CV) wash buffer (50mM KPi pH 7.5, 200mM NaCl, 50 mM imidazole, 0.04 % DDM) and incubated for 1 h at 4°C while gently agitating. The lysate was drained out of the column and the column was washed with 40 CV wash buffer. Bound protein was eluted from the column in 3 fractions with elution buffer (50mM KPi, pH 7.5, 200 mM NaCl, 350 mM imidazole, 0.04 % DDM). The sample was centrifuged for 3 min at 10.000 rpm to remove aggregates and loaded on a Superdex® 200 Increase 10/300 GL gel filtration column (GE Healthcare), which was equilibrated with SEC buffer (50mM KPi pH 7.5, 200 mM NaCl, 0.04% DDM). Peak fractions were combined and concentrated in a Vivaspin disposable ultrafiltration device (Sartorius Stedim Biotec SA).
MV saturation with hemoproteins

MVVs (120 mg total protein content) from RPMI were thawed and incubated for 10 min at RT with each of the following molecules: 5.6 µM heme, 476 µg/ml human hemoglobin, 437 µg/ml equine myoglobin, 5.6 µM hemopexin-heme, 476 µg/ml hemoglobin-haptoglobin. Further purification was performed as described above. After Ni²⁺ affinity chromatography the protein was concentrated and used to measure the peroxidase activity of heme (TMBZ staining).

TMBZ staining of heme

Protein content was determined by Bradford analysis (BIORAD) according to the manufacturer’s protocol. 15 µg protein sample was mixed 1:1 with native sample buffer (BIORAD) and loaded on a Mini-PROTEAN TGX Precast Gel (BIORAD). The PAGE was run at 4°C and low voltage for 2 h in Tris /Glycine buffer (BIORAD). The gel was rinsed with H₂O for 5 min and stained with 50 ml staining solution (15 ml 3,3',5,5'-tetramethylbenzidine (TMBZ) solution (6.3 mM TMBZ in methanol) +35 ml 0.25 M sodium acetate solution (pH 5)) for 1 h at room temperature (RT) while gently agitating. The gel was then incubated for 30 min at RT in the dark in presence of 30 mM H₂O₂. The background staining was removed by incubating the gel in a solution of isopropanol/0.25 M sodium acetate (3:7). Following scanning, the gel was completely destained in a solution of isopropanol/0.25 M sodium acetate (3:7) and stained with the BlueSafe stain (nzytech) for 10 min.

Preparation of human erythrocytes

Human blood was obtained from healthy volunteers and mixed 1:1 with MACS buffer (PBS w/o + 0.05 % BSA + 2 mm EDTA). Erythrocytes were pelleted by density gradient centrifugation in a histopaque blood gradient for 20 min 380 x g at RT. The erythrocyte pellet was washed 3 times with erythrocyte wash buffer (21 mM Tris, 4.7 mM KCl, 2 mM CaCl₂, 140.5 mM NaCl, 1.2 mM MgSO₄, 5.5 mM Glucose, 0.5 % BSA, pH 7.4). Cell count and viability was determined by using the trypan blue stain (BIO RAD).

Purification of human hemoglobin
Human/murine haemoglobin was purified by using standard procedures describe in detail elsewhere [66].

Preparation of saturated hemopexin and haptoglobin

Human hemopexin was dissolved in sterile PBS and saturated with porcine heme in a hemopexin:heme 1:1.3 molar ratio for 1 h at 37°C. This was followed by 48 h dialysis in a Slide-a-Lyzer chamber (ThermoFisher) with one buffer (1 x PBS) change. Haptoglobin was saturated by mixing 4.7 µg/ml haemoglobin with 8.4 µg/ml human haptoglobin for 30 min at 37°C.

Quantification of intracellular iron

Bacteria were grown at 37°C in RPMI + 1% casamino acids to an OD_{600} = 0.6. An aliquot of the culture was collected prior addition of 5 µM heme and 25 µM EDDHA and further incubation at 37°C for 3 hours. At this time point bacteria were collected and resuspended in buffer WB (10 mM Tris-HCl, pH 7, 10 mM MgCl₂, 500 mM sucrose) to an OD_{600} = 50. The bacterial pellet was collected by centrifugation at 8000 rpm for 7 min and resuspended in 1 ml buffer DB (10 mM Tris-HCl, pH 7, 10 mM MgCl₂, 500 mM sucrose, 0.6 mg/ml lysostaphin, 25 U/ml mutanolysin, 30 µl protease inhibitor cocktail (1 complete mini tablet dissolved in 1 ml H₂O (Roche), 1 mM phenylmethanesulfonylfluoride (Roth). The cell wall was digested by incubating at 37°C for 1.5 h, followed by centrifugation at 17000 x g for 10 min at 4°C. Pelleted protoplasts were washed with 1 ml buffer WB, centrifuged and resuspended in 200 µl buffer LB (100 mM Tris-HCl; pH 7, 10 mM MgCl₂, 100 mM NaCl, 100 µg/ml DNaseI, 1 mg/ml RNaseA). Protoplast lysis was performed through repeated cycles (3) of freezing and thawing. The lysate was centrifuged 30 min to pellet membrane fraction and recover the supernatant, which contained the cytosolic fraction and was used for quantification of total intracellular iron content.

Quantification of intracellular iron content by heme uptake was carried out according to Riemer et al. [20] with minor modifications. Briefly, 100 µl of the cytosolic fraction were mixed with 100 µl 50mM NaOH, 100 µl HCl, and 100 µl iron releasing reagent (1:1 freshly mixed solution of 1.4 M HCl and 4.5 % (w/v) KMnO₄ in H₂O). Samples were incubated for 2 h at 60°C in a fume hood. 30 µl iron
detection reagent (6.5 mM ferrozine, 6.5 mM neocuproine, 2.5 M ammonium acetate, 1 M ascorbic acid) was mixed with the samples and incubated for 30 min at 37°C while shaking (1100 rpm). Samples were centrifuged for 3 min at 12000 x g to remove precipitates. 150 µl of the supernatants were transferred to a 96-well microtiter plate and absorbance at 550 nm was measured in a plate reader (BMG Labtech). For determination of iron concentration, FeCl₃ standards in a range of 0 to 100 µM were prepared.

**Measurement of LhaS absorption spectra**

LhaS was purified from LB (holo LhaS) or RPMI (apo LhaS) as described above. 2µl protein sample were loaded on an Eppendorf µCuvette and absorptions spectra were measured at 260 - 620 nm with a BioPhotometer (Eppendorf).

**Characterization of LhaS and heme by mass spectrometry analysis**

MALDI-TOF mass spectra were recorded with a Reflex IV (Bruker Daltonics, Bremen, Germany) in reflector mode. Positive ions were detected and all spectra represent the sum of 50 shots. A peptide standard (Peptide Calibration Standard II, Bruker Daltonics) was used for external calibration. 2,5-dihydroxybenzoic acid (DHB, Bruker Daltonics) dissolved in water/acetonitrile/trifluoroacetic acid (50/49.05/0.05) at a concentration of 10 mg ml⁻¹ was used as matrix. Before the measurements, the samples Lhas-apo (317 µg ml⁻¹) and Lhas+heme (377 µg ml⁻¹) were centrifuged and diluted with MilliQ-H₂O (1:25). An aliquot of 1 µL of the samples was mixed with 1 µL of the matrix and spotted onto the MALDI polished steel sample plate. As the solution dried, the organic solvent evaporated quickly. At this point, the remaining mini droplet was removed gently with a pipette and the remaining sample was air-dried at room temperature.

High resolution mass spectra of Lhas-apo (317 µg ml⁻¹) and Lhas+heme (377 µg ml⁻¹) were recorded on a HPLC-UV-HR mass spectrometer (MaXis4G with Performance Upgrade kit with ESI-Interface, Bruker Daltonics). The samples were diluted with MilliQ-H₂O (1:25) and 3 µL were applied to a Dionex Ultimate 3000 HPLC system (Thermo Fisher Scientific), coupled to the MaXis 4G ESI-QTOF mass spectrometer (Bruker Daltonics). The ESI source was operated at a nebulizer pressure of 2.0 bar,
and dry gas was set to 8.0 L min⁻¹ at 200 °C. MS/MS spectra were recorded in auto MS/MS mode with collision energy stepping enabled. Sodium formate was used as internal calibrant. The gradient was 90 % MilliQ-H₂O with 0.1 % formic acid and 10 % methanol with 0.06 % formic acid to 100 % methanol with 0.06 % formic acid in 20 min with a flow rate of 0.3 mL/min on a Nucleoshell®EC RP-C₁₈ (150 x 2 mm, 2.7 μm) from Macherey-Nagel. [M+H]^+ calculated for C₃₄H₃₂FeN₄O₄^+: 616.1767; found 616.1778 (Δ ppm 1.78).

Calculation of binding stoichiometry

To calculate the putative binding stoichiometry of heme and LhaS the heme concentration in Fig.2A was determined utilizing the extinction coefficient of 58.4 mM⁻¹ cm⁻¹ at 384 nm for heme. The LhaS concentration was determined utilizing the extinction coefficient of 29910 M⁻¹ cm⁻¹ (calculated with ProtParam tool – ExPaSy) at 280 nm.

Human Cardiac Myocytes (HCM)

Primary human cardiac myocytes were purchased from PromoCell (the identity of the cell line was not verified; the culture was negative for mycoplasma) and in 75-cm² culture flasks in 20 ml of myocyte growth medium (PromoCell). Cells were detached with accutase, washed once with RPMI containing 200 µM EDDHA and resuspended in PRMI containing 200 µM EDDHA. 40000 cells per well were used for bacterial growth assays as described above.

Assessing hemolytic activity of S. lugdunensis culture supernatants

S. aureus and S. lugdunensis were grown overnight in TSB. Cells were pelleted and culture supernatants were filter sterilized using a 0.22 μM filter. A 100 μL volume of supernatant was added into 1 mL of PBS containing either 5% v/v murine or human red blood cells. Mixtures were incubated at room temperature without shaking for 48 hours.

Declarations:

Ethics approval and consent to participate: Animal experiments were performed in strict accordance with the European Health Law of the Federation of Laboratory Animal Science
Associations. The protocol was approved by the Regierungspräsidium Tübingen (IMIT1/17). Human Erythrocytes were isolated from venous blood of healthy volunteers in accordance with protocols approved by the Institutional Review Board for Human Subjects at the University of Tübingen. Informed written consent was obtained from all volunteers.

Consent for publication: not applicable

Availability of data and materials: The datasets gained during the current study are available over dryad at https://doi.org/10.5061/dryad.fqz612jqc

Competing interests: The authors declare that they have no competing interests.

Funding: We acknowledge the funding of this project by the Deutsche Forschungsgemeinschaft (DFG) in from of an individual project grant (HE8381/3-1) to SH. SH was supported by infrastructural funding from the Deutsche Forschungsgemeinschaft (DFG), Cluster of Excellence EXC 2124 Controlling Microbes to Fight Infections. DJS was supported by NWO (TOP grant 714.018.003). DEH acknowledges support from the Canadian Institutes of Health Research (PJT-153308). None of the funding bodies was involved in the design of the study, the performance of experiments, data evaluation, writing of the manuscript or the decision about submission.

Acknowledgements

We thank Timothy J. Foster and Libera Lo Presti for critically reading and editing this manuscript. We thank Andreas Peschel for helpful discussion. We thank Sarah Rothfuß and Vera Augsburger for excellent technical support and Imran Malik for the introduction to the ITC technology.

References


Figure 1. LhaSTA represents an iron-regulated heme transporter.

A) Schematic diagram of the isd operon of S. lugdunensis N920143. Coding sequences, direction of transcription and Fur-binding sites are indicated. ABC membrane-transporters are shown in green.

B) Iron-regulated expression of Lha: S. lugdunensis was grown overnight in TSB, TSB + 200 μM EDDHA or TSB + 200 μM EDDHA + 200 μM FeSO₄. Gene expression was quantified by qPCR. Expression was normalized to 5srRNA and to the TSB standard condition using the ΔΔCt method.
Fold differences in gene expression are shown. Data represent mean and SD of four independent experiments. Statistical evaluation was performed using students unpaired t-test (lhaS: t=7.045, df=6; lhaA: t=2.979, df=6)

C/D Growth curves of S. lugdunensis N920143 and isogenic mutants. The wild type (WT) S. lugdunensis N920143 strain and the indicated isogenic null mutant strains were grown in the presence of 20 µM FeSO₄ (C) or 150 nM heme (D) as a sole source of iron. 500 µl of bacterial cultures were inoculated to an OD₆₀₀ = 0,05 in 48 well plates and OD₆₀₀ was monitored every 15 min using an Epoch1 plate reader. For reasons of clarity values taken every 5 hours are displayed. Mean and SD of three experiments are shown. Statistical analysis was performed using one-way ANOVA followed by Dunett’s test for multiple comparisons. * - p < 0,05, **** p < 0,00001

Figure 1- figure supplement 1. LhaSTA dependent growth.

A/B) Proliferation of S. lugdunensis N920143 ∆isd pRB473 and ∆isd pRB473:lhaSTA strains. The indicated strains were grown in the absence of nutritional iron (A) or in the presence of 20 µM FeSO₄ (B). 500 µl of cultures were inoculated to an OD₆₀₀ = 0,05 in 48 well plates and OD₆₀₀ was monitored every 15 min using a Epoch1 plate reader. For reasons of clarity values taken every 5 hours are displayed. Mean and SD of three experiments are shown. Statistical analysis was performed using students unpaired t-test.

Figure 2. LhaS binds heme.

A) Ultraviolet-visible (UV-vis) spectrum of recombinant LhaS. C-terminal His-tagged LhaS was heterologously expressed in E. coli and purified from heme-containing LB medium or heme-free RPMI medium. The UV-vis spectrum of the purified LhaS was measured with a BioPhotometer.

B) and C) MALDI-TOF mass spectra of recombinant LhaS. LhaS (B) was purified out of LB medium and apo-LhaS (C) was purified out of RPMI medium. Mass spectra were recorded with a Reflex IV in reflector mode. All spectra are a sum of 50 shots. Prior to measurements the protein samples were mixed with a 2,5-dihydroxybenzoic acid matrix dissolved in
water/acetonitrile/trifluoroacetic acid (50/49.05/0.05) at a concentration of 10 mg ml\(^{-1}\) and spotted onto the MALDI polished steel sample plate.

Figure 2 – figure supplement 1. High resolution mass spectra of apo- and holo-LhaS.

Spectra of apo-LhaS isolated from *E.coli* grown in RPMI medium (upper panel), holo-LhaS isolated from LB-medium (middle panel) and a heme standard (lower panel) were recorded on a HPLC-UV-HR mass spectrometer. The samples were diluted with MilliQ-H\(_2\)O and applied to a Dionex Ultimate 3000 HPLC system that is coupled to the MaXis 4G ESI-QTOF mass spectrometer.

Figure 3. LhaSTA represents a functionally autonomous iron acquisition system.

A) LhaSTA-dependent proliferation. *S. lugdunensis* N920143 deletion mutant strains lacking the entire *isd* operon and expressing LhaSTA (Δ*isd pRB473:lhaSTA*) or not (Δ*isd pRB473*) from the plasmid pRB473 were grown in the presence of 200 nM heme as a sole source of iron. 500 µl of cultures were inoculated to an OD\(_{600}\) = 0.05 in 48 well plates and OD\(_{600}\) was monitored every 15 min using an Epoch1 plate reader. For reasons of clarity values taken every 5 hours are displayed. Mean and SD of three experiments are shown. Statistical analysis was performed using students unpaired t-test. *** p < 0.0001

B) Intracellular accumulation of iron. Strains were grown in iron limited medium to OD\(_{600}\)= 0.6 and 5 µM heme were added for 3 h. Cell fractionation of 1 ml OD\(_{600}\) = 50 was performed and the iron content of the cytosolic fraction was determined using the ferrozine assay. Data represent the mean and SD of three independent experiments. Statistical analysis was performed using students unpaired t-test (t=5.12729, df=4).

Figure 4. LhaSTA facilitates heme acquisition from a wide range of hemoprotein substrates.

A) Schematic diagram of known heme acquisition systems in the *S. lugdunensis* mutant strains lacking either the genes encoding LhaSTA (Δ*lhaSTA*, left) or the entire *isd* operon and expressing LhaSTA from the plasmid pRB473 (Δ*isd pRB473:lhaSTA*). ABC membrane transporters are shown in green. Cell wall-anchored proteins of the Isd-system are shown in yellow.
Heme/hemoglobin-binding NEAT motifs within each protein are indicated as black boxes. Black arrows indicate the transfer of heme. he: heme; hb: hemoglobin; PG: peptidoglycan.

**B) Growth of *S. lugdunensis* N920143 wild type (WT) and ΔlhaSTA.** Strains were grown in the presence of 20 µM FeSO₄ or 2.5 µg/ml human hemoglobin (hHb) or 10 µg/ml equine myoglobin (eqMb) or 117 nM hemoglobin-haptoglobin complex (Hb-Hap) as a sole source of iron. 500 µl of cultures were inoculated to an OD₆₀₀ = 0.05 in 48 well plates and OD₆₀₀ was measured after 30 h using an Epoch1 plate reader. Mean and SD of three experiments are shown. Statistical analysis was performed using students unpaired t-test. hHb - t=6.0007, df=4; eqMb – t=20.52, df=4; Hb-Hap – t=1.978, df=4.

**C) Growth of *S. lugdunensis* N920143 Δisd pRB473 and Δisd pRB473: lhaSTA.** Strains were grown in the presence of 20 µM FeSO₄ or 2.5 µg/ml hHb or 2.5 µg/ml murine hemoglobin (mHb) or 10 µg/ml human myoglobin (hMb) or 10 µg/ml eqMb or 200 nM human hemopexin (Hpx) or 117 nM Hb-Hap as a sole source of iron. 500 µl of cultures were inoculated to an OD₆₀₀ = 0.05 in 48 well plates and OD₆₀₀ was measured after 30 h using an Epoch1 plate reader. Mean and SD of three experiments are shown. Statistical analysis was performed using students unpaired t-test hHb – t=18.5, df=4; mHb – t=29.03, df=4; hMb – t=25.98, df=4; eqMb – t=3.442, df=4; Hpx – t=77.12 df=4; Hb-Hap t=2.758 df=4.

**D) TMBZ-H₂O₂ stain of TGX gels for heme-associated peroxidase activity.** Membrane vesicles were saturated with excess of hemoprotein (5.6 µM heme, 476 µg/ml hHb, 437 µg/ml eqMb, 5.6 µM Hpx, 476 µg/ml Hb-Hap) or no hemoprotein (-) for 10 min at RT. LhaS was purified, 15 µg protein was loaded on a TGX gel and stained for peroxidase activity with TMBZ-H₂O₂ (upper panel). Gels were destained and subsequently stained with BlueSafe (lower panel) to confirm the presence of the protein in all conditions.

**Figure 4 – figure supplement 1. Growth of *S. lugdunensis* N940135 Δisd pRB473 and Δisd pRB473: lhaSTA.** The indicated strains were grown in the presence of 20 µM FeSO₄ or 2.5 µg/ml human hemoglobin (hHb), 2.5 µg/ml or murine hemoglobin (mHb) or 10 µg/ml human myoglobin.
(hMb) or 10 µg/ml equine myoglobin (eqMb) or 200 nM human hemopexin (Hpx) or 117 nM Hb-Hap as a sole source of iron. 500 µl of cultures were inoculated to an OD$_{600}$ = 0.05 in 48 well plates and OD$_{600}$ was measured every 15 min. For reasons of clarity values taken every 5 hours are displayed. Mean and SD of three experiments are shown. Statistical analysis was performed using students unpaired t-test.

Figure 5. LhaSTA allows usage of host cells as an iron source.

A) Growth of S. lugdunensis N940135 Δisd pRB473:lhaSTA and Δisd pRB473 on human erythrocytes. Strains were grown in the presence of freshly isolated human erythrocytes (10$^5$ cells/ml) as a sole source of iron. 500 µl of cultures were inoculated to an OD$_{600}$ = 0.05 in 48 well plates and OD$_{600}$ was monitored every 15 min using an Epoch1 plate reader. For reasons of clarity values taken every 5 hours are displayed. Mean and SD of three experiments are shown. Statistical analysis was performed using students unpaired t-test. *** p < 0.0001

B) Growth of S. lugdunensis N940135 Δisd pRB473 and Δisd pRB473:lhaSTA on human cardiac myocytes. Strains were grown in the presence of 40000 primary human cardiac myocytes per well as a sole source of iron. Cardiac myocytes were detached and washed once with RPMI+200µM EDDHA prior addition to the wells. 500 µl of cultures were inoculated to an OD$_{600}$ = 0.05 in 48 well plates and OD$_{600}$ was monitored every 15 min using an Epoch1 plate reader. For reasons of clarity values taken every 5 hours are displayed. Mean and SD of three experiments are shown. Statistical analysis was performed using students unpaired t-test. * p < 0.05, ** p < 0.01

Figure 5 - figure supplement 1. Mouse systemic infection model.

C57BL/6 mice were infected either with 3 x 10$^7$ CFU per animal. Mice were sacrificed 72h post infection and CFUs within the indicated organs enumerated. Horizontal lines show the median. Statistical analysis was performed using Mann Whitney test.

Figure 5 - figure supplement 2. Hemolysis of human and mouse erythrocytes.
Human (left and middle panels) or mouse erythrocytes (right panel) were incubated with the filtrated culture supernatants of *S. aureus* (left panel) or *S. lugdunensis* (middle and right panels) for 24 and 48 h.

**Figure 6. Model of heme acquisition in *S. lugdunensis*.**

ABC membrane transporters are shown in green. Cell wall-anchored proteins of the Isd-system are shown in yellow. Heme/hemoglobin-binding NEAT motifs within each protein are indicated as black boxes. Black arrows indicated the transfer of heme. he: heme; hb: hemoglobin; PG: peptidoglycan; Mem: Membrane; Hap: Haptoglobin; Hpx: Hemopexin; Atl: Autolysin.

**Supplementary File 1. Key Resource Table PCR primers.**

PCR and qPCR primers sequences used in this study.
Figure A: Gene regulation diagram showing the expression of the genes involved in iron uptake and storage. The fur boxes indicate the regulatory regions.

Figure B: Bar graph showing the fold change in gene expression compared to TSB. The p-values are 0.0004 and 0.0247 for lhaS and lhaA, respectively.

Figure C: Graph showing the OD600 over time with 20 µM FeSO4. The graph compares WT, ΔisdEFL, ΔlhaSTA, and ΔisdEFLΔlhaSTA.

Figure D: Graph showing the OD600 over time with 150 nM heme. The graph compares WT, ΔisdEFL, ΔlhaSTA, and ΔisdEFLΔlhaSTA.
A

![Absorbance spectrum graph]

- **LhaS from LB**
- **LhaS from RPMI**

B

- **LhaS from LB**
  - LhaS: 24074.437 Da

- **LhaS from RPMI**
  - LhaS: Heme 616.1767 Da

C

- **LhaS from LB**
  - LhaS: Heme 616.1767 Da

- **LhaS from RPMI**
  - LhaS: Heme 616.1767 Da
(A) 200 nM heme

- \( \Delta isd \) pRB473: lhaSTA
- \( \Delta isd \) pRB473

(B) Iron content [µM]

- Prior heme
- After heme

\( p = 0.3272 \) \( p = 0.02595 \)
**A** ΔlhaSTA

**Δisd** pRB473: lhaSTA

- Heme transfer
- Proposed transfer

**B**

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<td></td>
</tr>
</tbody>
</table>

**C**

- p = 0.00039
- < 0.0001
- = 0.0039
- = 0.0191
- < 0.0001
- < 0.0001
- < 0.0001
- = 0.0262
- < 0.0001

**D**

- Heme: 25 kDa, 15 kDa
- Hb: 15 kDa
- eqMb: 25 kDa, 15 kDa
- Hpx: 25 kDa, 15 kDa
- Hb-Hap: 25 kDa, 15 kDa
A  erythrocytes

B  cardiac myocytes

**Δ*isd pRB473:*lhaSTA**

***Δ*isd pRB473:**

\[ \text{OD}_{600} \]

**time [h]**

---

**time [h]**
A) FeSO₄  

B) heme  

C) hHb  

D) mHb  

E) hMb  

F) eqMb  

G) Hpx  

H) Hb-Hap
kidneys

\[ \text{Δ} \text{isd pRB473: lhaSTA} \]

CFU / 2 kidneys

\[ p = 0.8182 \]

spleen

\[ \text{Δ} \text{isd pRB473: lhaSTA} \]

CFU / spleen

\[ p = 0.0542 \]

heart

\[ \text{Δ} \text{isd pRB473: lhaSTA} \]

CFU / heart

\[ p = 0.0931 \]

liver

\[ \text{Δ} \text{isd pRB473: lhaSTA} \]

CFU / g liver

\[ p = 0.8182 \]
Human Erythrocytes

Mouse Erythrocytes

0h

24h

48h

S. aureus

S. lugdunensis

S. lugdunensis