Hepcidin-mediated iron sequestration protects against bacterial dissemination during pneumonia

Kathryn R. Michels,1 Zhimin Zhang,2 Alexandra M. Bettina,1 R. Elaine Cagnina,2 Debora Stefanova,3 Marie D. Burdick,2 Sophie Vaulont,4 Elizabeta Nemeth,5 Tomas Ganz,5 and Borna Mehrad1,2,6

1Departments of Microbiology, Immunology, and Cancer Biology, 2Division of Pulmonary & Critical Care Medicine, Department of Medicine, University of Virginia, Charlottesville, Virginia, USA. 3Departments of Molecular, Cellular, and Integrative Physiology, University of California, Los Angeles, California, USA. 4INSERM U1016, Cochin Institute, Descartes University, Paris, France. 5Department of Medicine, David Geffen School of Medicine, University of California, Los Angeles, California, USA. 6Beirne B. Carter Center for Immunology, University of Virginia, Charlottesville, Virginia, USA.

Introduction

Infections caused by aerobic Gram-negative bacteria are prevalent and dangerous. As a group, these organisms are the most common etiology of nosocomial infections, including most cases of hospital-acquired pneumonia, ventilator-associated pneumonia and healthcare-associated pneumonia, and they are important causes of morbidity and death in hospitalized patients (1). Dissemination of these organisms from the primary site of infection to the bloodstream is especially associated with mortality (2, 3). The progressive rise in antibiotic resistance among Gram-negative bacteria, particularly in the healthcare setting, has critically diminished the available treatment options for these infections. Better mechanistic understanding of the host-pathogen interplay during Gram-negative infections has the potential to lead to novel therapeutic strategies to augment the diminishing armamentarium of traditional antibiotics.

Iron is an essential component of many proteins and a required nutrient for nearly all organisms. In biological systems, free iron ions are present at extremely low concentrations, and the competition for iron between pathogens and the host is important to the outcome of many infections (4). While the iron acquisition mechanisms of aerobic Gram-negative bacteria are known to be essential to their virulence (5), the host mechanisms of iron restriction in these infections are incompletely defined.

Under homeostatic conditions, plasma iron is rapidly turned over with a half-life of < 1 hour, being continuously utilized in erythropoiesis and replenished by splenic macrophages that scavenge iron from senescent erythrocytes (6). The hormone hepcidin is the key regulator of plasma iron concentration that acts by mediating the degradation of the only known cellular iron exporter, ferroportin, thus trapping...
During homeostasis, hepcidin is induced by elevated extracellular and intracellular iron levels via SMAD4 signaling, and is inhibited by anemia and hypoxia. During inflammation, hepcidin is induced as an acute-phase reactant by IL-6 and Activin B via STAT3 and SMAD4, respectively (8, 9). In the context of inflammation, hepcidin mediates acute hypoferremia, which is posited to be a defense mechanism against infectious disease; continual hepcidin production in this setting can infamously cause anemia of chronic inflammation.

The role of hepcidin in disorders of iron metabolism is well established, but its contribution to antimicrobial host defenses is an emerging field. For example, hepcidin appears to protect the host against malaria but to promote the growth of pathogens that reside within the intracellular niche of macrophages (10). Hepcidin-mediated hypoferremia is essential for protection of the host against *Vibrio vulnificus* in a murine model (11), consistent with the clinical observation that individuals with iron-overload conditions are uniquely susceptible to this organism (12). In contrast, the role of hepcidin as a general defense against common bacterial infections, such as those caused by Enterbacteriaceae, remains unknown.

We reasoned that the bloodstream dissemination of microorganisms from the primary site of infection is a critical complication in Gram-negative bacterial infections. In the context of pneumonia, as well as other focal infections caused by aerobic Gram-negative bacilli, bacteremia is strongly associated with sepsis and death (13, 14). We therefore tested the hypothesis that hepcidin-mediated depletion of extracellular iron during Gram-negative pneumonia protects the host by limiting dissemination of bacteria to the bloodstream.

**Results**

*Plasma iron is suppressed during bacterial pneumonia.* We used a well-described model of *Klebsiella pneumonia* caused by a virulent strain of bacteria, delivered directly into the trachea of sedated experimental mice (15, 16). We first characterized the effect of *Klebsiella pneumonia* on iron metabolism in WT mice. As expected, plasma iron concentration decreased quickly in WT mice following infection (Figure 1A), similar to prior reports with acute inflammatory stimuli (9). To assess the availability of iron to bacteria in the lungs, we also measured heme-associated and non–heme-associated iron levels in whole lung homogenates (which includes both intra- and extracellular iron), and in the extracellular space, in the bronchoalveolar lavage (BAL) supernatant (Figure 1, B and C). We found that most lung iron was intracellular, and extracellular iron levels were low and did not change appreciably during infection.
Increased iron availability worsens the outcome of infection. To determine whether iron availability influences host defense during the course of infection, we tested the effect of transient increase in extracellular iron availability, achieved by i.p. injection of ferric ammonium citrate, on the outcome of *Klebsiella pneumonia*. Mice treated with iron prior to intrapulmonary *K. pneumoniae* challenge developed transient elevation in plasma iron that resolved completely within 12 hours (Figure 1D) without influencing lung iron (Supplemental Figure 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.92002DS1), consistent with prior studies indicating that excess iron is rapidly cleared from the blood and redistributed to the BM, spleen, and liver (6). This transient increase in iron availability during the first 24 hours after infection was associated with a markedly increased mortality from pneumonia associated with increased bacterial burden in the lung and the blood (Figure 1, E and F). Since WT mice developed hypoferrremia upon infection with *K. pneumoniae*, and excess iron was associated with poor infection outcome, we hypothesized that endogenous iron regulation was important for controlling infection.

*Hepcidin regulates plasma iron during bacterial pneumonia by an IL-6–dependent mechanism*. To assess the role of hepcidin in mediating hypoferrremia during pneumonia, we began by characterizing hepcidin expression during the infection. Infection resulted in a marked increase in hepcidin protein levels in plasma, in parallel with the development of hypoferrremia (Figure 2A). Hepcidin protein concentration also increased in the

![Figure 2. Hepcidin induction during *Klebsiella pneumonia*. Time 0 represents naive animals. (A) Hepcidin protein levels in the plasma and whole lung homogenate following infection. Data represent mean ± SEM. (B–D) Hepcidin mRNA in the liver, buffy coat, and lung normalized to GAPDH expression and then to day 0. Box and whisker plots show median (line within box), upper and lower quartiles (upper and lower box boundaries), and total range (bars); n = 4–6 animals per time point in each panel; *, **, and *** denote P values of <0.05, <0.01, and <0.001, respectively (1-way ANOVA, multiple comparison for linear trend for panels A–C, t test for panel D).](https://doi.org/10.1172/jci.insight.92002)

![Figure 3. The role of IL-6 in hepcidin induction during *Klebsiella pneumonia*. (A) Animals were treated daily with an isotype control or IL-6 neutralizing monoclonal antibody and assessed for bacterial burden on day 2 of infection. Horizontal lines represent median, and each circle represents 1 animal; animals with no detectable bacteria are reported to have a bacterial burden of 1 CFU on the logarithmic scale; * and **** denote P values of <0.05 and <0.0001 by Mann-Whitney U test, respectively. (B and C) Plasma hepcidin protein and iron levels in uninfected and infected animals treated with isotype or IL-6 neutralizing antibody. Box and whisker plots show median (line within box), upper and lower quartiles (upper and lower box boundaries), and total range (bars) combined from 2 experiments; n = 8–9 animals per group; **P < 0.01, one way ANOVA Tukey post-test.](https://doi.org/10.1172/jci.insight.92002)
Hepcidin mediates hypoferremia during bacterial pneumonia and is required for survival. Reduced availability of plasma iron in the context of acute inflammation can occur as a result of both hepcidin-dependent and -independent mechanisms (23–25). We therefore tested the contribution of hepcidin in regulation of plasma iron levels in the context of bacterial pneumonia. In contrast to reduction in plasma iron during infection in WT mice, plasma iron in hepcidin-deficient mice became more elevated during the infection (Figure 4A), indicating that hepcidin is essential for controlling plasma iron levels following infection. Consistent with reports of tissue iron overload in hepcidin-deficient mice (18, 26), we found elevated lung iron levels in hepcidin-deficient as compared with WT mice, but these values did not change over the infection period (Figure 4B).

We next assessed the contribution of hepcidin-mediated iron sequestration to host defense against *K. pneumoniae* and found that hepcidin-deficient animals were markedly more susceptible to bacterial pneumonia, demonstrating 100% mortality compared with 50% mortality in WT controls (Figure 4C), comparable to mice treated with exogenous iron during infection (Figure 1, D and E). Hepcidin-deficient mice exhibited 23% greater incidence of bacteremia and >100-fold higher median blood bacterial concentration, as well as ~10-fold higher bacterial burden in the lungs and liver as compared with WT mice (Figure 4D).

Since the hepatocytes in hepcidin-deficient animals are iron loaded, we reasoned that the increase of iron in infected hepcidin−/− mice could originate from greater release of iron from damaged hepatocytes. We therefore quantified alanine aminotransferase and aspartate aminotransferase in the blood of infected mice to assess liver damage and found neither enzyme to be elevated in hepcidin−/− as compared with WT mice in the context of infection (Supplemental Figure 2), arguing against disproportionate

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**Figure 4. The role of hepcidin in *Klebsiella* pneumonia and dissemination.** (A and B) Comparison of plasma or lung nonheme iron levels in WT and hepcidin−/− mice before and 2 days after infection. Box and whisker plots show median (line within box), upper and lower quartiles (upper and lower box boundaries), and total range (bars) of data from 2 experiments; n = 4–9 per group; ** and **** denote P values of <0.01 and <0.0001 by one-way ANOVA with Tukey post test, respectively. (C) Survival of WT and hepcidin−/− mice during pneumonia. n = 26–30 mice per group, combined from 3 independent experiments; ***P < 0.001 Mantel-Cox test. (D) Bacterial burden on day 2 of infection in WT and hepcidin−/− mice in indicated organs. Pooled data from 3 independent experiments; n = 23–26. Horizontal lines represent median and each circle represents one animal; animals with no detectable bacteria are reported to have a bacterial burden of 1 CFU on the logarithmic scale; ** and **** denote P values of <0.01 and <0.0001, respectively, using the Mann-Whitney U test.
hepatic injury as a contributor to worse outcomes in hepcidin-deficient hosts.

In order to assess whether the beneficial effects of hepcidin are dependent on iron, we used established protocols to induce severe iron-deficiency in hepcidin-deficient mice (27). We found that iron-depleted hepcidin–/– animals did not develop hyperferremia during infection, had lower body iron stores than WT mice, and were no more susceptible to infection compared with WT mice (Supplemental Figure 3, A and B). Taken together, these data suggest that the hepcidin-mediated control of bacterial growth during infection is iron dependent.

Excess iron promotes the growth of \( K.\) pneumoniae in plasma. To assess whether the increased susceptibility of hepcidin-deficient mice to infection was attributable to increased bacterial growth in the high-iron environment of the plasma, we assessed the effect of iron supplementation on the in vitro growth of \( K.\) pneumoniae in human plasma. Supplementing agar composed of 90% human plasma with ferric ammonium citrate at physiologically relevant concentrations incrementally raised the transferrin saturation of the sample, resulting in the appearance of non–transferrin-associated labile pool iron (Figure 5A). The appearance of labile pool iron at transferrin saturation greater than 70% is consistent with prior reports (28, 29). \( K.\) pneumoniae growth was suppressed in plasma but was permitted by iron supplementation, consistent with reports that transferrin inhibits bacterial growth (30) (Figure 5B). Interestingly, we found that, in plasma preparations from 3 independent donors, addition of ferric iron resulted in \( K.\) pneumoniae outgrowth coincided with the appearance of labile iron in the plasma, rather than with the total amount of iron citrate added. These data suggest that ferric ammonium citrate supports \( K.\) pneumoniae growth in plasma by providing iron, rather than acting as a carbon or nitrogen source.

We next sought to test the hypothesis that, during in vivo infection, elevated plasma iron is sufficient to enhance growth of blood-borne bacteria. To test this, we subjected WT mice with an i.p. injection of PBS or ferric ammonium citrate prior to i.v. challenge with \( K.\) pneumoniae. After 24 hours, mice treated with ferric ammonium citrate had higher blood bacterial content than mice treated with PBS, indicating that
Elevated plasma iron directly mediates increased growth of *K. pneumoniae* in the bloodstream (Figure 5C). Interestingly, mice treated with ferric ammonium citrate also had higher bacterial burdens in the lungs. We interpret this as evidence that, in mice with increased bacteremia, bacterial seeding of the lung from the bloodstream contributes to higher lung bacterial content. Similar to prior reports (31), mice treated with iron also had marked elevation of plasma hepcidin as compared with mice treated with PBS in response to acute iron loading (Figure 5D), indicating that hepcidin induction alone is not sufficient to protect mice from fatal infection if plasma iron levels are high and supporting the hypothesis that the beneficial effects of hepcidin are mediated via induction of hypoferremia, rather than an iron-independent effect.

Hepcidin-deficient mice have intact cellular immunity against *Klebsiella*. Iron deficiency and iron overload have complex effects on immune responses (32), and phagocytes from iron-overloaded hosts may have altered antibacterial properties that could contribute to impaired host defenses of hepcidin-deficient mice independent of the effects of acute hyperferremia on bacteria. In order to assess the functionality of the innate immune response in hepcidin-deficient mice, we analyzed leukocyte populations in the blood and BAL (Figure 6, A–F). We found similar concentrations of circulating neutrophils, and Ly6<sup>Ch</sup> and Ly6<sup>Cl</sup> monocytes in WT and hepcidin<sup>−/−</sup> mice at baseline and 2 days following infection. The number of BAL alveolar macrophages, neutrophils, and monocyte-derived macrophages were also similar between WT and hepcidin-deficient hosts, both at baseline and following infection, arguing against a defect in leukocyte quantity or recruitment in the absence of hepcidin. Since oxidative burst is an iron-dependent process, we compared the generation of reactive oxygen species in lung and blood leukocyte populations and found no difference between infected WT and hepcidin-deficient mice (Supplemental Figure 4). Similarly, neutrophils from WT and hepcidin-deficient mice exhibited comparable ex vivo bacterial killing (Figure 6G). While these data do

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**Figure 6. Leukocytes responses during *Klebsiella pneumonia*. (A–F) BAL and blood were harvested from WT and hepcidin<sup>−/−</sup> mice before and 2 days following infection. Box and whisker plots show median (line within box), upper and lower quartiles (upper and lower box boundaries), and total range (bars). n = 4–5 per group, one way ANOVA Tukey post-test. No significant differences were found for any WT to hepcidin<sup>−/−</sup> comparison. (G) Freshly isolated neutrophils from WT and hepcidin<sup>−/−</sup> mice were coincubated with 2 × 10<sup>5</sup> CFU of *K. pneumoniae* for 1 hour. Total surviving bacteria were enumerated from each well. n = 3 independent experiments per group; data were analyzed using a 2-way ANOVA, and no differences were found between groups.
not formally exclude the possibility that hepcidin deficiency impairs the immune system in some other way, they suggest intact cellular innate immune mechanisms against *K. pneumoniae* in hepcidin–/– mice.

**Restoring hypoferrremia is sufficient to protect hepcidin-deficient mice from infection.** Impaired hepcidin production is common in the clinical setting — for example, in patients with liver disease (33–36) — and our data suggests a mechanism by which these hosts may be predisposed to Gram-negative bacterial infections. As proof-of-principle of a potential therapeutic approach in this population, we assessed the effect of minihepcidin, a hepcidin analogue, on the outcome of *K. pneumoniae* infection in hepcidin-deficient hosts. We used a previously described synthetic peptide based on the 9 N-terminal amino acids of the hepcidin molecule, which are necessary and sufficient to induce the degradation of ferroportin (37, 38). Daily treatment with the hepcidin analogue beginning on the day of infection resulted in markedly lowered plasma iron levels in infected hepcidin-deficient mice as compared with vehicle-treated controls without affecting the total lung iron content (Figure 7, A and B), consistent with utilization of plasma iron in erythropoiesis (6). The treatment also resulted in >99% reduction in median lung and blood bacterial burden in infected hepcidin-deficient mice and 40% reduction in mortality compared with vehicle alone (Figure 7, C and D). These data provide evidence that the acute induction of hypoferrremia during infection in hepcidin-deficient hosts may represent a plausible therapeutic strategy.

**Discussion**

Although the competition for iron is thought to be a general feature of antimicrobial host response, the mechanisms and contribution of iron restriction in most infections are undefined. In particular, hepcidin-mediated hypoferrremia has only been implicated in host defense against *V. vulnificus*, an organism with specific affinity for iron-overloaded hosts, but its role in defense against common pathogens and in normal hosts is not known (10). We showed that hepcidin was necessary for protection against bacterial dissemination in Gram-negative pneumonia and that treatment with iron citrate was sufficient to permit bacterial outgrowth in WT mice. Finally, we showed that treatment with minihepcidin restored hypoferrremia and rescued iron-overloaded, hepcidin-deficient hosts from the infection. Taken together, these data suggest that hepcidin-mediated iron sequestration protects against Gram-negative pneumonia.

Our data suggest that the hepcidin-dependent protection was mediated by reducing plasma iron, as indicated by our observation that WT mice treated with iron citrate were susceptible to *K. pneumoniae* infection despite robust hepcidin induction (Figure 1, E and F) and that hepcidin-deficient mice were protected from
infection if starved of iron (Supplemental Figure 3B). While there may be iron-independent effects of ferric ammonium citrate administration that we have not accounted for, such as bacterial fermentation of citrate, we are encouraged that the phenotype of iron citrate–treated mice is remarkably similar to that of hepcidin−/− mice in terms of death and bacterial burden. Furthermore, we found that hepcidin-deficient mice treated with minihepcidin were robustly protected from infection despite having tissue iron overload, suggesting that restriction of iron in the plasma is sufficient for protection (Figure 7, C and D). Hepcidin exhibits antimicrobial activity against a broad array of pathogens in vitro, and an iron-independent direct microbicidal role for hepcidin has thus been suggested (17, 39, 40). Arguing against such a direct microbicidal role, we report plasma hepcidin levels at concentrations much lower than those required to mediate killing in vitro (39). Similarly, administration of a hepcidin analogue, which lacks the β-defensin motif required for its antimicrobial activity (37), rescued the phenotype of hepcidin-deficient mice (Figure 7, C and D).

Although hepcidin-independent mechanisms of hypoferremia have been reported after challenge with inflammatory stimuli (23, 25, 41), we found that hypoferremia during pneumonia was completely abrogated in hepcidin-deficient mice, and in fact, plasma iron levels became elevated following infection in hepcidin-deficient mice. Our use of an infectious agent, rather than sterile inflammatory stimuli used in other reports, may account for this difference. We speculate that this increase in plasma iron levels reflects decreases in iron consumption by cytokine-suppressed erythropoiesis (24) and release of iron from intracellular sources damaged by infection, such as the liver (42). We thus propose a model in which infections result in a tendency toward increased extracellular iron that benefits the pathogen, and the protective role of inflammation-mediated hepcidin induction is to counteract this phenomenon. Furthermore, while hepcidin can be produced by leukocytes, airway epithelial cells and other tissues (19–21), our results indicate that hepcidin is predominantly produced by the liver and circulates systemically in the context of bacterial pneumonia. Consistent with this, we observed that hepcidin induction was dependent on IL-6, similar to observations with other systemic inflammatory stimuli (43). In addition, we found that IL-6 neutralization disrupted host defense against K. pneumoniae (Figure 3A), in corroboration with previous reports (22). While IL-6 induces many acute phase reactants, such as C reactive protein, which are crucial to the innate immune response, we speculate that hepcidin induction may be a component of IL-6–mediated protection.

These data provide the host counterpart to the existing literature on the role of the bacterial siderophore system in the virulence of Gram-negative pathogens (30, 44). The hypervirulent strain of K. pneumoniae used in this study produces yersiniabactin, which supports bacterial growth in the respiratory tract, and glycosylated enterobactin, which promotes bacterial growth in the bloodstream by evading lipocalin-2, a host-derived molecule that sequesters nonglycosylated enterobactin and limits microbial iron uptake (45–49). Our findings indicate that even bacteria equipped with these high-affinity acquisition systems are susceptible to a host mechanism that limits iron availability in the host environment. As such, we speculate that these findings may be broadly applicable to infections caused by other phylogenetically related aerobic Gram-negative pathogens (50).

This work has several implications for future research. First, our results may have implications for the therapeutic use of hepcidin antagonists, which are currently under development for the treatment of anemia of inflammation. In this context, we posit that relieving the iron-restricted state could predispose some individuals to Gram-negative sepsis. Second, impaired hepcidin production is a feature of several common clinical entities, including alcohol intoxication and chronic liver disease (33–36), illnesses associated with striking susceptibility to Gram-negative bacteremia (51, 52). Our data suggest that impaired hepcidin production may represent an unappreciated mechanism predisposing these hosts to infection. Finally, hepcidin agonists may represent a novel therapeutic strategy in illnesses where hepcidin production is diminished, such as liver disease and hereditary hemochromatosis, and possibly more broadly in acquired iron-overload states.

Methods

Supplemental Methods are available online with this article.

Animals and in vivo procedures. C57BL/6 mice, purchased from the Jackson Laboratories. Hepc1−/− mice generated on a C57BL/6 background (26) — acquired from the University of California, Los Angeles, California, USA — were bred at the University of Virginia under specific-pathogen–free conditions for a minimum of 2 generations, and offspring were used in the experiments. To minimize any effect of differing microbiota between groups of animals, all animals were bred and maintained in the same room in the vivarium and used bedding from cages of breeders, and animals to be used in experiments were intermixed at weekly intervals. Age- and sex-matched male and female 6- to 12-week-old mice were used in experiments.
Standard rodent chow has artifactually high iron content that causes near-maximal expression of hepcidin at baseline in WT animals (43). To correct this, all animals were conditioned on a low-iron diet (2–4 ppm iron; Teklad diet TD.80396, Envigo) for 5–7 days prior to, and throughout the duration of, all experiments. In our pilot studies and published reports (43, 53), this protocol normalizes baseline hepcidin expression without inducing iron deficiency.

Experimental bacterial pneumonia was induced as previously described (15, 16), by intratracheal inoculation of 500–1,500 CFU of *K. pneumoniae* strain 43816 (American Type Culture Collection). In some experiments, 1,500 CFU of *K. pneumoniae* was delivered i.v. in 100 μl PBS. To induce transient hyperferremia without iron overload, 900 μg of ferric ammonium citrate in 100 μl PBS was delivered i.p., using a previously characterized protocol (54, 55). In some experiments, mice were given 1 mg anti–IL-6 antibody or isotype control i.p. 3 hours prior to and 24 hours following infection (clones MP5-20F3 and HRPN1, BioXcell). In other experiments, 100 nmol of the synthetic hepcidin analogue PR73 (manufactured in house), referred to as minihepcidin in text, or vehicle was administered via i.p. injection beginning 5 hours prior to infection and then every 24 hours until day 5 after infection. PR73 was synthesized using Fmoc solid-phase peptide synthesis and was solubilized for in vivo use with SL220 (a polyethylene glycosylated-phospholipid based solubilizer; NOF America Corporation) as previously described (11, 37, 38).

**Tissue harvest and determination of bacterial burden.** Animals were euthanized with an overdose of a ketamine/xylazine solution, blood was collected from the right ventricle into heparinized syringes, and pulmonary vasculature was perfused with 2 ml of 2 mM EDTA solubilized in PBS. The median liver lobe was collected for RNA analysis. Lungs and liver were homogenized in 1 ml PBS. BAL was performed by serially inflating the lungs via an intratracheal catheter with 1 ml of 2 mM EDTA solubilized in PBS and recovering the fluid 4 times (56). For bacterial enumeration, lung homogenates and blood were serially diluted in sterile water and cultured on blood agar plates, as previously described (57).

**RNA, protein, and iron analysis.** Tissue RNA was extracted using TRIzol and chloroform (Thermo-Fisher Scientific). Buffy coat RNA was extracted using a commercial kit (Qiagen). A commercial kit was used to synthesize cDNA (Promega). Hepcidin transcript was quantified by quantitative PCR in duplicate using SYBR green (Bioline). Expression was calculated using the ΔΔC(t) method normalized to glyceraldehyde 3-phosphate dehydrogenase, and then relative to uninfected animals. Primers were custom synthesized (Sigma Aldrich) as previously published (35): hepcidin 5′ TGCAGAAGAGAAGGAGAGAGACA 3′ (forward) and 5′ CACACTGGGAATTGTTACAGCATT 3′ (reverse); glyceraldehyde 3-phosphate dehydrogenase 5′ TTCACCCACCATGGGAAGGCG 3′ (forward) and 5′ GGCAAGCTCTGTGCTCATGA 3′ (reverse) (58). The thermal cycling conditions were as follows: 2 minutes at 95°C, 10 minutes at 95°C, 5 seconds at 95°C, and 10 seconds at 55°C for 40 cycles. Reactions were performed on a BioRad iQ5 Cycler. Protein concentration of hepcidin was quantified using a commercial ELISA kit (Intrinsic Life Sciences) according to the manufacturer instructions. Plasma iron was quantified with a commercial kit (Sekisui Diagnostics) according to manufacturer instructions with minor modifications — namely, samples were scaled to volume and centrifuged at 10,000 g for 10 minutes between steps to clear insoluble material. Transferrin saturation was determined by measuring the unsaturated iron binding capacity (UIBC) of each sample using a commercial kit (Sekisui Diagnostics). UBBC values were added to plasma iron values in each sample to calculate the total iron binding capacity (TIBC). Transferrin saturation was calculated as (plasma iron/TIBC) × 100%.

**Tissue heme and nonheme iron levels were quantified using the ferrozine assay, as previously described (59–61).** Whole lungs and left lateral lobe of the liver were homogenized in 1 ml of water. BAL was collected into 1 ml of 2 mM EDTA solubilized in PBS. Briefly, 100 μl sample were heated in 100 μl buffer containing 1 N hydrochloric acid and 10% trichloroacetic acid with (total iron) or without (nonheme iron) 2.25% KMnO4 at 60°C for 2 hours. Samples were centrifuged at 16,000 g for 10 minutes to remove insoluble material and were incubated with 50 μl iron detection reagent (6.5 mM ferrozine, 6.5 mM neucoproine, 1 M L-ascorbic acid, 2.5 M ammonium acetate) for 10 minutes and then plated in duplicate and read at A590 nm on a Dynex Triad microplate reader (Dynex Technologies). Heme-iron values were calculated by subtracting nonheme iron values from total iron content. Labile plasma iron (LPI), the major component of non–transferrin-bound iron, was measured using a published protocol with minor modifications (62): briefly, HEPES buffered saline was iron-depleted by incubation with Chelex-100 chelating resin (Bio-Rad) for 1 hour, filtered, and supplemented with 60 nM deferoxamine (Sigma-Aldrich). For LPI measurements, 10 μl of plasma samples were mixed with either 140 μl of HBS/50 μM dihydrorhodamine/40 μM ascorbic acid/0.5 mM nitroacetic acid (−DFP) or 140 μl of the same buffer supplemented with 100 μM deferrirhodamine (+DFP).
Changes in fluorescence were measured over 40 minutes, and the slope was calculated for each sample. The difference between –DFP and +DFP was used as a measure of LPI. The experiment was repeated a total of 3 times using different plasma donors with similar results.

**Flow cytometry.** Cell suspensions from blood and BAL fluid were prepared and analyzed by flow cytometry as previously described, with minor modifications (16, 63, 64). The following antibodies were used for surface antigen staining: anti–MHC-II APC (clone M5/114.15.2), anti–CD11b Pe-Cy7 (clone M170), and anti–CD115 APC (clone AF598) from eBioscience; anti–CD11c PerCP (clone HL3) and anti–CD64 brilliant violet 421 (clone XS5-5/7.1) from BioLegend; and anti–CD16/CD32 (clone 2.4G2), anti–Ly6G PE (clone 1A8), and anti–CD45 AmCyan (clone 30-F11) from BD Biosciences. After extracellular staining, cells were washed in 1% FCS in PBS once and resuspended in the same buffer. Data were acquired on a FACS Canto II instrument using BD FACS Diva software (version 8.0; BD Biosciences) and analyzed with FlowJo software (version 8.8.6; Tree Star Inc.). The absolute cell number was calculated as the product of the cell type frequency (as determined by gating analysis) and the total number of cells in the sample (as determined by manually counting under a hemocytometer).

**In vitro experiments.** BM neutrophils were isolated by grinding the tibia, femur, pelvis, and spine from uninfected mice in a sterile mortar and pestle as previously described (15, 57), followed by positive selection using anti-Ly6G magnetic beads, according to the manufacturer’s instructions (MACS; Miltenyi Biotec). Isolated populations were found to have > 90% viability by examination in trypan blue. Neutrophil bacterial killing was quantified using a published protocol with minor modifications (57). Briefly, *K. pneumoniae* was opsonized at a concentration of 8 × 10^6 CFU/ml in 20% fresh mouse serum and HBSS for 30 minutes at 37°C, washed once in cold water, once in HBSS, and resuspended in HBSS. Aliquots of 2 × 10^5 bacteria were added to sterile U-bottom 96-well plates and coincubated with 5 × 10^3, 2 × 10^3, 2 × 10^2, 2 × 10^1, or no neutrophils from WT or hepcidin-deficient mice in triplicate in a total volume of 200 μl HBSS. Samples were incubated in a humidified chamber with 5% CO_2 at 37°C for 1 hour. Number of surviving bacteria was determined by hypotonic lysis of neutrophils in cold water and serial dilution and culture.

Human plasma was inactivated (30 min at 56°C) and incubated overnight at 37°C with 0–60 μM ferric ammonium citrate (Sigma-Aldrich), followed by quantification of transferrin saturation and LPI. The samples were then used to generate 90% plasma agar solid media, poured in sterile 12-well tissue culture plates. *K. pneumoniae* was suspended at 1 × 10^4 CFU/ml and 5 μl plated in triplicates on plasma agar wells and incubated for 24 hours at 37°C; bacterial colonies were counted and photographed.

**Statistics.** Data were analyzed using Prism software (version 6.0, GraphPad Software). The Mantel-Cox test was used to analyze survival data. A one-way ANOVA with a post-test for linear trend was used to analyze change in a single parameter over time, and one-way ANOVA with a Tukey multiple comparison post-test was used to analyze differences between 3 or more groups. A 2-way ANOVA with Tukey multiple comparison post-test was used to compare changes over time between 2 groups. An unpaired, 2-tailed t test was used to compare 2 groups for RNA, protein, and iron levels. A Mann-Whitney U test was used to compare bacterial burden between groups. P <0.05 was considered statistically significant.

**Study approval.** Animal experiments were performed in compliance with NIH Guide for the Care and Use of Laboratory Animals, The US Animal Welfare Act, and PHS Policy on Humane Care and Use of Laboratory Animals and was approved by the Institutional Animals Care and Use Committee at the University of Virginia (protocol 3571).

**Author contributions**

KRM, EN, TG, and BM conceived the project, designed experiments, and interpreted data; KRM, ZZ, AMB, REC, DS, and MDB performed experiments; KRM and DS generated the figures; SV, EN, and TG designed and contributed materials; and KRM and BM wrote the first draft of the manuscript. All authors revised the manuscript.

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Address correspondence to: Borna Mehrad, P.O. Box 800546, Charlottesville, Virginia 22908, USA. Phone: 434.243.4845; E-mail: mehrad@virginia.edu.


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42. Kell DB, Pretorius E. Serum ferritin is an important inflammatory disease marker, as it is mainly a leakage product from damaged cells. *Metallomics*. 2014;6(4):748–747.