Genomic Analyses Identify Manganese Homeostasis as a Driver of Group B Streptococcal Vaginal Colonization

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ABSTRACT

Group B Streptococcus (GBS) is associated with severe infections in utero and in newborn populations, including pneumonia, sepsis, and meningitis. GBS vaginal colonization of the pregnant mother is an important prerequisite for transmission to the newborn and the development of neonatal invasive disease; however, our understanding of the factors required for GBS persistence and ascension in the female reproductive tract (FRT) remains limited. Here, we utilized a GBS mariner transposon (Krmit) mutant library previously developed by our group and identified underrepresented mutations in 535 genes that contribute to survival within the vaginal lumen and colonization of vaginal, cervical, and uterine tissues. From these mutants, we identified 47 genes that were underrepresented in all samples collected, including mtsA, a component of the mtsABC locus, encoding a putative manganese (Mn2+) -dependent ATP-binding cassette transporter. RNA sequencing analysis of GBS recovered from the vaginal tract also revealed a robust increase of mtsA expression during vaginal colonization. We engineered an ΔmtsA mutant strain and found by using inductively coupled plasma mass spectrometry that it exhibited decreased concentrations of intracellular Mn2+, confirming its involvement in Mn2+ acquisition. The ΔmtsA mutant was significantly more susceptible to the metal chelator calprotectin and to oxidative stressors, including both H2O2 and paraquat, than wild-type (WT) GBS. We further observed that the ΔmtsA mutant strain exhibited a significant fitness defect in comparison to WT GBS in vivo by using a murine model of vaginal colonization. Taken together, these data suggest that Mn2+ homeostasis is an important process contributing to GBS survival in the FRT.

IMPORTANCE

Morbidity and mortality associated with GBS begin with colonization of the female reproductive tract (FRT). To date, our understanding of the factors required for GBS persistence in this environment remain limited. We identified several necessary systems for initial colonization of the vaginal lumen and penetration into the reproductive tissues via transposon mutagenesis sequencing. We determined that mutations in mtsA, the gene encoding a protein putatively involved in manganese (Mn2+) transport, were significantly underrepresented in all in vivo samples collected. We also show that mtsA contributes to Mn2+ acquisition and GBS survival during metal limitation by calprotectin, a metal-chelating protein complex. We further demonstrate that a mutant lacking mtsA is hypersusceptible to oxidative stress induced by both H2O2 and paraquat and has a severe fitness defect compared to WT GBS in the murine vaginal tract. This work reveals the importance of Mn2+ homeostasis at the host-pathogen interface in the FRT.

KEYWORDS

Group B Streptococcus, intrauterine infection, manganese, vaginal colonization

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Infant morbidity and mortality resulting from adverse pregnancy outcomes remain a significant health burden, even in developed countries (1, 2), and one of the leading causes of adverse pregnancy outcomes is intrauterine infection (3). *Streptococcus agalactiae*, also known as group B *Streptococcus* (GBS), is a Gram-positive bacterium that asymptptomatically colonizes the lower gastrointestinal tract and vaginal tract of 25 to 30% of healthy women. However, during pregnancy, GBS poses a risk to the mother and the developing fetus, as it has potential to cause severe infections (4). In pregnant women, the presence of GBS in the vaginal tract, particularly at high burdens, can result in ascending spread through the reproductive tract, causing intrauterine infection (5, 6). This has been shown to contribute to miscarriage, stillbirth, and preterm premature rupture of membranes (PPROM) (7–9), with an estimated 3.5 million preterm births due to GBS infection per year (7).

In addition to the impact of GBS vaginal colonization on maternal and fetal morbidity and mortality, GBS remains a primary threat to newborn babies. Approximately 50 to 70% of vaginally delivered infants of GBS-colonized mothers will become colonized, and 1 to 2% of colonized infants develop GBS invasive diseases, including pneumonia, sepsis, and meningitis (10, 11). These cases result from ascending infection of the bacterium through the placental membranes to initiate infection in utero, or alternatively, GBS may be acquired during passage through the birth canal by aspiration of infected vaginal fluids (9). During intrauterine GBS infection, localized inflammatory reactions allow bacteria to invade either the blood vessels of the fetus or the amniotic fluid, where the bacteria can gain entry to the fetus by aspiration or direct invasion (3, 12). In rare cases, GBS can lead to maternal sepsis (13, 14). To combat the risk of GBS infection in newborns, many countries have implemented universal or risk-based screening and the use of prophylactic antibiotics administered to colonized pregnant mothers at the time of delivery (15). But before late gestational GBS screening occurs, the pregnant mother and the fetus are susceptible to GBS colonization and ascension to the higher reproductive tissues and penetration of the placental membranes, through largely unknown mechanisms.

Transposon sequencing (Tn-seq) provides a high-throughput method to determine how individual genes contribute to bacterial fitness by combining untargeted mutagenesis with next-generation sequencing (16). Recent advances in the field of streptococcal pathogenesis have utilized transposon sequencing to determine gene fitness and essentiality in several pathogens both in vitro and in vivo (17–25). Previous work from our laboratory performed the first transposon sequencing experiment in the presence of calprotectin, a metal-chelating protein complex that is abundant in the neutrophilic cytosol, to identify some of the genes involved in the global response to metal availability in vitro (26). In this study, we sought to use our saturated transposon mutant library in a murine model of GBS vaginal colonization to identify the systems important for GBS fitness in the female reproductive tract (FRT). We identified 535 mutants that were underrepresented in comparison to the input library and important for colonization of the vaginal lumen or invasion into the higher reproductive tissues. Assignment of clusters of orthologous groups (COGs) revealed a majority of these underrepresented systems to contribute to nutrient transport or transcriptional regulation. We further identified that *mtsA*, the gene encoding a putative Mn²⁺-binding protein, was one of 47 genes that were underrepresented in all sites within the reproductive tract and contributes to Mn²⁺ import, survival during calprotectin-mediated metal chelation, resistance to oxidative stress, and GBS vaginal colonization. Collectively, these data support the conclusion that Mn²⁺ homeostasis is important for evasion of the host response and fitness in the FRT.

**RESULTS**

**Genome-wide analysis of GBS factors involved in colonization and ascending infection.** To determine the systems important for GBS fitness in the FRT, we utilized our previously established murine model of vaginal colonization (4, 27). Triplicate groups of mice (n = 3/group) were synchronized with 17β-estradiol, and mice were challenged intra-vaginally with triplicate mid-logarithmic cultures of 10⁷ CFU of saturated *Kmr* transposon
On subsequent days, vaginal swabs were collected, and on day 3 post-colonization, reproductive tissues were harvested. All samples were plated on selective medium to assess colonization and to recover transposon mutants for sequencing analysis (Fig. 1). Briefly, recovered transposon mutants from luminal swabs collected on days 1 and 3 and vaginal, cervical, and uterine tissues harvested day 3 post-colonization were pooled, and genomic DNA was isolated. Genomic DNA samples were sheared and enriched for Kmit transposon insertions by PCR, and Illumina adaptors were added (see Fig. S1A in the supplemental material). Samples were sequenced as previously described (26), reads were mapped to the Streptococcus agalactiae CJB111 genome (28), and differential analyses were performed using Tn-seq TRANSIT and DESeq2 by comparing in vivo samples to the input library (26, 29).

We detected a total of 535 underrepresented mutants with a significant defect in colonization of the vaginal lumen and/or higher reproductive tissues (Fig. 1B to G; Table S1). We identified underrepresented mutations in 250 genes in luminal samples collected day 1 post-colonization (Fig. 1B) and 192 genes in day 3 post-colonization samples (Fig. 1C). We further identified mutations in the samples collected from the reproductive tissues, with underrepresented mutations occurring in 277 genes in vaginal tissues, 237 genes in cervical tissues, and 182 genes in uterine tissues (Fig. 1D to F). Of the Tn mutants that were identified as underrepresented in the FRT, clusters of orthologous groupings (COGs) were identified for 145 and 117 genes in the luminal samples from days 1 and 3 post-colonization, respectively, and for 158, 149, and 101 genes in the vaginal, cervical, and uterine tissues (Fig. 1G and H).
tissues, respectively (Fig. 1H; Fig. S2). The most highly represented COGs of known function for the luminal samples were those pertaining to carbohydrate and amino acid transport and metabolism and to replication, recombination, and repair, while the vaginal, cervical, and uterine tissues had the highest abundance for COGs involved in transcription, cell wall biogenesis, and inorganic ion transport and metabolism (Fig. 1H). We also observed overrepresented mutations, including 53 genes that were overrepresented for transposon insertion mutations in all samples collected (Fig. S1B), with COGs showing contribution of these genes to nutrient transport, transcription, and translation (Fig. S1C).

From these analyses, we observed underrepresented mutations in several systems that are involved or hypothesized to contribute to GBS colonization or virulence potential (Table 1), including pili, the newly characterized type VII secretion system (30), hydrolases, proteases, and bacteriocin immunity proteins. We also identified a large number of underrepresented mutations in systems involved in sensing and responding to environmental cues, including transcriptional regulators, two-component systems, phosphotransferase (PTS) systems, ATP-binding cassette transporters (ABC transporters), and metal homeostasis systems (Table 1). To understand the factors required specifically for GBS colonization of the vaginal lumen or the uterus, we further categorized the genes by those that were significantly underrepresented in the vaginal lumen and in all subsequent tissues (Fig. 2A, shown in blue) and those that were uniquely underrepresented in the uterine tissues (Fig. 2A, shown in red). We identified a unique set of 47 genes with underrepresented mutations detected in all samples collected (Table S2), indicating that GBS luminal persistence is an important prerequisite for invasion into the reproductive tissues. An additional 47 genes were uniquely found to contribute to penetrating the cervicovaginal barrier and accessing the uterus (Table S3). Similar COG classifications were found to be important for colonization of both the vaginal lumen and the uterus, including amino acid and carbohydrate metabolism and inorganic ion transport and metabolism (Fig. 2B and C). We identified unique COGs for each set of samples, with lipid transport and metabolism unique to colonization of all the FRT (Fig. 2B) and cell wall biogenesis, energy production and conversion, and signal transduction mechanisms unique to uterine colonization (Fig. 2C). From these subsets of data, we identified systems involved in the maintenance of copper, iron, Mn\(^{2+}\), zinc, and magnesium homeostasis as differentially important for colonization of specific sites in the FRT (Fig. 2D). One of the genes that was significantly underrepresented in the lumen and all tissues was mtsA (Fig. 2D). mtsA encodes the substrate-binding protein of a putative Mn\(^{2+}\)/Fe-dependent ATP-binding cassette transporter, MtsABC. A previous study performed RNA sequencing of GBS strain A909 recovered from murine vaginal lavage fluid on day 2 post-colonization and found that mtsA was upregulated in the vaginal lumen (31). We performed a similar experiment using the GBS CJB111 strain and sequenced bacteria recovered from vaginal swab samples collected days 1 and 3 post-colonization and also observed a robust induction of mtsA at both time points (Fig. S3; Table S4). We provide the complete data set of our RNA sequencing experiment with CJB111 as an additional community resource (Table S4).

**Role of mtsA in Mn\(^{2+}\) homeostasis.** To confirm the importance of mtsA to Mn\(^{2+}\) homeostasis, we constructed a ΔmtsA mutant strain as described in Materials and Methods. We first performed inductively coupled plasma mass spectrometry (ICP-MS) on bacterial pellets grown in Todd-Hewitt broth (THB) medium as described in Materials and Methods. The ΔmtsA mutant had significantly lower intracellular concentrations of Mn\(^{2+}\) than the wild-type (WT) GBS control, with a mean of 821.9 μg/mL and a standard error of the mean (SEM) of ±32.8 μg/mL in the WT strain and a mean of 699.7 μg/mL and a SEM of ±10.88 μg/mL in the ΔmtsA mutant strain, with all strains normalized by cell dry weight (mg) (Fig. 3A). The levels of intracellular Mn\(^{2+}\) were restored in the complemented (ΔmtsA+) strain compared to the WT GBS control (Fig. S4A). These data support the hypothesis of the involvement of mtsA in Mn\(^{2+}\) transport. Previous work from our laboratory demonstrated that calprotectin, a protein complex produced by the host, inhibited GBS growth by limiting nutrient metal ions (26). Calprotectin is known to possess two metal-binding sites, and previous studies have described site 1 to form a unique His6 site that serves as a high-affinity binding site for Mn\(^{2+}\) ions (32–34). We next hypothesized that expression of genes involved in Mn\(^{2+}\)
| Functional category and gene no. (CJB111 genome) | Gene name | Description | Fold change in reproductive tract |
|-----------------------------------------------|-----------|-------------|----------------------------------|
| Transcriptional regulators                    |           |             | D1  | D3  | VG  | CX  | UT  |
| ID870_00525                                  | CtsR family transcriptional regulator |             | −9.94 | −11.33 | −25.05 |
| ID870_00635                                  | LacI family DNA-binding transcriptional regulator |             | −5.10 |         |       |
| ID870_00850                                  | MerR family transcriptional regulator |             | −8.19 |         |       |
| ID870_01280                                  | LacI family DNA-binding transcriptional regulator |             | −5.36 |         |       |
| ID870_02045                                  | LysR family transcriptional regulator |             | −5.52 |         |       |
| ID870_02800                                  | LysR family transcriptional regulator |             | −9.51 | −7.83  | −8.05 |
| ID870_02815                                  | Transcriptional regulator |             | −4.35 |         |       |
| ID870_03090                                  | TetR/AcrR family transcriptional regulator |             | −8.59 |         |       |
| ID870_03570                                  | Helix-turn-helix transcriptional regulator |             | −6.92 | −6.59  | −9.99 |
| ID870_04400                                  | Spx/MgsR family RNA polymerase-binding regulatory protein |             | −6.29 | −6.05  | −4.13 |
| ID870_05775                                  | FadR family transcriptional regulator |             | −6.22 | −8.42  | −10.78 |
| ID870_06770                                  | ArgR family transcriptional regulator |             | −8.12 | −12.14 | −18.06 |
| ID870_07170                                  | AraC family transcriptional regulator |             | −4.38 | −9.28  | −7.60 |
| ID870_07370                                  | Peroxide-responsive transcriptional repressor |             | −5.04 |         |       |
| ID870_07715                                  | Sugar-binding transcriptional regulator |             | −3.28 | −2.35  |       |
| ID870_08260                                  | Helix-turn-helix transcriptional regulator |             | −4.81 |         |       |
| ID870_09330                                  | MurR/RpiR family transcriptional regulator |             | −8.81 |         |       |
| ID870_10200                                  | ArgR family transcriptional regulator |             | −8.69 |         |       |
| ID870_10525                                  | PadR family transcriptional regulator |             | −7.00 |         |       |
| Two-component systems                        |           |             | D1  | D3  | VG  | CX  | UT  |
| ID870_00210                                  | fspR      | Response regulator (TCS-16) | −4.78 | −4.48 | −7.15 |
| ID870_00330                                  | Response regulator (TCS-15) |             | −6.19 |         |       |
| ID870_03010                                  | HAMP domain-containing histidine kinase (TCS-12) |         | −2.79 |         |       |
| ID870_04300                                  | ltrR      | LytTR family transcriptional regulator |             | −5.35 | −7.56 |
| ID870_05675                                  | yycF      | Response regulator transcription factor (TCS-8) |             | −3.92 |         |       |
| ID870_07355                                  | haeS      | HAMP domain-containing histidine kinase |             | −2.66 | −6.47  | −7.61 |
| ID870_07360                                  | saeR      | Response regulator transcription factor |             | −5.89 |         |       |
| ID870_08500                                  | lytS      | Sensor protein |             | −3.68 |         |       |
| ID870_10420                                  | HAMP domain-containing histidine kinase (TCS-19) |             | −5.24 |         |       |
| Metal systems                                |           |             | D1  | D3  | VG  | CX  | UT  |
| ID870_00250                                  | adcAII    | Zinc ABC transporter substrate-binding protein |             | −5.66 |         |       |
| ID870_02005                                  | mtsR      | Metal-dependent transcriptional regulator |             | −4.89 |         |       |
| ID870_02010                                  | mtsA      | Metal ABC transporter substrate-binding protein |             | −4.95 | −6.71  | −10.29 |
| ID870_02080                                  | nikC      | ABC transporter permease |             | −7.11 | −10.16 | −7.95 |
| ID870_02090                                  | nikD      | ABC transporter ATP-binding protein |             | −6.23 | −7.75  | −5.92 |
| ID870_02665                                  | fhuB      | Iron ABC transporter permease |             | −2.89 | −6.31  | −9.13 |
| ID870_07025                                  | Magnesium transporter CorA family protein |             | −4.98 |         |       |
| ID870_07140                                  | sczA      | MerR family transcriptional regulator |             | −11.26 | −23.68 |       |
| ID870_07395                                  | copZ      | Heavy-metal-associated domain-containing protein |             | −9.54 | −16.55 | −18.27 |

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| Functional category and gene no. (CJB111 genome) | Gene name                      | Description                                      | Fold change in reproductive tract |
|-------------------------------------------------|--------------------------------|--------------------------------------------------|-----------------------------------|
|                                                 |                                |                                                  | D1      | D3      | VG      | CX      | UT      |
| Transport systems                                |                                |                                                  |         |         |         |         |         |
| Phosphotransferase systems                       |                                |                                                  |         |         |         |         |         |
| ID870_00140                                     | PTS transporter subunit EIIC   |                                                  | -4.92   |         |         |         |         |
| ID870_00190                                     | PTS sugar transporter subunit II B |                                              | -9.85   | -6.35   | -7.73   | -8.63   | -5.88   |
| ID870_00265                                     | PTS sugar transporter subunit II A |                                               |         |         |         |         |         |
| ID870_00430                                     | PTS system mannose/fructose/N-acetylgalactosamine-transporter subunit II B |                               | -7.30   |         |         |         |         |
| ID870_00990                                     | PTS fructose transporter subunit II B |                                         | -9.43   | -6.94   | -19.55  | -21.30  |
| ID870_00995                                     | PTS sugar transporter subunit II A |                                          | -7.71   | -6.60   |         |         |         |
| ID870_01290                                     | PTS glucose transporter subunit II A |                                         | -3.29   | -3.53   | -3.21   |         |         |
| ID870_01665                                     | Transporter substrate-binding domain-containing protein |                                      | -6.37   | -6.37   | -5.75   | -4.38   |         |
| ID870_07535                                     | PTS mannose/fructose/sorbose transporter family subunit IID |                                 | -3.95   | -6.86   | -8.25   | -10.86  | -30.53  |
| ATP-binding cassette transporters               |                                |                                                  |         |         |         |         |         |
| ID870_01555                                     | ECF-type riboflavin transporter substrate-binding protein |                               | -3.90   |         |         |         |         |
| ID870_01670                                     | Amino acid ABC transporter permease |                                             | -8.20   | -9.11   | -9.11   |         |         |
| ID870_01820                                     | ABC transporter ATP-binding protein |                                         | -2.92   | -3.35   |         |         |         |
| ID870_02315                                     | Amino acid ABC transporter ATP-binding protein |                              | -5.60   |         |         |         | -7.11   |
| ID870_02408                                     | Transporter substrate-binding domain-containing protein |                                 | -3.74   | -7.05   | -10.51  | -11.00  | -12.24  |
| ID870_02675                                     | ABC transporter ATP-binding protein |                                         | -2.53   | -5.25   | -8.06   | -5.70   | -4.94   |
| ID870_02990                                     | ABC transporter ATP-binding protein |                                         | -3.51   | -5.86   | -3.42   |         |         |
| ID870_04360                                     | ABC transporter ATP-binding protein |                                         | -4.48   |         |         |         |         |
| ID870_04650                                     | ABC transporter ATP-binding protein |                                         | -5.64   | -7.43   | -4.99   |         |         |
| ID870_04655                                     | ABC transporter ATP-binding protein |                                         | -4.63   |         |         |         |         |
| ID870_05620                                     | ABC transporter ATP-binding protein |                                         | -5.38   | -6.01   | -8.57   |         |         |
| ID870_05695                                     | Amino acid ABC transporter permease |                                         | -3.81   |         |         |         |         |
| ID870_05935                                     | ABC transporter ATP-binding protein |                                         | -7.10   |         |         |         |         |
| ID870_05965                                     | ABC transporter permease |                                                   | -2.69   | -10.28  | -9.16   | -7.53   |         |
| ID870_06520                                     | ABC transporter permease |                                                   | -3.53   |         |         |         |         |
| ID870_06635                                     | ECF transporter S component |                                                   | -2.76   | -3.63   |         |         |         |
| ID870_07335                                     | bioY                           | Biotin transporter |                                    |         |         |         |         | -8.85   |
|                                                 |                                |                                                  |         |         |         |         |         |
| Virulence potential                              |                                |                                                  |         |         |         |         |         |
| ID870_01235                                     | comE                           | Competence protein |                                    | -8.79   |         | -7.62   |         |         |
| ID870_02600                                     | piliB                          | PI-2a pilus major subunit |                 | -3.57   | -3.52   |         |         |         |
| ID870_02605                                     | Class C sortase |                                              | -6.62   |         | -4.78   |         |         |

(Continued on next page)
| Functional category and gene no. (CJB111 genome) | Gene name      | Description                                      | Fold change in reproductive tract |
|-------------------------------------------------|----------------|--------------------------------------------------|-----------------------------------|
| ID870_02615                                     | pilC           | Pil-2a pilus subunit                             | D1: 4.25                         |
|                                                 |                |                                                  | D3: -8.42                        |
|                                                 |                |                                                  | VG: -13.29                       |
|                                                 |                |                                                  | CX: -10.63                       |
|                                                 |                |                                                  | UT: -9.08                        |
| ID870_03655                                     | lrgB           | LrgB family protein                              | D1: -3.25                        |
|                                                 |                |                                                  | D3: -4.69                        |
|                                                 |                |                                                  | VG: -9.12                        |
|                                                 |                |                                                  | CX: -8.51                        |
|                                                 |                |                                                  | UT: -7.60                        |
| ID870_04185                                     | essA           | Type VII secretion protein                       | D1: -7.42                        |
|                                                 |                |                                                  | D3: -10.17                       |
|                                                 |                |                                                  | VG: -9.95                        |
|                                                 |                |                                                  | CX: -8.09                        |
| ID870_04625                                     | srtC1          | Class C sortase                                  | D1: -6.09                        |
|                                                 |                |                                                  | D3: -9.97                        |
|                                                 |                |                                                  | VG: -6.00                        |
|                                                 |                |                                                  | CX: -5.60                        |
| ID870_05305                                     | serine protease|                                                   | D1: -10.70                       |
|                                                 |                |                                                  | D3: -10.35                       |
|                                                 |                |                                                  | VG: -9.43                        |
| ID870_06020                                     |                  |                                                  |                                  |
| ID870_07305                                     |                  |                                                  |                                  |
| ID870_07950                                     |                  |                                                  |                                  |
| ID870_08585                                     |                  |                                                  |                                  |
| ID870_09920                                     |                  |                                                  |                                  |
| ID870_09980                                     |                  |                                                  |                                  |
| ID870_10015                                     |                  |                                                  |                                  |
| ID870_10190                                     |                  |                                                  |                                  |
| ID870_10550                                     |                  |                                                  |                                  |
| ID870_10565                                     |                  |                                                  |                                  |

aD1, vaginal lumen on day 1 postcolonization; D3, vaginal lumen on day 3 postcolonization; VG, tissue from the vagina; CX, tissue from the cervix; UT, tissue from the uterus.
acquisition would increase following exposure to calprotectin. We therefore assessed expression of \textit{mtsA} by quantitative reverse transcriptase PCR (qRT-PCR) analysis following treatment with calprotectin or with calprotectin plus MnCl\textsubscript{2} supplementation. Fold changes in \textit{mtsA} expression were calculated by the \textit{DD}_{CT} method with \textit{gyrA} serving as an internal control and were compared to expression observed in untreated controls. Following exposure to calprotectin, we observed a significant induction of \textit{mtsA} expression, nearly 28-fold upregulation, while supplementation with MnCl\textsubscript{2} following exposure to calprotectin restored expression to control levels, at nearly 4-fold downregulation compared to that of untreated controls (Fig. 3B). Having confirmed the importance of \textit{mtsA} to Mn\textsuperscript{2+} import, we then hypothesized that the \textit{DmtsA} mutant strain would exhibit increased sensitivity to chelation by calprotectin. We assessed growth (as measured by optical density at 600 nm [OD\textsubscript{600}]) of GBS WT and \textit{DmtsA} mutant strains in the presence of 720 and 960 mg/mL calprotectin with or without the addition of 10 \mu M MnCl\textsubscript{2} (Fig. 3C and D). Similar to what was previously described by our laboratory (26), we observed a significant inhibition of growth in WT CJB111 grown in the presence of calprotectin (Fig. 3C), and further, we identified a significant increase in susceptibility to calprotectin-mediated metal chelation in the \textit{DmtsA} mutant strain compared to that of WT GBS (Fig. 3C). Notably, we found that supplementing all samples with 10 \mu M MnCl\textsubscript{2} rescued growth of WT GBS grown in the presence of calprotectin, while growth was still impaired in the GBS \textit{DmtsA} mutant regardless of MnCl\textsubscript{2} supplementation (Fig. 3D).

\textbf{Mn\textsuperscript{2+} transport as an important driver of GBS vaginal colonization.} The findings reported thus far led us to hypothesize that MtsA would contribute to GBS colonization and persistence within the FRT. Using our previously described murine model of vaginal colonization, we colonized female CD-1 mice intravaginally with both the WT CJB111 (serotype V) GBS and the \textit{DmtsA} mutant strain in competition. The \textit{DmtsA} mutant strain exhibited a significant fitness defect compared to WT GBS as early as day 1 post-colonization.
By day 3 post-colonization, the WT strain had outcompeted the ΔmtsA mutant and all 19 mice had cleared the ΔmtsA mutant strain from the vaginal lumen, while 14/19 mice were still colonized with WT CJB111 at the experimental endpoint (Fig. 4A). To assess GBS burden in the higher reproductive tissues, we harvested tissues on day 6 post-colonization and enumerated GBS CFU. We observed a complete clearance of the ΔmtsA strain from the vagina, cervix, and uterus, although WT GBS was detected at a high burden in all tissues (Fig. 4B to D). We similarly found that the WT outcompeted the ΔmtsA mutant strain in vitro when grown in coculture (Fig. S4B). All vaginal colonization experiments were repeated in a second GBS genetic background, and we observed similar phenotypes in the serotype Ia WT A909 and A909 ΔmtsA strains (Fig. 4E to H). Consistent with what we observed in the CJB111 background, we detected complete clearance of the ΔmtsA mutant strain from the vaginal lumen at day 4 post-colonization, while WT A909 was recovered from the lumen of 8/19 mice at the time of tissue collection (Fig. 4E to H). We also observed during single challenge that the ΔmtsA mutant strain exhibited a reduced ability to colonize the FRT compared to the WT strain (Fig. S4C to F). To determine if phenotypes observed between the WT and the ΔmtsA mutant were due to the presence of calprotectin in the FRT during GBS colonization, we colonized age-matched WT C57BL/6 and S100A9−/− calprotectin knockout mice with WT and ΔmtsA GBS in competition or in single challenge. We found that the ΔmtsA mutant strain still exhibited a significant decrease in fitness in colonization and ascension of both C57BL/6 WT and S100A9−/− mice compared to the WT
GBS during competition (Fig. 5A to D) or single challenge (Fig. 5E to H). These data confirm that GBS MtsA is important for colonization of the FRT, in both the presence and absence of calprotectin.

**Mn**

homeostasis impacts GBS sensitivity to oxidative stressors. As Mn**2+** homeostasis has been previously shown to contribute to the ability of bacterial pathogens to withstand oxidative stress (33, 35, 36), we hypothesized that the ΔmtsA mutant would exhibit increased sensitivity to stress. To understand the contribution of mtsA to GBS survival during oxidative stress, we assessed the growth of the WT and the ΔmtsA mutant strain following addition of H2O2 and the superoxide stressor paraquat. We observed a significant decrease in growth of the ΔmtsA mutant strain in the presence of both H2O2 (Fig. 6A) and paraquat (Fig. 6B) in comparison to that of WT GBS.

**DISCUSSION**

Vaginal colonization by GBS is an important prerequisite for maternal and fetal infections. Despite the abundance of data showing the contribution of GBS to maternal-fetal morbidity and mortality, our understanding of the factors involved in colonization or ascension in the FRT remain to be fully elucidated. In this study, we sought to utilize a high-throughput transposon mutagenesis screen to identify systems involved in colonization of the lumen and invasion into the reproductive tissues using our previously described mariner-GBS transposon mutant library (26). From this in vivo screen, we identified 525 underrepresented mutants in systems involved in transcriptional regulation, cellular metabolism, and nutrient import and efflux and systems involved in GBS virulence. As mtsA transposon mutants were found to be underrepresented in all samples collected, we further characterized the role of mtsA in GBS survival during calprotectin and oxidative stress and persistence within the vaginal tract. These data...
provide new insight into the importance of metal homeostasis for bacterial colonization and ascension of the FRT.

GBS systems of particular interest that were identified as loss-of-function mutations, or underrepresented mutations, included regulatory systems such as two-component regulatory systems or transcriptional regulators (Table 1). We observed underrepresented mutations in several two-component systems (TCS) that were previously annotated, including TCS-8, -12, -15, -16, and -18 (37). Of these, few have been characterized in GBS and their regulon is unknown, with the exception of TCS-16 (FspRS). FspRS has been shown to be induced by and necessary for GBS growth in the presence of fructose-6-phosphate and is important for persistence within the murine vaginal tract (37).

We also observed underrepresented mutations in SaeRS and LtdR, two recently studied TCS in GBS (31, 38). Expression of SaeRS was previously shown to be induced during GBS vaginal colonization (31), and mutations in genes regulated downstream of SaeRS led to significant decreases in GBS fitness in the vaginal tract (31). Further, LtdR is a response regulator that our laboratory has previously shown to contribute to regulation of virulence (38). Previous studies demonstrated that a ΔltdR strain was cleared more quickly from the murine vaginal tract than WT GBS (38), confirming the underrepresentation we observed in our transposon screen.

In addition to systems involved in gene regulation, we also observed a significant underrepresentation of known and putative systems involved in GBS virulence. We identified underrepresented mutants in the lumen and in vaginal and cervical tissues in pilus components of the pilus island-2a (PI-2a), including the major and minor pilus

FIG 5  GBS Mn²⁺ homeostasis is independent of host calprotectin stress in the FRT. Female C57BL/6 and S100A9−/− mice were colonized with 1 × 10⁷ CFU of GBS CJB111 WT and ΔmtsA strains in competition (A to D) or as single challenge (E to H). Recovered CFU were quantified from vaginal lumen (A and E) or vaginal (B and F), cervical (C and G), or uterine (D and H) tissue homogenates. Significance was determined by two-way ANOVA with Sidák’s multiple-comparison test (A and E) or one-way ANOVA with Sidák’s multiple-comparison test (B to D and F to H). *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001; ns, not significant.
subunits and the pilus 2a cell wall anchoring C-class sortase. GBS strains encode three different pilus islands, PI-1, PI-2a, and PI-2b, with most strains expressing two different types. Studies from our laboratory have previously described the importance of GBS pili in interactions with reproductive epithelial cells and in vaginal colonization in vivo (39, 80). Interestingly, PI-1 was unaltered in our transposon mutagenesis screen, and it has been previously shown that PI-1 does not promote GBS interactions with the reproductive epithelium (40). The type VII secretion system was recently described in GBS and was shown to contribute to invasive disease through EsxA pore formation-induced endothelial cell cytotoxicity (30). Although the GBS type VII secretion system has not yet been shown to contribute to colonization of FRT, we observed underrepresented mutations in \( \text{essA} \), a gene encoding a core machinery component, and in a gene encoding a putative effector WXG100 family protein in samples collected from the lumen and other tissues, likely suggesting an important role in the reproductive tract as well.

In addition to underrepresented mutations, we also detected mutations that are enriched or overrepresented during GBS colonization (see Table S1 in the supplemental material). Transposon sequencing screens provide a high-throughput approach to determine biological fitness in a given environment (16), although this competitive environment does not always correlate with data observed in single mutations. For example, our screen identified \( \text{hylB} \) as a gene with significantly overrepresented mutations (Table S1), suggesting that mutations in \( \text{hylB} \) increase GBS fitness in the female reproductive tract; however, it has been previously shown that hyaluronidase (encoded by \( \text{hylB} \)) is a secreted factor that contributes to vaginal colonization and reproductive tract ascension (41, 42). Another example of an overrepresented mutation found in our data set was \( \text{cylE} \), the gene that encodes the GBS \( \beta \)-hemolysin/cytolysin. Interestingly, \( \text{cylE} \) mutants have previously been shown to exhibit a decreased persistence in a murine model of vaginal colonization (43). Thus, there are limitations to the sensitivity of transposon sequencing, specifically as it is unable to detect the contribution of individual mutations in genes encoding products secreted into the environment (44, 45). Also in this study, rather than declaring gene essentiality, we aimed to focus our analyses on GBS transposon mutant fitness. This could allow for some mutants that are classified herein as underrepresented to have essential roles in vivo, with some examples including DNA polymerase, tRNAs, and genes encoding ribosomal proteins.

Another group of genes that were well represented in our data set were systems involved in the maintenance of metal ion homeostasis. Some of the underrepresented metal-dependent systems of interest were those involved in homeostasis of zinc, copper, a putative nickel system, and Mn\(^{2+}\). Underrepresented mutations in the zinc transport machinery, including \( \text{adcAII} \) and \( \text{adcB} \) (26, 46, 47), were detected only in the cervical and uterine tissue samples, suggesting that the vaginal lumen may in fact be a zinc-replete environment where GBS may
need not rely on these systems for fitness, although it is important to note that we and others identified genes involved in zinc import to be upregulated in the vaginal lumen (31) (Table S4). While previous work from our laboratory has shown that the zinc import machinery is crucial for the development of invasive disease in vivo (26), the contribution of these systems to GBS vaginal colonization has not yet been described. We also identified copZ as underrepresented in all samples collected. CopZ is a gene that encodes a copper-binding chaperone protein contributing to regulation of the copY repressor, and the downstream, copA are collectively involved in copper efflux (48–50). CopZ and the ortholog CupA of Streptococcus pneumoniae have been shown to reduce Cu2+ to Cu1+, which contributes to regulatory signaling, to growth in the presence of copper, and to interactions with host cells (51, 52). Components of the putative nik operon (ID870_02070 to ID870_02090), including nikC and nikE, were also underrepresented in luminal swabs, as well as vaginal, cervical, and uterine tissues, and were upregulated during vaginal colonization in the GBS CJB111 genetic background reported here and in a previously published data set in the GBS A909 background (31). The function of this putative system remains to be studied.

To maintain intracellular Mn2+ homeostasis, bacteria rely largely on ABC transport systems or natural resistance-associated macrophage proteins (NRAMPs) (35, 53–55). GBS expresses both the mtsABC and mntH systems. The GBS MtsABC transporter is conserved across GBS strains and other streptococcal pathogens (56); however, other pathogenic streptococci, including S. pyogenes and S. pneumoniae, are devoid of NRAMP transporters (57, 58). We detected an underrepresentation of mutations in mtsA across the lumen and all tissue samples collected, and previous analysis of the GBS transcriptome identified mtsA as strongly induced during vaginal colonization (31). We were able to confirm this robust induction of mtsA in another GBS background by performing RNA sequencing during vaginal colonization.

The host has also evolved an elegant strategy, termed nutritional immunity, to limit bioavailable metals from invading pathogens (59, 60). Calprotectin is a unique chelating molecule due to the presence of a distinctive His6 site that creates a high-affinity Mn2+ binding site (32, 33). Our laboratory has previously conducted a transposon mutant screen of GBS grown in the presence of calprotectin, and interestingly, several of the systems we observed as important in vivo were also deemed important for calprotectin survival in vitro, including the mtsABC system. We show here that the ΔmtsA mutant strain is more susceptible to chelation by calprotectin than WT GBS, and while supplementation of extracellular Mn2+ rescues growth of calprotectin-treated WT GBS, the ΔmtsA mutant strain remains susceptible to chelation despite Mn2+ supplementation. While our work has focused on the role of MtsA in Mn2+ homeostasis, in light of previous work showing the ability of calprotectin to also induce iron stress (61–63) and knowledge that expression of the GBS mtsABC locus is induced under both Mn2+ and iron limitation (64), it will be of interest to examine whether the mtsABC locus is also involved in iron uptake and how this might contribute to GBS colonization of the FRT.

The impact of bacterial Mn2+ homeostasis in the FRT had not been described. Here, we utilized a murine model of vaginal colonization previously established in our laboratory (4, 27) to compare fitness of WT GBS and the ΔmtsA mutant strains. Using two different GBS genetic backgrounds, CJB111 (serotype V) and A909 (serotype Ia), we demonstrate that the ΔmtsA mutant strain has a significant decrease in fitness in the FRT compared to WT GBS during competition and single challenge, confirming the importance of Mn2+ homeostasis for GBS vaginal colonization and persistence. Interestingly, we observed more dramatic phenotypes of the ΔmtsA mutant during competition with the WT strain both in vitro and in vivo. This suggests that manganese homeostasis is an important factor for GBS fitness and may also significantly contribute to GBS competition with other members of the vaginal microbiota. We further performed these animal studies in WT and S100A9−/− mice that lack functional calprotectin. We observed a similar decrease in fitness of the ΔmtsA mutant strain in S100A9−/− mice, suggesting that the availability of this metal remains limited even in the absence of calprotectin. These data could provide insight into the limitation of bioavailable Mn2+ within the vaginal mucosa, and studies on how the vaginal microbiota and
host mucosal immunity impact Mn\(^{2+}\) levels warrants further investigation. Mn\(^{2+}\) is also a known metal ion cofactor for bacterial proteins that function in response to oxidative stress (35, 36, 65). This is consistent with our observation that the ΔmtsA mutant was significantly more sensitive to superoxide (paraquat) and H\(_2\)O\(_2\) treatment. GBS is known to express a single Mn\(^{2+}\)-dependent superoxide dismutase (66), and future studies will seek to elucidate the role of this and other GBS Mn\(^{2+}\)-dependent enzymes during colonization of the FRT.

In this study, we utilized for the first time a saturated GBS transposon library during \textit{in vivo} colonization to begin to understand the important factors for GBS persistence in the vaginal lumen and ascension to reproductive tract tissues. We identified numerous factors involved in GBS fitness in the FRT, including MtsA, which we confirmed contributes to GBS persistence in the vaginal tract. Our results further suggest that MtsA has a role in the maintenance of Mn\(^{2+}\) homeostasis and resistance to calprotectin-mediated metal chelation and contributes to the ability of GBS to withstand oxidative stress. MtsA is highly conserved across Gram-positive pathogens, is a component of the GBS core genome (67), and is essential for fitness in the female vaginal tract, thus warranting a more thorough understanding of its role in GBS physiology and suggesting its potential as a future therapeutic target.

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions.** 

\textit{Streptococcus agalactiae} (GBS) isolates CJ8111 (serotype V) and A909 (serotype Ia) and mutant strains CJ8111 ΔmtsA and A909 ΔmtsA were cultured in Todd-Hewitt broth (THB) at 37°C. Mutant strains were constructed as previously described (30). Briefly, genomic 5' and 3' regions flanking the \textit{mtsA} gene were amplified and fused with a spectinomycin cassette by FailSafe PCR (Lucigen). Fragments and pHY304 vector were digested with restriction enzymes XbaI and XmaI and ligated using a Quick Ligation kit (NEB). The ligation reaction product was transfected into \textit{Escherichia coli} MC1061 and selected on LB agar with ampicillin. pHY304 plasmids were purified from \textit{E. coli} and electroporated into GBS CJ8111 and A909 genetic backgrounds and selected on Todd-Hewitt agar (THA) plates with 100 \(\mu\)g/mL spectinomycin. The WT and mutant strains grew to the same optical density at stationary phase (−5 h) in nutrient-rich THB (see Fig. S5 in the supplemental material). The ΔmtsA/ΔmtsA complemented strain (ΔmtsA\(^{+}\)) was generated using the pDCERM expression vector, digested with XbaI and EcoRI-HF, and electroporated into GBS ΔmtsA strains, and mutants were selected on THA with erythromycin at 5 \(\mu\)g/mL. The complemented strain exhibited a significant growth defect during log phase (Fig. S5). All constructs were confirmed by PCR and sequencing. Primers used in the construction of the \textit{mtsA} mutant and complemented strains are listed in Table S5.

**In vivo transposon library screening during vaginal colonization.** All animal experiments were conducted under the approval of the Institutional Animal Care and Use Committee (approval no. 00316) at the University of Colorado Anschutz Medical Campus and performed using accepted veterinary standards. We utilized a murine model of vaginal colonization as previously described (4, 27), in which female CD-1 mice were synchronized on day 1 with 17β-estradiol by intraperitoneal injection. On day 0, triplicate overnight cultures of GBS pooled \textit{Km} library were back-diluted into THB with kanamycin at 300 \(\mu\)g/mL and grown to OD\(_{600}\) 0.4. Cultures were normalized in triplicate to 1 \(\times\) 10\(^{5}\) CFU/10 \(\mu\)L and inoculated by pipetting directly into the vaginal lumen of triplicate groups with 3 mice/group (\(n = 9\) total). Swabs of the vaginal lumen were collected on days 1, 2, and 3, and on day 3 post-colonization, mice were euthanized and vaginal, cervical, and uterine tissues were harvested. Samples collected from daily swabs and tissue homogenates were used to quantify the GBS CFU burden over time, and the remainder of each sample was spread plated onto CHROMagar Strep B with 300 \(\mu\)g/mL kanamycin to collect recovered transposon insertion mutants. Bacterial growth from spread plates of daily swabs and tissues were collected and pooled into triplicate groups, and genomic DNA was extracted using a ZymoBiotics DNA miniprep kit (Zymo Research).

**Transposon library sequencing.** Library preparation and sequencing were performed as previously described (68) by the Microarray and Genomics Core at the University of Colorado Anschutz Medical Campus. Briefly, genomic DNA was sheared to approximately 340-bp fragments and processed through the Ovation ultralow V2 DNA-Seq library preparation kit (Tecan), and 9 ng of each library was used as a template for enrichment by PCR (16 cycles) for the transposon insertions using \textit{Km}it-specific (TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGCCGGGGACTTATCATCCAACC) and Illumina P7 primers. The enriched PCR products were diluted 1:100, and 20 \(\mu\)L was used as a template for an indexing PCR (9 cycles) using the TruSeq P5 indexing and P7 primers. Sequencing was performed to obtain roughly 20 million reads per sample using an Illumina NovaSeq 6000 system in a 150-base paired-end format.

**In vivo Tn-seq bioinformatic analyses.** Analysis of transposon sequencing data was performed as previously described with minor modifications (26), including the use of the recently improved \textit{Streptococcus agalactiae} CJ8111 (CP063198.1) genome as a reference for these reads. A range of 6 to 61 million raw reads/sample (518 million in total) were queried for quality with FastQC (69), and of those, a range of 2.6 to 31 million reads/sample (255 million in total) were deemed suitable and mapped to the GBS CJ8111 (CP063198.1 RNA) genome. To identify \textit{mariner} inverted terminal repeats (ITRs) and transposon insertion TA sites, a Bio::SeqIO-based script...
was written (bin/consolidate_reads.pl), and when the ITR to TA was observed on the negative strand, the sequence was reverse complemented to trim to the insert TA. The number of observed insertion points on each strand was counted, before removal of the mariner ITR leading sequences with cutadapt (70). In a further attempt to verify that the samples were sufficiently robust, we used the TPP preprocessing tools provided by the TRANSIT package (29). This used bowa (71) rather than bowtie/htisal2 for mapping and generating wig files suitable for the downstream TN-seq TRANSIT analytical methods. Invocations and preprocessing methods were recorded in the markdown document, tnseq_transit.Rmd. The libraries were quantified with respect to relative coverage, similarity, and saturation with respect to available TA insertion points by using the bowtie-derived counts (72, 73). The resulting alignments were converted to sorted/compressed binary alignments (74) and counted (75) against the reference genome coding DNA sequence (CDS) and intergenic regions. Comparison and normalization of control (input) and experimental (output) libraries were performed using DESeq2. Many of the postprocessing tasks were handled via the hpqtool R package (76), and functions used are italicized and suffixed with “fl”. Clustering of orthologous groups of proteins (COGs) were assigned using EggNOG 5.0.0 (77), and Venn diagrams were calculated and plotted using BxToolBox (https://www.bioinforx.com).

In vivo RNA sequencing during vaginal colonization and bioinformatic analyses. For collection of control RNA from GBS grown in vitro, GBS was grown in triplicate in THB to an OD600 of 0.4, pelleted, resuspended in 1 mL TRIzol reagent, and stored at −80°C. For collection of RNA from GBS in vivo, we utilized a murine model of vaginal colonization as previously described (4, 27), in which female CD-1 mice were synchronized on day −1 with 17β-estradiol by intraperitoneal injection, and the next day, mice were vaginally inoculated with 10^7 CFU of GBS CJB111. On days 1 and 3 postinoculation, the vaginal lumen of mice was swabbed, and swabs were placed into 100 μL of TRIzol reagent (Thermo Fisher), vortexed to dissociate bacteria from swabs, and stored at −80°C. Swab samples from 6 mice were pooled, and bacteria were lysed by bead beating for 2 min (30-s intervals with 30 s of ice treatment between each bead-beating cycle) at maximum speed. RNA was isolated following the manufacturer’s protocol using a Direct-zol RNA miniprep plus kit (Zymo Research).

Total RNA samples were sequenced by Illumina sequencing at the University of Colorado Anschutz Medical Campus Genomics and Microarray Core. One hundred twenty micrograms of total RNA was depleted of rRNAs, and libraries underwent 9 cycles of PCR prior to 1× AMPure bead purification. Libraries were sequenced using the Illumina NovaSeq 6000 sequencing platform with 75-base single reads at 40 million reads/sample. Raw sequencing reads in fastq format were aligned and annotated to a file containing rRNA reads from the clinical group B streptococcal isolate CJB111 reference genome (CP063198_sRNA) using the following QiaGen CLC Genomics Workbench default settings (version 21.0.5): mismatch cost, 2; insertion and deletion cost, 3; and length and similarity fraction, 0.8. Unmapped genes were then mapped to the CJB111 reference genome (CP063198_sRNA) for RNA-seq analysis in CLC. Volcano plots for differentially expressed genes were generated using GraphPad Prism 9.0.

ICP-MS. Overnight cultures of GBS CJB111 WT, ΔmctsA, and ΔmctsA+ strains were back diluted into 10 mL fresh THB and were grown to an OD600 of 0.4. Bacterial cells were pelleted, rinsed in 1 mL nuclease-free H2O, pelleted again, and desiccated at 65°C for 1 h. For inductively coupled plasma mass spectrometry (ICP-MS), bacterial pellets were digested at 65°C overnight in 50 μL nitric acid. Samples were centrifuged and diluted 1:10 into an Agilent ICP-MS 7500cx autosampler. Mn55 concentrations in parts per billion (μg/mL) were determined by plotting samples against a standard curve and were normalized across bacterial strains by cell dry weight. The data are represented as the mean Mn55 concentration from four independent cultures of WT or ΔmctsA mutant or five independent cultures of GBS WT, ΔmctsA, and ΔmctsA+ strains.

Calprotectin growth assays. Briefly, GBS cultures grown overnight for 18 h in THB were diluted 1:50 into 100 μL in a 96-well microtiter plate with 38% 3× THB and 62% calprotectin buffer (20 mM Tris [pH 7.5], 100 mM NaCl, 3 mM CaCl2) and 0 to 960 μg/mL recombinant calprotectin (78, 79). The range of recombinant calprotectin was previously established for assaying bacterial survival in vitro (33, 54). At 6 h postinoculation, growth was assessed by measuring the optical density (OD600). In experiments where Mn55 was supplemented to counter calprotectin chelation, MnCl2 was added at 10 μM.

Quantitative reverse transcriptase PCR (qRT-PCR). GBS were grown to mid-logarithmic phase (OD600 0.4) in 3× THB medium plus calprotectin buffer and incubated with calprotectin at 960 μg/mL alone for 30 min or calprotectin at 960 μg/mL for 15 min followed by 50 μM MnCl2 supplementation for an additional 15 min. Following incubation, bacteria were centrifuged at 5,000 × g for 5 min, total RNA was extracted (Macherey-Nagel), and cDNA was synthesized (QuantA Biosciences) per the manufacturers’ instructions. Primers used in this study are listed in Table SS.

Oxidative stress experiments. Overnight cultures of GBS CJB111 WT and ΔmctsA were back-diluted 1:50 into fresh THB control medium or THB with added 0 to 0.01% H2O2 or 0.25 mM paraquat and were grown at 37°C for 6 h. Following incubation, optical density (OD600) was measured. All experiments were performed in technical triplicate wells and were repeated in three biologically independent experiments. The mean result for the technical replicate wells was calculated for each independent experiment, and the data shown are the mean calculated from all three biologically independent experiments.

Statistical analyses. Significance was determined using an unpaired two-tailed Student’s t test for ICP-MS experiments and CD-1 tissue burden data, one-way analysis of variance (ANOVA) with Tukey’s multiple-comparison test for qRT-PCR experiments, two-way ANOVA with Tukey’s multiple-comparison test for calprotectin growth experiments, and two-way ANOVA with Sidák’s multiple-comparison test for colonization data. C37BL/6 and 5t049Δ/mctsA tissue burden, and oxidative stress experiments. Statistical significance was accepted when the P value was <0.05, with α equal to 0.05. All statistical analyses were performed using GraphPad Prism 9.3.1.

Data availability. Sequencing reads from the transposon sequencing analyses are available in the NCBI Sequence Read Archive (SRA) under accession numbers PRJNA820592 (Tn-seq) and PRJNA821062 (RNA-seq).
SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, TIF file, 1 MB.
FIG S2, TIF file, 1 MB.
FIG S3, TIF file, 0.5 MB.
FIG S4, TIF file, 0.6 MB.
FIG S5, TIF file, 0.3 MB.
TABLE S1, XLSX file, 0.1 MB.
TABLE S2, DOCX file, 0.03 MB.
TABLE S3, DOCX file, 0.02 MB.
TABLE S4, XLSX file, 0.1 MB.
TABLE S5, DOCX file, 0.01 MB.

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