Manganese Homeostasis in Group A *Streptococcus* Is Critical for Resistance to Oxidative Stress and Virulence

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**ABSTRACT**  *Streptococcus pyogenes* (group A *Streptococcus* [GAS]) is an obligate human pathogen responsible for a spectrum of human disease states. Metallobiology of human pathogens is revealing the fundamental role of metals in both nutritional immunity leading to pathogen starvation and metal poisoning of pathogens by innate immune cells. Spy0980 (MntE) is a paralog of the GAS zinc efflux pump CzcD. Through use of an isogenic mntE deletion mutant in the GAS serotype M1T1 strain 5448, we have elucidated that MntE is a manganese-specific efflux pump required for GAS virulence. The 5448ΔmntE mutant had significantly lower survival following infection of human neutrophils than did the 5448 wild type and the complemented mutant (5448ΔmntE::mntE). Manganese homeostasis may provide protection against oxidative stress, explaining the observed *ex vivo* reduction in virulence. In the presence of manganese and hydrogen peroxide, 5448ΔmntE mutant exhibits significantly lower survival than wild-type 5448 and the complemented mutant. We hypothesize that MntE, by maintaining homeostatic control of cytoplasmic manganese, ensures that the peroxide response repressor PerR is optimally poised to respond to hydrogen peroxide stress. Creation of a 5448ΔmntE-ΔperR double mutant rescued the oxidative stress resistance of the double mutant to wild-type levels in the presence of manganese and hydrogen peroxide. This work elucidates the mechanism for manganese toxicity within GAS and the crucial role of manganese homeostasis in maintaining GAS virulence.

**IMPORTANCE** Manganese is traditionally viewed as a beneficial metal ion to bacteria, and it is also established that most bacteria can tolerate high concentrations of this transition metal. In this work, we show that in group A *Streptococcus*, mutation of the mntE locus, which encodes a transport protein of the cation diffusion facilitator (CDF) family, results in accumulation of manganese and sensitivity to this transition metal ion. The toxicity of manganese is indirect and is the result of a failure of the PerR regulator to respond to oxidative stress in the presence of high intracellular manganese concentrations. These results highlight the importance of MntE in manganese homeostasis and maintenance of an optimal manganese/iron ratio in GAS and the impact of manganese on resistance to oxidative stress and virulence.

Manganese is important for defense against oxidative stress, especially in Gram-positive bacteria (1, 2), and the acquisition of manganese has been shown to be crucial for survival and virulence of *Streptococcus pneumoniae* (3) and *Streptococcus pyogenes* (group A *Streptococcus* [GAS]) (4). Competition for transition metal ions has emerged as a significant component of the innate immune defense system against pathogens (5), and the ability of the host to withhold metal ions (zinc, iron, and manganese) from bacterial pathogens is an important aspect of nutritional immunity, defined here as the restriction of bacterial survival through control of available metal nutrients (6, 7). This is exemplified by the action of the host protein calprotectin, which exerts antimicrobial effects against bacterial pathogens via its sequestration of zinc and/or manganese (8).

In excess, the transition metal ions copper, zinc, and iron are toxic to the cell, and thus, bacteria have evolved sophisticated systems that sense excess transition metal ions and remove these by efflux from the cell (copper or zinc) or sequestration of excess metal ions inside the cell (iron and zinc) (9–12). In contrast to the other major first-row transition metal ions, manganese is usually regarded as being well tolerated by bacteria and can be accumulated to millimolar concentrations within *Escherichia coli* and many lactobacilli without any apparent deleterious effects on the cell (13, 14). However, recently it was observed that deletion of the manganese cation diffusion facilitator (CDF) protein, MntE, in *S. pneumoniae* resulted in increased hydrogen peroxide production and decreased virulence in a murine model of infection (15). Also, the high level of sensitivity of *Neisseria gonorrhoeae* to manganese in comparison to *Neisseria meningitidis* was correlated with the absence of a functional manganese efflux pump, MntX, in the former (16, 17). These observations suggest that in some bacteria, manganese efflux and homeostasis are important for cellular survival.
GAS is an obligate human pathogen responsible for a wide variety of diseases ranging from mild infections such as impetigo and pharyngitis to life-threatening invasive diseases such as streptococcal toxic shock syndrome (STSS) and necrotizing fasciitis (18). There are more than 18 million cases of severe diseases caused by GAS infections which result in over 500,000 deaths each year (19). Since the 1980s, there has been an increase in the incidence of severe invasive GAS diseases, and despite sensitivity to penicillins, emerging resistance to macrolide antibiotic treatment has been reported (20). To date, there are no licensed vaccines against GAS (21). During infection, the host innate immune cells, such as neutrophils and macrophages, mobilize to the site of infection, to control GAS by phagocytosis (18). Recently, we showed that the CDF transporter, CzcD, is essential for efflux of zinc from GAS and is important for protection against neutrophil killing and virulence in a mouse model of infection (22), indicating that the innate immune system can use zinc as an antimicrobial agent. In this study, we characterize a second CDF family transporter in GAS, Spy0980, and describe its role in linking manganese and iron homeostasis with resistance to oxidative stress and virulence.

RESULTS

GAS Spy0980 is a homologue of S. pneumoniae MntE and functions as a manganese efflux pump. During our investigation of the GAS zinc efflux pump CzcD, we observed that a second CDF transporter, Spy0980, is encoded in the GAS genome. Spy0980 exhibited approximately 60% amino acid identity (NCBI BLASTp) to the manganese efflux pump MntE from S. pneumoniae, compared to only 20% amino acid identity to the zinc efflux pump CzcD from GAS, S. pneumoniae, or Cupriavidus metallidurans. Phylogenetic analysis revealed that the CzcD and MntE-related proteins clustered into distinct clades (Fig. 1). Hence, GAS Spy0980 was designated MntE and hypothesized to be involved in manganese efflux.

To investigate the role of mntE, an isogenic deletion mutant and complemented mutant were constructed in the well-characterized GAS M1T1 strain 5448 (23). In the absence of manganese, all three strains grew at comparable rates (Fig. 2A), but upon the addition of increasing concentrations of manganese, growth of the 5448ΔmntE mutant was inhibited (Fig. 2B to D). The defective growth phenotype in the 5448ΔmntE mutant was also observed on solid medium (Fig. 2E). Complementation of the deletion mutant fully restored the ability to grow in the presence of manganese (Fig. 2B to E), confirming that this phenotype is attributable to the mntE deletion. While other CDF proteins, such as CzcD and EmfA from S. pneumoniae, C. metallidurans, and Rhizobium etli, provide resistance to multiple metals (24–26), no differential inhibition of the 5448ΔmntE mutant was observed in the presence of cadmium, cobalt, copper, iron, nickel, and zinc (see Fig. S1 in the supplemental material). Hence, in GAS, MntE appears to be specific for manganese tolerance.
To better understand the biochemical basis of the effect of manganese on the growth of the 5448ΔmntE mutant, we determined the intracellular content of manganese using inductively coupled plasma mass spectrometry (ICP-MS) for 5448 wild type (WT), 5448ΔmntE, and 5448ΔmntE::mntE grown in the absence or presence of 1 mM manganese. When grown in Todd-Hewitt broth supplemented with yeast extract (THY broth), the intracellular manganese contents remained identical for the 3 strains (P > 0.05) (Fig. 3A). However, the addition of manganese to the growth medium resulted in a significant accumulation of intracellular manganese in the 5448ΔmntE mutant compared to wild-type and complemented strains (P < 0.001) (Fig. 3A). This result is consistent with a role for MntE in manganese efflux.

Additionally, we tested mntE relative gene expression levels in the presence of manganese. Expression of mntE in the wild type and complemented mutant is 2-fold higher when they are grown in the presence of 1 mM manganese than when grown in THY broth alone (P < 0.001) (Fig. 3B). Furthermore, mntE gene expression was not induced by other metal ions tested (Cd, Co, Cu, Fe, Ni, and Zn) (see Fig. S2 in the supplemental material), indicating that mntE gene expression is specifically induced by manganese.
Acumulation of intracellular manganese reduces tolerance of GAS to oxidative stress. Previous studies have suggested that manganese plays a role in resistance of GAS to oxidative stress (4). We therefore examined the effect of manganese accumulation in 5448ΔmntE on tolerance to oxidative stress, using drop tests performed with increasing hydrogen peroxide and manganese concentrations. At sublethal concentrations of manganese, increasing hydrogen peroxide concentration resulted in reduced survival of the 5448ΔmntE mutant compared to wild-type and complemented strains (Fig. 4A).

PerR mediates resistance to hydrogen peroxide stress in GAS (27). PerR can bind both iron and manganese, but when bound to iron, PerR forms a less stable repressor (Fig. 5A), and it is more sensitive to the presence of hydrogen peroxide (28, 29). As such, we hypothesized that addition of iron would rescue the growth defect of 5448ΔmntE in the presence of manganese. The addition of iron partially restored growth of 5448ΔmntE at otherwise growth-inhibitory concentrations of manganese (Fig. 4B). This suggested that the toxicity of manganese toward GAS might be exerted via a dysfunction in PerR-dependent regulation.

Manganese efflux protects GAS against PerR-regulated oxidative stress. In view of our findings, we have developed a model that describes the way in which manganese efflux would be critical for the PerR-regulated oxidative stress response. Under normal physiological conditions, the iron-loaded PerR responds to peroxide stress by dissociation from the Per box, inducing expression of peroxide-responsive genes (Fig. 5A). However, manganese-loaded PerR (similar to conditions experienced by the 5448ΔmntE mutant in the presence of manganese) results in stabilization of the PerR complex and inability to respond to peroxide stress (Fig. 5B). To test this model, we created a 5448ΔmntEΔperR double deletion mutant that we predicted would be unable to efflux manganese and yet would retain the ability to defend against peroxide stress. This is because the PerR regulon would be constitutively expressed, rescuing growth of the double mutant in the presence of manganese and peroxide. Drop test analysis with manganese and hydrogen peroxide illustrated that growth of the 5448ΔmntEΔperR double deletion mutant was restored to levels approaching those of the wild type (Fig. 5C). This result indicates that the mechanism of manganese toxicity is via the stabilization of the PerR complex, resulting in reduced ability to mount a response to oxidative stress in GAS.

Loss of manganese efflux in GAS results in decreased expression of the DNase Sda1 and increased killing by human neutrophils. The key phage-encoded virulence determinant of M1T1 GAS, DNase Sda1 (30), is crucial for escape from neutrophil extracellular traps (NETs) (31) and is regulated by PerR (32). Western blot analysis demonstrated that Sda1 is uniformly secreted when the strains are grown in the absence of manganese (Fig. 6A). However, when the 5448ΔmntE mutant was grown in the presence of manganese, Sda1 production was reduced compared to the wild type and complemented mutant (Fig. 6A). These results suggest that the accumulation of manganese results in constitutive repression by PerR, thus affecting the expression of PerR-regulated genes such as sda1.

Neutrophils are recognized as the first line of host cellular innate immune defense against GAS, and the DNase Sda1 plays an important role in protecting GAS from neutrophil killing (33). Thus, we examined the level of resistance of 5448ΔmntE to neutrophil killing. 5448ΔmntE exhibited reduced survival compared to the 5448 wild type and complemented mutant (P < 0.001 and 0.05, respectively) (Fig. 6B).

Finally, we assessed the virulence of the 5448ΔmntE mutant using a transgenic humanized plasminogen mouse model of invasive GAS infection (34). Following subcutaneous infection, no significant difference (P = 0.067) was observed in murine survival between 5448 wild type and 5448ΔmntE (Fig. 6C). Taken together, these results indicate that there may be biological niche-specific differences in manganese requirements for GAS.

DISCUSSION

Manganese has been traditionally viewed as a beneficial metal ion for bacterial pathogens, particularly in Gram-positive streptococci (4, 35, 36). The acquisition of manganese is critical for the assembly of defenses against oxidative stress. This includes production of superoxide dismutase (SodA) and the protection of metalloenzymes against oxidative damage (37, 38). In order to acquire manganese, streptococcal species such as S. pneumoniae and GAS possess high-affinity inducible uptake systems. Expression of high-affinity manganese acquisition systems PsaABC in S. pneumoniae (39) and MtsABC in GAS (4) is under the control of the DtxR family protein MtsR (PsaR in S. pneumoniae) (40, 41), which ensures that expression of manganese uptake is repressed when there is sufficiency. However, our results indicate not only that manganese homeostasis is maintained by the control of manganese uptake but that an efflux pump for manganese, MntE, also plays an important role.

Since it is well known that manganese usually exerts antioxidant rather than pro-oxidant effects, it was not immediately obvious why this ion would be toxic to GAS. The answer appears to be linked to the mechanism of action of the peroxide-responsive repressor, PerR (42). In the model for the action of PerR, develop-
opoped in *Bacillus subtilis*, this repressor can exist in two peroxide-responsive states depending on whether it binds ferrous iron or manganese at the reversible ion-binding site (43). The iron-loaded form of PerR is highly sensitive to hydrogen peroxide, and this leads to derepression of oxidative stress defense systems. In contrast, when manganese is present in the reversible binding site, PerR is more stable in the presence of hydrogen peroxide and thus continues to act as a repressor (29). This mechanism of action enables PerR to act as a sensor for hydrogen peroxide and also respond to the intracellular ratio of the pro-oxidant ferrous iron and antioxidant manganese. Our results are consistent with this model; when manganese is in excess in the 5448/H9004 mntE mutant, GAS is not able to defend against exogenously produced hydrogen peroxide (Fig. 5). Our observation that the 5448/H9004 mntE mutant is sensitive to manganese in the absence of externally added peroxide suggests that the MntE and PerR system also has a role in protection against production of hydrogen peroxide produced endogenously by GAS (44). Support for this model also comes from our observation that addition of ferrous iron restored resistance to manganese in the 5448ΔmntE mutant, consistent with the view that excess manganese leads to constitutive repression of PerR-regulated peroxide defenses. An additional secreted protein was observed in the 5448ΔmntE mutant in the presence of manganese (Fig. 6A). We hypothesize that the elevated expression of this protein results from the significant stress placed on the cell physiology of the 5448ΔmntE mutant under these conditions. MntE-type transporters are also present in other bacteria, but their impact on cell physiology may be different, depending on the regulatory networks that control the physiology of the bacterium. In *S. pneumoniae*, the mntE mutant was also sensitive to manganese and produced more hydrogen peroxide. The mutant exhibited reduced virulence in a mouse model of infection (15). However, *S. pneumoniae* lacks the PerR regulator, and so the influence of MntE on cell physiology and pathogenicity must be exerted via a distinct alternate mechanism.

During infection, GAS is subject to peroxide stress as a consequence of the action of innate immune cells such as neutrophils and macrophages (18). Thus, it is crucial that enzymes under PerR
Proposed model of PerR-manganese displacement in GAS. (A) In wild-type 5448, excess manganese can exit the cell and PerR maintains iron as a cofactor. Under hydrogen peroxide stress, metal-catalyzed oxidation occurs to the iron cofactor, leading to damage of PerR, dissociation from Per boxes, and subsequent transcription of peroxide response genes. (B) In the 5448ΔmntE mutant, the presence of manganese and hydrogen peroxide stress results in non-redox-active manganese displacement of iron within PerR. This stabilizes the complex, resulting in continued repression of peroxide response genes. (C) Drop test analysis of 5448, 5448ΔmntE, 5448ΔmntE::mntE, 5448ΔperR, and 5448ΔmntE-ΔperR on THY agar supplemented with increasing concentrations of manganese vertically (0, 0.25, 0.50, and 0.75 mM) and increasing concentrations of hydrogen peroxide horizontally (0, 1, 2, and 4 mM). Cells were grown and adjusted to an OD$_{600}$ of 0.6 and serially diluted, and 5-μl drops were spotted onto the plate from concentrations of $10^0$ (top) to $10^{-3}$ (bottom).
and failure to induce the PerR regulon. This model is consistent with the observation that in an ex vivo human neutrophil killing assay, the 5448ΔmntE deletion mutant is attenuated. On the other hand, in the subcutaneous infection model used in this study, there may be a niche-specific effect occurring, resulting in no significant difference between the virulence of the mutant and that of the WT. Loss of mntE in manganese-depleted and iron-replete conditions subcutaneously may lift selection against the 5448ΔmntE mutant compared to 5448 WT. Such conditions may result due to the action of calprotectin at the site of infection (45). Thus, manganese homeostasis in GAS as well as the ratio of manganese to iron in the bacterial cell may play a significant role in GAS pathogenesis and protection against oxidative stress.

It is also established that mutation of high-affinity manganese acquisition systems in S. pneumoniae (46) and GAS (4) leads to reduced growth and survival of these human pathogens in a mouse model of infection. Manganese may provide protection from oxidative stress through the displacement of redox-sensitive ferrous iron within key enzymes (38, 47), and manganese is also a cofactor in superoxide dismutase in bacterial pathogens such as S. pneumoniae (48) and GAS (37). Recently, it was also observed that in S. pneumoniae, zinc toxicity is a result of the inability of the bacteria to acquire manganese through the high-affinity manganese solute protein, Psaa (49). These observations indicate that there is a dynamic interplay of manganese, iron, and zinc at the host-pathogen interface. Bacterial pathogens are finely tuned to adjust their adaptive responses to altered availability of these transition metal ions via control of acquisition and efflux systems. This is likely to be a key factor in the regulation of virulence in GAS and other streptococci.

MATERIALS AND METHODS

Bacterial strains and growth conditions. The S. pyogenes MIT1 clinical isolate 5448 (23) and isogenic derivatives were routinely grown on 5% horse blood agar or statically in liquid cultures at 37°C in Todd–Hewitt broth supplemented with 1% yeast extract (THY broth). Escherichia coli MC1061 was grown in Luria-Bertani medium (LB). Where required, erythromycin was used at 2 μg/ml (GAS) or 500 μg/ml (E. coli) and spectinomycin was used at 100 μg/ml (both GAS and E. coli). GAS isogenic mutants were constructed as previously described (31). All bacterial strains and plasmids are listed in Table S1 in the supplemental material.

DNA manipulation and genetic techniques. GAS mutants 5448ΔmntE, 5448ΔperR, and 5448ΔmntE::ΔperR were constructed by deletion replacement. All PCR primer sequences are provided in Table S2 in the supplemental material. The 1-kb regions upstream of mntE and perR were amplified using primers mntE1 and mntE4 and perR1 and perR2, respectively. The 1-kb downstream regions were amplified using primers mntE3 and mntE4 and primers perR3 and perR4, respectively. The kanamycin cassette was amplified using primers kan-F and kan-R, while the spectinomycin cassette was amplified using primers spec-F and spec-R. For 5448ΔmntE, the three PCR fragments (5′- and 3′-flanking regions with kanamycin cassette) were joined together with primers mntE1 and mntE4, and for 5448ΔperR, the three fragments (5′- and 3′-flanking regions with spectinomycin cassette) were joined with primers perR1 and perR4. The subsequent fragment was cloned into the pH304 shuttle vector (50) and transformed into E. coli MC1061. Electrotransformation and allelic replacement mutagenesis were undertaken using standard protocols (51, 52). Complementation of 5448ΔmntE was performed by marker rescue. The entire wild-type region of mntE, including the 1-kb upstream and downstream sequences, was amplified using primers mntE1 and mntE4. The construct was cloned into pH304 via restriction ligation and transformed into E. coli MC1061. Electrotransformation and allelic replacement mutagenesis were undertaken as described above, and
complemented strains were screened for the loss of kanamycin resistance. All strains were confirmed by DNA sequencing (Australian Equine Genome Research Centre, University of Queensland, Brisbane, Australia).

**Phylogenetic analysis.** Phylogenetic analyses of 59 cation diffusion facilitator family protein sequences, selected on the basis of BLAST (NCBI) analysis against functionally characterized CDF family proteins, CzcD, EmfA, and MntE, were used to determine the clustering of the different CDF families. An amino acid alignment was generated using Clustal Omega v1.2.1 (53). A best-fit phylogenetic tree was estimated using PhyML v3.0 (54) based on the WAG substitution model with gamma correction of among-site rate variation. One hundred nonparametric bootstraps were applied.

**Growth curve analyses.** Overnight cultures grown in THY broth were diluted to an optical density at 600 nm (OD 600) of 0.05 in fresh THY broth supplemented with various amounts of manganese. The cells were statically grown in a 96-well microtiter plate, and OD 595 was measured half-hourly using the FLUOstar Optima (BMG Labtech) plate reader at 37°C.

**Drop test plate sensitivity assays.** Drop tests were performed as previously described (22). Antimicrobial agents tested were cobalt(II) (CoSO₄·4H₂O), copper(II) (CuSO₄·5H₂O), iron(II) (FeSO₄·7H₂O) and iron(III) (Fe(NO₃)₃·9H₂O), manganese (MnSO₄·4H₂O), nickel(II) (NiSO₄·6H₂O), and zinc(II) (ZnSO₄·7H₂O). These assays were also previously described (22). Antimicrobial agents tested were cobalt(II) (CoSO₄·4H₂O), copper(II) (CuSO₄·5H₂O), iron(II) (FeSO₄·7H₂O) and iron(III) (Fe(NO₃)₃·9H₂O), manganese (MnSO₄·4H₂O), nickel(II) (NiSO₄·6H₂O), and zinc(II) (ZnSO₄·7H₂O). These assays were also repeated in the presence of hydrogen peroxide. Plates were photographically documented following overnight incubation at 37°C. Drop tests are representative of biological replicates performed on at least 3 separate occasions.

**Intracellular metal concentration measurement.** Cells from an overnight THY agar plate, grown in the absence or presence of 1 mM MnSO₄·4H₂O, were resuspended, washed three times with phosphate buffer, 0.25 M EDTA and three times with phosphate buffer, resuspended in 80% nitric acid, and incubated at 80°C for 24 h. The samples were then diluted to 2% nitric acid and submitted for inductively coupled plasmid mass spectrometry (ICP-MS) analysis at the School of Earth Sciences, University of Queensland. The final value was normalized to the amount of cells present by measuring the total protein content in accordance with the QuantiPro bicinchoninic acid (BCA) assay kit (Sigma) instructions.

**Quantitative gene expression studies.** RNA was isolated from cells harvested under the desired growth phase grown in the presence or absence of 1 mM MnSO₄·4H₂O, in accordance with the RNEasy minikit (Qiagen) with the additional mechanical lysis step in lysing matrix B tubes (MP Biomedicals). The isolated RNA was DNase treated using the RNase-free DNase set (Qiagen) and quantified using a NanoDrop instrument (Thermo Scientific). One microgram of RNA was converted to cDNA using the SuperScriptIII first-strand synthesis system for reverse transcription-PCR (RT-PCR) (Invitrogen). Real-time reverse transcription-PCR was performed using the primers specified in Table S2 in the supplemental material. The PCR was performed using SYBR green master mix (Applied Biosystems) according to the manufacturer’s instructions. All data were analyzed using the ViiA7 software (Applied Biosystems). Relative gene expression was calculated using the threshold cycle (2^{-ΔΔCt}) method with proS as the reference gene (55). All reactions were performed in triplicate from 3 independently isolated RNA samples.

**Sda1 Western blotting.** Detection of Sda1 by Western blotting from cell supernatants grown in the presence or absence of 0.5 mM MnSO₄·4H₂O to mid-logarithmic phase was performed using standard protocols (56).

**Neutrophil killing assay.** GAS survival following incubation with human neutrophils ex vivo was assayed as previously described (31). Experiments were performed in triplicate using mid-logarithmic-phase (A 600 = 0.4) GAS at a multiplicity of infection of 10:1 (GAS-to-neutrophil ratio).

**Virulence of GAS in a humanized plasminogen transgenic mouse model.** Transgenic humanized plasminogen mice heterozygous for the human plasminogen transgene (Alb→PBG172→) were infected with a dose of 2 × 10^8 to 4 × 10^8 CFU of either 5448, 5448ΔmntE, or 5448ΔmntE::mntE. Mice (n = 20 for 5448 WT and 5448ΔmntE::mntE, n = 19 for 5448ΔmntE) were subcutaneously infected with freshly prepared GAS strains in 100 μl of 1X phosphate-buffered saline (PBS), and virulence was assessed as previously described (31).

**Ethics approval.** All animal experiments were conducted according to the Guidelines for the Care and Use of Laboratory Animals (National Health and Medical Research Council, Australia) and were approved by the University of Queensland Animal Ethics Committee. Human blood donation for use in neutrophil killing assays was conducted in accordance with the National Statement on Ethical Conduct in Human Research, complied with the regulations governing experimentation on humans, and was approved by the University of Queensland Medical Research Ethics Committee.

**Statistical analyses.** Differences in intracellular metal ion concentration, relative gene expression, and neutrophil survival were analyzed using unpaired Student’s t test or 1-way analysis of variance (ANOVA) with Bonferroni’s posttest as necessary (GraphPad Prism 6). Murine survival curves were analyzed using the Mantel-Cox log rank test (GraphPad Prism 6).

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at http://mbio.asm.orglookup/suppl doi:10.1128/mBio.00278-15/-/DCSupplemental.

**Characteristics of selected CDF family members.**

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