Abstract

Background: In prokaryotes, the ureases are multi-subunit, nickel-containing enzymes that catalyze the hydrolysis of urea to carbon dioxide and ammonia. The *Brucella* genomes contain two urease operons designated as *ure*1 and *ure*2. We investigated the role of the two *Brucella suis* urease operons on the infection, intracellular persistence, growth, and resistance to low-pH killing.

Results: The deduced amino acid sequence of urease-α subunits of operons-1 and -2 exhibited substantial identity with the structural ureases of alpha- and beta-proteobacteria, Gram-positive and Gram-negative bacteria, fungi, and higher plants. Four *ure* deficient strains were generated by deleting one or more of the genes encoding urease subunits of *B. suis* strain 1330 by allelic exchange: strain 1330Δ*ure*1K (generated by deleting *ureD* and *ureA* in *ure*1 operon), strain 1330Δ*ure*2K (*ureB* and *ureC* in *ure*2 operon), strain 1330Δ*ure*2C (*ureA*, *ureB*, and *ureC* in *ure*2 operon), and strain 1330Δ*ure*1KΔ*ure*2C (*ureD* and *ureA* in *ure*1 operon and *ureA*, *ureB*, and *ureC* in *ure*2 operon). When grown in urease test broth, strains 1330, 1330Δ*ure*2K, and 1330Δ*ure*2C displayed maximal urease enzyme activity within 24 hours, whereas, strains 1330Δ*ure*1K and 1330Δ*ure*1KΔ*ure*2C exhibited zero urease activity even 96 h after inoculation. Strains 1330Δ*ure*1K and 1330Δ*ure*1KΔ*ure*2C exhibited slower growth rates in tryptic soy broth relative to the wild type strain 1330. When the BALB/c mice were infected intraperitoneally with the strains, six weeks after inoculation, the splenic recovery of the *ure* deficient strains did not differ from the wild type. In contrast, when the mice were inoculated by gavage, one week after inoculation, strain 1330Δ*ure*1KΔ*ure*2C was cleared from livers and spleens while the wild type strain 1330 was still present. All *B. suis* strains were killed when they were incubated in-vitro at pH 2.0. When the strains were incubated at pH 2.0 supplemented with 10 mM urea, strain 1330Δ*ure*1K was completely killed, strain 1330Δ*ure*2C was partially killed, but strains 1330 and 1330Δ*ure*2K were not killed.

Conclusion: These findings suggest that the *ure*1 operon is necessary for optimal growth in culture, urease activity, resistance against low-pH killing, and in vivo persistence of *B. suis* when inoculated by gavage. The *ure*2 operon apparently enhances the resistance to low-pH killing in-vitro.
Background

A number of environmentally and medically important bacteria produce the enzyme urease (urea amidohydrolase) [1], which catalyzes the hydrolysis of urea, leading to the production of carbamate and ammonia. In an aqueous environment, the carbamate rapidly and spontaneously decomposes to yield a second molecule of ammonia and one of carbonic acid. The carbonic acid equilibrates in water, as do the two molecules of ammonia, which become protonated to yield ammonia hydroxide ions. The reaction results in an increased pH of the environment [reviewed in [2-5]]. In sites where microorganisms colonize epithelial surfaces, such as the normal flora of the oral cavity or intestines, or when certain pathogenic bacteria infect tissues, the metabolism of urea by microbial ureases can have a profound impact on tissue integrity, microbial ecology, and the overall health of the host.

The ureases of most microbes are composed of three subunits α, β, and γ that are encoded by ureA, ureB and ureC genes respectively. The plant jack bean produces a single-subunit urease [12], whereas, in gastroduodenal pathogen H. pylori, the ureA and ureB genes are sufficient to encode urease. Nevertheless the UreAB subunits of H. pylori can be aligned with the UreABC subunits of other ureolytic bacteria and with the single polypeptide of the jack bean urease. The crystal structure of the Klebsiella aerogenes urease reveals a trimeric configuration [13]. Biochemical analyses of ureases by gel filtration have shown that other bacterial ureases are multimeric and probably have similar stoichiometry [4].

Ureases are structurally complex enzymes, and additional urease subunits are required for the production of a catalytically active holoenzyme in-vivo. Ureases are among the few enzymes that require nickel for activity. Biogenesis of a functional urease in prokaryotes requires the presence and expression of four urease accessory genes, ureDFEG. In vitro experiments using purified accessory proteins support the idea that UreE likely acts as a carrier of nickel [14] and that UreDFG form a chaperone-like complex that keeps the apoenzyme in a configuration competent to accept nickel [15].

Urease activity can be a critical factor in the colonization, persistence and pathogenesis of bacteria. Considering the products produced by urease, it would be logical to assume that one of the enzyme’s functions is to allow nitrogen assimilation. In fact, urea represents an assimilable nitrogen source for bacteria that can colonize the human body and there is evidence suggesting that ammonia assimilation from urea occurs in-vivo. A significant proportion of the urea produced in the liver ends up in the intestines, where it can be hydrolyzed and assimilated by several different species of anaerobic, ureolytic bacteria [3]. Similarly, the oral bacterium, Actinomyces naeslundii can use urea as a primary nitrogen source for growth [6]. So there is little doubt that nitrogen acquisition as the result of urease activity can be important in the ecology of complex populations colonizing the human body. However, it is an open question as to whether the capacity to assimilate ammonia produced by urease contributes to the pathogenic potential of bacteria. Instead, it appears that the release of the strongly alkaline ammonia released by urease is a major cause of the damage to the host tissue, and in some cases, a key factor in persistence of pathogens [reviewed in [2]]. Jubier-Maurin et al. (30) identified the nikABCDE operon encoding the specific transport system for nickel in B. suis. Insertional inactivation of nikA strongly reduced the activity of the nickel metalloenzyme urease, which was restored by addition of nickel excess. Intracellular growth rates of the B. suis wild-type and nikA mutant strains in human monocytes were similar, indicating that nikA was not essential for this type of infection.

The Brucellae are gram-negative, facultative intracellular bacterial pathogens of a wide range of vertebrates [7]. This pathogen is the etiologic agent of the disease brucellosis and the pathological manifestations of brucellosis include abortion and sterility in animals [7], and meningitis, endocarditis, spondylitis and arthritis in humans [8]. Paulsen et al. [9] annotated the genome of B. suis strain 1330 (biovar 1), and discovered that unlike many other organisms, Brucella have two urease gene operons located on chromosome I (GenBank accession no. NC_004310). Urease activity is important for the nitrogen assimilation and persistence of other bacterial species like Helicobacter pylori [10,11]. We investigated the role of the two B. suis urease operons on the infection, intracellular persistence, growth, and resistance to low-pH killing. We report that the B. suis ure1 operon, in contrast to ure2, appears to be principally responsible for determining urease activity, optimum growth and resistance to low-pH killing in-vitro and persistence in-vivo.

Results

Organization, and nucleotide and amino acid sequences of urease genes

The ure1 and ure2 operons are located on the chromosome I of B. suis strain 1330 (GenBank accession number NC_004310). The ure1 operon is 5284-bp long and composed of seven coding sequences (CDS). The ure2 operon is 6571-bp long and comprised eight CDS (Figure 1). The ureA gene was the same in size in both operons (302-bp). All the other genes of ure2 operon were slightly longer than their counterparts in ure1 operon. The ureC gene was the longest in each operon (1712-bp in operon-1 and 1721-bp in operon-2). The G+C content of each ure gene was compared with that of its counterpart of the other
operon and found not differ substantially between ure genes of operon-1 and operon-2 (Table 1). The identity of each ure gene was compared with that of its counterpart of the other operon. The ureA, ureB, ureC, and ureG genes of the two operons exhibited 52 to 60% identity, whereas the ureD, ureE, and ureF genes did not share significant identity (Table 1).

The deduced amino acid sequences encoded by the ureA, ureB, and ureC genes in both operons displayed great identity with the structural urease subunits of a vast range of organisms including Gram-positive bacteria, Gram-negative bacteria, photosynthetic bacteria, fungi, and higher plants (see Table 5). For instance, urease subunits of other organisms exhibited up to 81% identity with the UreC of ure1 and up to 69% identity with the UreC of ure2. The ureases of alpha and beta-proteobacteria exhibited the greatest identity with UreC of operon-1, whereas, the ureases of all species of Yersinia exhibited the greatest identity with the UreC of operon-2.

Real-time PCR assays produced amplicons in sizes exactly identity with the UreC of operon-2. In qualitative urease assay, urease test broth started turning positive within 4 h after either strain 1330, 1330Δure2K, or 1330Δure2C were introduced, and acquired a bright pink color after approximately 24 h (Figure 3 and Table 2). In contrast, strains 1330Δure1K and 1330Δure1KΔure2C failed to cause a pink color in the urease test broth even after 96 h of incubation (Figure 3 and Table 2).

Strains 1330Δure1K and 1330Δure1KΔure2C, both urease negative, grew approximately 25% slower than wild type strain 1330. In contrast, strains 1330Δure2K and 1330Δure2C, both urease positive, did not display any measurable differences in growth rate compared to strain 1330 (Table 2).

Survival of B. suis strains in macrophage cell lines
When used to infect J774A.1 or H36.12a [Pixie 12a] mouse macrophage cell lines, the recovery of all the B. suis strains declined 2–3 log_{10} cfu between 0 and 4 h post-inoculation. During the next 20 h, all the B. suis strains increased 1–2 log_{10} cfu. There were no significant differences between the wild type and the urease mutant strains in terms of their ability to replicate in macrophages (data not shown).

Survival of B. suis strains in BALB/c mice
Following an intraperitoneal inoculation, the recovery of ure mutants from spleens did not differ from the wild type strain at 6 wks post-infection (Table 3). When the mice were inoculated by gavage, one week after inoculation, strain 1330 was recovered from spleens (Figure 4) as well as from livers (Figure 5). When the mice were inoculated with strain 1330 supplemented with 10 mM urea, nearly 2.2 log_{10} greater cfu was recovered from spleens and nearly 3.5 log_{10} greater cfu was recovered from livers. However, when the mice were inoculated with strain 1330Δure1KΔure2C, with or without urea supplementation, no cfu were recovered from spleens (Figure 4) but nearly 2.5 log_{10} cfu was recovered from livers only when the inoculum was supplemented with 10 mM urea (Figure 5).

Resistance of B. suis strains against low-pH killing
The wild type and the ure mutants did not differ with respect to the survival after 90 min incubation at pH 4.0 or 7.0 (data not shown). All the strains including the wild type were killed when incubated at pH 2.0 for 90 min.
Table 1: Sequence identities between the two B. suis urease operons

| Gene | Operon-1 | Operon-2 | Gene comparison | Identity (%) |
|------|----------|----------|-----------------|--------------|
| ureA | 60.3     | 57.3     | ure1A vs ure2A  | 52           |
| ureB | 58.4     | 58.1     | ure1B vs ure2B  | 60           |
| ureC | 60.4     | 59.3     | ure1C vs ure2C  | 55           |
| ureD | 62.0     | 58.1     | ure1D vs ure2D  | 23           |
| ureE | 59.3     | 58.9     | ure1E vs ure2E  | Not significant |
| ureF | 63.3     | 59.1     | ure1F vs ure2F  | Not significant |
| ureG | 57.4     | 56.5     | ure1G vs ure2G  | 54           |
| ureT | -        | 60.4     | -               | -            |

(Figure 6). When the strains were supplemented with 5 mM urea during incubation at pH 2.0, more than 6.0 log<sub>10</sub> cfu of strains 1330 and 1330<sub>Δure2K</sub> were recovered. In comparison to strain 1330, the recovery of the strain 1330<sub>Δure2C</sub> was nearly 1.5 log<sub>10</sub> lower at 5 mM urea concentration and nearly 1.0 log<sub>10</sub> lower at 10 mM urea. In contrast to strains 1330, 1330<sub>Δure2K</sub> and 1330<sub>Δure2C</sub>, strain 1330<sub>Δure1K</sub> was not recovered after incubation at pH 2.0 supplemented at any urea concentration (Figure 6). Addition of urea did not change the pH of the incubation media.

Discussion

The ure<sub>A</sub>, ure<sub>B</sub>, and ure<sub>C</sub> genes of B. suis (Figure 1) encode the γ, β, and α subunits respectively, and the urease holoenzyme of B. suis is likely to be assembled in a trimeric configuration. The total predicted mass of the B. suis urease holoenzyme (UreA+B+C) is 91-kDa. The native
polyacrylamide gel reveals urease activity at approximately 95-kDa (Figure 2) and supports a trimeric configuration of this enzyme. The *Brucella* genome also contains *ure*DEFG genes (Figure 1) in each of *ure*1 and *ure*2 operons and are predicted to produce the UreD, UreE, UreF and UreG proteins. Unlike many other microorganisms, *Brucella* contains two operons encoding urease subunits (Figure 1) located on the chromosome I. Based on the similarities of G+C contents among genes in *ure*1 and *ure*2 operons, it is unlikely that any of the operons were acquired by horizontal gene transfer. The genes of the *ure*1 operon shared less than 60% identity with their counterparts of the *ure*2 operon (Table 1). In particular, the *ureE* and *ureF* genes of the *ure*1 operon did not share considerable similarity with those genes in the *ure*2 operon. Based on the relatively low identity among genes between *ure*1 and *ure*2 operons, it seems unlikely that they were the result of a recent duplication event. However, further analyses are required to confirm these predictions.

We generated a series of mutants by disrupting the first few genes encoding structural subunits of each urease. All seven genes of *ure*1 operon appear to be transcribed in a single direction. The gaps between individual *ure* genes are extremely small (Figure 1), so that all or most of the genes are possibly expressed under a single, common promoter – leading to a polycistronic mRNA. The *ureG* is the last gene of the *ure*1 operon. All three genes downstream of the *ureG* are transcribed in the opposite direction, from the complementary strand (Figure 1). The closest non-*ure*

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**Table 2: B. suis strains: generation time (doubling time, h) in TSB and urease activity in urease test broth.**

| Strain                  | Doubling time (h)* | Urease activity |
|-------------------------|--------------------|-----------------|
|                         |                    | Qualitative**   | Quantitative*** |
| 1330                    | 5.1                | +               | 9.28            |
| 1330*ure1K              | 5.8                | -               | ~0              |
| 1330*ure2K              | 5.3                | +               | 9.38            |
| 1330*ure2C              | 5.1                | +               | 8.28            |
| 1330*ure1K1330*ure2C    | 6.5                | -               | ~0              |

*A* A representative experiment was used to calculate the generation time.

**+** represents positive or pink color and – represents negative or yellow color of the uninoculated test broth; the color remained unchanged even 96 h after inoculation

***Specific activity μmoles/min/mg of protein.
gene to ureG encodes an Arg-tRNA. The genes downstream of this tRNA gene encode a serine histidine kinase (CDS Shk) and a hypothetical protein. The distance from the stop codon of ureG to the stop codon of Arg-tRNA (transcribed in opposite direction) is 124-bp. Thus it does not seem likely that the insertion of the antibiotic resistance gene into ure1 influences the expression of genes downstream of the operon. In addition, it is not apparent that these genes have any regulatory role on urease expression.

The deletion of genes within the ure1 operon caused the disappearance of urease activity on a native polyacrylamide gel (Figure 2). Even though the ure2 operon was present in the ure1 mutant, it failed to produce urease activity as measured by either of two assays. In addition, a disruption of the ure2 operon did not have any impact on production of a detectable urease activity.

The deletion of genes within the ure1 operon caused the slower growth and loss of urease activity (Table 2) suggest-

| Strain Injected dosage (log₁₀ cfu/mouse) | cfu 6 weeks after inoculation (log₁₀/spleen) |
|----------------------------------------|---------------------------------------------|
| 1330 (wild)                            | 5.24                                        | 4.41 ± 0.18†                             |
| 1330Δure1K                             | 5.24                                        | 4.61 ± 0.20‡                             |
| 1330Δure2K                             | 5.22                                        | 4.40 ± 0.18†                             |
| 1330ΔctpA (control)                    | 5.25                                        | 2.04 ± 0.89§                             |
| 1330 (wild)                            | 4.11                                        | 4.25 ± 0.23                              |
| 1330Δure2C                             | 4.16                                        | 4.17 ± 0.25                              |
| 1330Δure1KΔure2C                       | 4.28                                        | 4.19 ± 0.31                              |

Table 3: Splenic recovery of B. suis strains six weeks after intraperitoneal inoculation in BALB/c mice

| Inoculation group | Splenic recovery of brucellae (log₁₀ cfu/spleen) |
|-------------------|-----------------------------------------------|
| 1330 with urea    | 100†                                          |
| 1330              | 100‡                                          |
| 1330-ure1K-ure2C  | S                                            |
| 1330-ure1K-ure2C  | $                                            |

Figure 4
Recovery of Brucella cfu from spleens one week after BALB/c mice were inoculated by gavage with wild type strain 1330 or strain 1330Δure1KΔure2C with or without urea supplementation. P value for the difference among mean values was <0.01. The mean values that share the same symbol do not differ from one another significantly; and the mean values designated by different symbols differ from one another significantly.
ing that the genes are necessary for maximal growth and urease activity of *B. suis*. Strains 1330Δure2K and 1330Δure2C made by deleting genes of *ure2* operon did not display any change in growth rate or urease activity, suggesting that the genes of *ure2* operon are not required for these functions. Overall, *B. suis* is apparently capable of exhibiting urease activity even without involvement of *ure2* operon. This observation raises the question as to whether the genes within *ure2* are actually being expressed. Our measurements using RT-PCR of RNA extracted from *B. suis* 1330 cultured in TSB (data not shown) suggest that all the genes in *ure2* are being expressed. It is apparent that much more work will have to be done to determine the role of the *ure2* operon. Except for a detectable role in resistance to acidic pH *in vitro*, there were no other detectable phenotypes associated with the *ure2* mutants under the various conditions employed. One obvious question is whether there is any post-translational interaction going on between the subunits of both *ure* operons. Given that there is no urease activity in an *ure1* mutant, it is possible to conclude that the corresponding Ure2 subunits are not acting to restore Ure1 activity.

In addition to the ureolytic bacteria [3] that can use urea as a primary nitrogen source for growth, the urinary tract pathogen *Ureaplasma urealyticum* [16] and some alkalophilic bacteria [17] can use ureolysis to generate ATP. Urea, particularly at millimolar concentrations can readily enter the cell [4]. Thus, in bacteria that utilize urea for nitrogen assimilation or ATP generation, the urease activity is expected to occur intracellularly. In most organisms, the ureases are found in the cytoplasm, although there is a report of urease membrane association and cell surface localization as well [4]. The ureases for which primary sequence information is available do not have characteristics consistent with being integral membrane proteins or secreted through the general secretory pathways. Consistent with the ureases of other bacteria, the deduced *B. suis ure* encoded subunits (with exception of UreB encoded by the *ure2* operon) are predicted to localize in the cell cytoplasm (data not shown). Thus it may be possible that urease functions within the cytoplasm in order to assimilate nitrogen. Further work is needed to characterize the exact location of urease subunits and whether they play a role in nitrogen uptake.

The *ure* mutants did not exhibit any decline in persistence in BALB/c mice when inoculated intraperitoneally (Table 3), suggesting that the urease activity is not critical once the pathogen has entered the host. Using *ure* mutants of *Bordetella bronchiseptica*, Monack and Falkow [18]
reported that urease is not essential for the colonization of the guinea-pig respiratory and digestive tracts. One of the common ways that humans acquire brucellosis is through consumption of contaminated milk. *Brucella* needs to have a mechanism to resist the low-pH killing in the gastrointestinal tract. When the mice were inoculated by gavage, the wild type strain was recovered from livers and spleens, whereas, strain 1330Δure1KΔure2C was not. It is noteworthy that when the infecting doses of *B. suis* were supplemented with gavage, the wild type strain was recovered from livers and spleens, whereas, strain 1330Δure1KΔure2C was not. It is noteworthy that when the infecting doses of *B. suis* were supplemented with urea, the recovery of strain 1330 from livers and spleens was increased, but the mutant strain 1330Δure1KΔure2C was recovered only from livers. These findings suggest that the urease activity and sufficient substrate is needed for *B. suis* to cope with the low pH in the gastrointestinal tract i.e. either stomach or upper intestine. Similar observations have been recently reported by Sangari et al., [37] in urea-mutant strains of *B. abortus*. Thus, similar to the urease of *H. pylori* [10,11], that of *B. suis* appears to be a factor in coping with the pH of the gastrointestinal tract. Even though the ure1, ure2 mutant *B. suis* exhibited no urease activity, urea supplementation raised the recovery of this strain from livers. This is possible because other bacterial species in stomach and gastrointestinal tract may have utilized the supplemented urea to reduce the acidity and facilitated enhanced survival of *Brucella*. It is worth noting that inoculation via an oral route versus direct inoculation in the stomach may be more representative of a natural infection. However, we chose to deliver a known dose of *Brucella* and measure uptake through the gastro-intestinal tract (i.e., into the spleen and liver).

For an intracellular bacterium like *Brucella* that replicates inside phagocytic cells associated with the various tissues including the reproductive tract [19], urease may not be important for regulating the pH in an intracellular milieu. Correspondingly, the lack of differences between wild type and ure mutants inoculated intraperitoneally during splenic clearance was not surprising (Table 3). Nevertheless, in the case of *H. pylori* and perhaps other organisms, urease is an important factor in survival in severely acidified environments, probably because neutralization of the extracellular milieu around the cells is needed to prevent irreversible membrane damage. Acidification and subsequent alkalinization in the phagosomal compartment is a prerequisite for a successful *Brucella* infection in-vitro [20].
However, the amount of urea present in macrophages maybe low enough as not to cause a significant pH change. As such, the impact of urease activity on macrophage pH can be considered insignificant, and therefore, similar survival rates in macrophages between wild-type and ure mutants was not unexpected.

Conclusion
The B. suis genome contains two operons encoding urease. The genes in the ure1 operon are important for optimal growth in culture, and indispensable for urease activity, resistance to low-pH killing and survival of the pathogen when inoculated by gavage. The genes in the ure2 operon seem to be dispensable for the above functions, but slightly enhance the resistance to low-pH killing measured in-vitro. The apparent lack of urease activity encoded by ure2 is consistent with the observation by Hooper and Berg who reviewed over 20 microbial genomes with respect to gene innovation following gene duplication (38). They concluded that most gene copies are deleted but of the ones retained, they function in weak or ancillary roles. Thus it is possible that ure2 in B. suis is diverged enough to encode a new function that has yet to be defined.

Methods
DNA and protein sequence analyses
The nucleotide sequence of the urease genes was analyzed with DNASTAR software (DNASTAR, Inc., Madison, Wis.). The destination of the deduced proteins upon translation and processing was predicted using the Subloc v1.0 server of the Institute of Bioinformatics of the Tsinghua University http://www.bioinfo.tsinghua.edu.cn/. Identity of the ureA, ureB, and ureC genes of B. suis ure1 and ure2 operons with sequences of the EMBL/GenBank/ DDBJ databases was analyzed using the BLAST software [22] at the National Center for Biotechnology Information (Bethesda, MD).

Bacterial strains, plasmids, and reagents
B. suis strain 1330 was obtained from our culture collection. Escherichia coli strain Top10 (Invitrogen Life Technologies, Carlsbad, Calif.) was used for producing plasmid constructs. E. coli were grown in Luria-Bertani (LB) broth or on LB agar (Difco Laboratories, Sparks, MD). Brucella were grown either in Trypticase soy broth (TSB) or on Trypticase soy agar (TSA) plates (Difco) at 37°C in the presence of 5% CO2 as previously described [23]. The plasmids used in this study are listed in Table 4. Bacteria containing plasmids were grown in the presence of ampicillin or kanamycin at a 100-μg/ml concentration (Table 4).

All experiments with live Brucella were performed in a Biosafety Level 3 facility in the Infectious Disease Unit of the Virginia-Maryland Regional College of Veterinary Medicine per standard operating procedures approved by the Centers for Disease Control and Prevention.

Recombinant DNA methods
Genomic DNA was isolated from B. suis strain 1330 by use of a QIAGEN blood and tissue DNA kit (QIAGEN Inc., Valencia, Calif.).

Table 4: Description of the plasmids and bacterial strains used in this study

| Plasmid or strain | Description | Source or reference |
|-------------------|-------------|---------------------|
| pCR2.1 | TA cloning vector, 3.9-kb, Amp' | Invitrogen |
| pCR2.1 | pCR2.1 with 2.2-kb insert containing the B. suis ure1D, ure1A, ure1B, ure1C genes; Amp' | This study |
| pCR2.1Δ | pCR2.1Δ with 2.2-kb insert containing the B. suis ure1D, ure1A, ure1B, ure1C genes; Amp' | This study |
| pGEM-3Z | Cloning vector, 2.74-kb, Amp' | Promega |
| pGEMure1 | pGEM-3Z with 2.2-kb insert containing the B. suis ure1D, ure1A, ure1B, ure1C genes; Amp' | This study |
| pGEMure2ABC | pGEM-3Z with 2.2-kb insert containing the B. suis ure2A, ure2B, ure2C genes; Amp' | This study |
| pGEMure2ABCΔ | pGEM-3Z with 2.2-kb insert containing the B. suis ure2A, ure2B, ure2C genes; Amp' | This study |
| pUC4K | Cloning vector, 3.9-kb, Kan', Amp' | Pharmacia |
| pGEMure1K | pGEMure1 with 0.9-kb Ncol fragment deleted and blunt ended and a 1.6-kb Pvu-cut and blunt-ended Kan' cassette from pUC4K ligated, Kan', Amp' | This study |
| pGEMure2ABCΔ | pGEMure2 with 0.6-kb SacI fragment deleted and blunt ended and a 1.6-kb Pvu-cut and blunt-ended Kan' cassette from pUC4K ligated, Kan', Amp' | This study |
| pGEMure2ABCΔ | pGEMure2 with 1.2-kb ClaI and MfeI fragment deleted and blunt ended and a 1.7-kb EcoS2I plus KpnI-cut and blunt-ended Cm' cassette from pBBR4MCS ligated, Cm', Amp' | This study |
| pBRR4MCS | Broad-host-range vector; Cm' | [27] |
| Strains | | |
| Escherichia coli | Top10 | F- mcrA (mrn-hsdRMS-mcrBC) Δ(lac-proAB) leuB66 lacY1 galK116 lacZΔM15 ΔlacIq X74 deoR recA1 araD139 (rha-leu) 7697 galU galK rpsL (StrR) endA1 nupG | Invitrogen |

B. suis

| Strain | Description | Source or reference |
|--------|-------------|---------------------|
| 1330   | Parent-type, smooth strain | G.G. Schurig |
| 1330Δure1K | ure1A deleted mutant of 1330, Kan' | This study |
| 1330Δure2K | ure2BC deleted mutant of 1330, Kan' | This study |
| 1330Δure2C | ure2ABC deleted mutant of 1330, Cm' | This study |
| 1330Δure1KΔure2C | ure1A and ure2ABC deleted mutant of 1330, Cm', Kan' | This study |
Table 5: Identity of B. suis urease-1α and urease-2α sequences with the urease α subunits or urease proteins in GenBank

| Accession number | Organism                          | Identity% |
|------------------|-----------------------------------|-----------|
| YP_221060.1      | Brucella abortus                  | 99%       |
| NP_540569.1      | Brucella melitensis               | 99%       |
| NP_105696.1      | Mesorhizobium loti                | 81%       |
| AAL83830.1       | Rhizobium leguminosum             | 78%       |
| NP_386576.1      | Sinorhizobium meliloti            | 79%       |
| NP_355353.1      | Agrobacterium tumefaciens         | 78%       |
| YP_166953.1      | Silicibacter pomeroyi             | 75%       |
| EAQ05022.1       | Oceanicola baltensis              | 75%       |
| BAB21067.1       | Rhodobacter capsulatus            | 75%       |
| ZP_01056362.1    | Roseobacter sp.                   | 73%       |
| ZP_00962028.1    | Sulfitobacter sp.                 | 73%       |
| YP_109255.1      | Burkholderia pseudomallei         | 70%       |
| AAA25151.1       | Klebsiella Aerogenes              | 68%       |
| NP_0090659.1     | Vibrio splendidus                 | 69%       |
| NP_286680.1      | Escherichia coli                  | 67%       |
| NP_886007.1      | Bordetella parapertussis          | 66%       |
| ZP_00504504.1    | Clostridium thermocellum          | 63%       |
| NP_176922.1      | Arabidopsis thaliana              | 64%       |
| YP_248248.1      | Haemophilus influenzae            | 62%       |
| NP_391545.1      | Bacillus subtilis                 | 62%       |
| ZP_00133792.2    | Actinobacillus Pleuroptunmoniae   | 62%       |
| ABC74584.1       | Yersinia enterocolitica           | 57%       |
| AAK32714.1       | Helicobacter pylori               | 59%       |
| NP_336355.1      | Mycobacterium tuberculosis        | 62%       |
| AAZ99164.1       | Streptococcus vestibularis        | 57%       |
| AAQ85883.1       | Glycine max                       | 63%       |
| BAB78715.1       | Oryza sativa                      | 62%       |
| XP_750204.1      | Aspergillus fumigatus             | 62%       |

With urease-2α

| Accession number | Organism                          | Identity% |
|------------------|-----------------------------------|-----------|
| NP_539564.1      | Brucella melitensis               | 99%       |
| YP_222047.1      | Brucella abortus                  | 99%       |
| ZP_00828648.1    | Yersinia frederiksenii            | 69%       |
| ZP_00797116.1    | Yersinia pestis (and all species of Yersinia) | 69% |
| NP_929433.1      | Photobacter luminescens           | 67%       |
| NP_929433.1      | Photobacter luminescens           | 67%       |
| NP_241120.1      | Bacillus halodurans               | 59%       |
| YP_237504.1      | Pseudomonas syringae              | 59%       |
| NP_391545.1      | Bacillus subtilis                 | 57%       |
| YP_368247.1      | Burkholderia sp.                  | 56%       |
| ZP_00612021.1    | Mesorhizobium sp.                 | 57%       |
| NP_333073.1      | Agrobacterium tumefaciens         | 57%       |
| YP_295218.1      | Ralstonia eutropha                | 56%       |
| EAQ70376.1       | Synechococcus sp.                 | 57%       |
| IO5696.1         | Mesorhizobium loti                | 56%       |
| YP_248248.1      | Haemophilus influenzae            | 56%       |
| NP_997995.1      | Bacillus cereus                   | 56%       |
| NP_440403.1      | Synechocystis sp.                 | 56%       |
| AAG52306.1       | Arabinodipolis thaliana           | 57%       |
| ZP_00133792.2    | Actinobacillus Pleuroptunmoniae   | 55%       |
| ZP_00990659.1    | Vibrio splendidus                 | 56%       |
| NP_286680.1      | Escherichia coli                  | 56%       |
| YP_204056.1      | Vibrio fischeri                   | 55%       |
| AAP51176.1       | Helicobacter pylori               | 55%       |
| BAB78715.1       | Oryza sativa                      | 56%       |
| AAR21273.1       | Streptococcus thermophilus        | 54%       |
| NP_336355.1      | Mycobacterium tuberculosis        | 55%       |
| AAC46128.1       | Bordetella bronchiseptica         | 52%       |
| XP_658035.1      | Aspergillus nidulans              | 54%       |
Plasmid DNA was isolated using either plasmid Mini- or Midiprep purification kits (QIAGEN). Restriction digests, Klenow reactions, and ligations of DNA were performed as described elsewhere [24]. Restriction enzymes, Klenow fragment, and T4 DNA ligase enzyme were purchased from Promega Corporation (Madison, WI). Ligated plasmid DNA was transformed into E. coli Top10 cells by heat shock per the guidelines of the supplier (Invitrogen). Plasmid DNA was electroporated into B. suis with a BTX ECM-600 electroporator (BTX, San Diego, CA), as described previously [25].

### Mutation of ure1 operon

A 2,241-bp region including the whole length of the ure1A, ure1B, and ure1C genes and a portion of ure1C gene (Figure 1) was amplified via PCR using the genomic DNA of B. suis strain 1330 and the primers UreaseONE-Forward and UreaseONE-Reverse (RansomHill Bioscience, Inc., Ramona, CA) (see Table 6). The amplified gene fragment was cloned into the pCR2.1 vector to produce plasmid pCRure1. From this plasmid the ure1 region was isolated by EcoRI digestion and cloned into pGEM-3Z (Promega) and the resulting 5.0-kb plasmid was designated pGEMure1. The suicide vector pGEMure1K was constructed as follows: the plasmid pGEMure1 was digested with Ncol to delete a 575-bp region from the ure1 region. The Ncol sites on the 4.4-kb plasmid were filled in by reaction with Klenow enzyme and ligated to the 1.6-kb PvuII fragment of pUC4K (also blunt ended) containing the Tn903 npt gene [26], which confers kanamycin resistance (Kan') to B. suis. The resulting suicide vector was designated pGEMure1K. The E. coli Top10 cells carrying the recombinant plasmid were picked from TSA plates containing ampicillin (100 μg/ml) to determine whether a single- or double-crossover event had occurred. Five of the colonies did not grow on ampicillin-containing plates, suggesting that a double-crossover event had occurred. PCR with the primers UreaseONE-Forward and UreaseONE-Reverse (see Table 6) confirmed that a double-crossover event had taken place in all five transformants. One of these strains was chosen for further analyses and designated 1330Δure1K.

### Mutation of ure2B and ure2C

A 2,214-bp region including the whole length of the ure2A and ure2B genes, and a portion of the ure2C gene was amplified via PCR using the primers UreaseTWO-Forward and UreaseTWO-Reverse (see Table 6). The amplified gene fragment was cloned into the pCR2.1 vector to produce plasmid pCRure2ABC. From this plasmid, the ure2ABC region was isolated by BamHI and Xbal digestion and cloned into the same sites of plasmid pGEM-3Z (Promega). The resulting 5.0-kb plasmid was designated pGEMure2ABC. The suicide vector pGEMure2ABCK was constructed as follows: the plasmid pGEMure2ABC was digested with SacI to delete a 940-bp region from the ure2ABC region, sticky sites filled in with Klenow enzyme, and ligated to the 1.6-kb PvuII fragment of pUC4K. The resulting suicide vector was designated pGEMure2ABCX. One microgram of pGEMure2ABC was used to electroporate B. suis strain 1330, and a transformed strain containing a double-crossover event was verified by PCR and designated 1330Δure2K.

### Mutation of ure2A, ure2B and ure2C

A 2,923-bp region including the whole length of ure2A and ure2B genes, and a portion of ure2C gene (Figure 1) was amplified via PCR using the primers Ure-2-AB-Forward and Ure-2-AB-reverse (see Table 6). The amplified gene fragment was purified using a Qiagen PCR purification kit (Qiagen), digested with BamHI and Xbal, cloned into the same sites of plasmid pGEM-3Z (Promega) to produce the 5.7-kb plasmid pGEMure2ABC-2. The suicide

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**Table 6: Sequences of primers (5' to 3') used to amplify ure genes**

| Primer name          | Sequence (5' to 3') |
|----------------------|---------------------|
| UreaseONE-Forward    | CGACGCCTAGGTAATTC   |
| UreaseONE-Reverse    | CTAGTTAGCCGCTGCTG   |
| UreaseTWO-Forward    | GCTTGCCCTGATTTGCTG  |
| UreaseTWO-Reverse    | ATCTGCGAATTTGCCGA   |
| Ure-2-AB-Forward     | CGGGGATCCCATCAATCG  |
| Ure-2-AB-reverse     | CGGTCTAGAATGGCGGA   |

(Continued from the previous page)
vector pGEM\textit{ure}2\textit{ABCC} was constructed as follows: the plasmid pGEM\textit{ure}2ABCC-2 was digested with \textit{C}la\textit{i} and \textit{M}fe\textit{l} to delete a 1215-bp region from the \textit{ure}2\textit{ABC} region, and the ends were filled in with Klenow enzyme. The 1.7-kb gene encoding resistance to chloramphenicol (Cm\textsuperscript{r}) was isolated by digesting the plasmid pBBR1MCS [27] with \textit{Eco}52\textit{I} and \textit{K}pn\textit{l}, and the ends were filled in with Klenow enzyme. The larger fragment of the plasmid pGEM\textit{ure}2ABC-2 was ligated with the Cm\textsuperscript{r} gene, to make the suicide vector pGEM\textit{ure}2ABCC. One ug of pGEM\textit{ure}2ABC was used to electroporate \textit{B. suis} strain 1330 and the transformants were picked from TSA plates containing chloramphenicol (30 \text{mg/mL}). A transformed \textit{B. suis} strain with a double-crossover event was verified by PCR and designated 1330\textit{\text{Δ}ure}2\textit{C}.

**Generation of an \textit{ure}1, \textit{ure}2\textit{strains}**

One ug of suicide vector pGEM\textit{ure}2ABCC was used to electroporate mutant \textit{B. suis} strain 1330\textit{\text{Δ}ure}1\textit{K}. The transformants were picked from TSA plates containing kanamycin (100 \text{μg/mL}) plus chloramphenicol (30 \text{μg/mL}). A transformant \textit{B. suis} strain with a double-crossover event was verified by PCR and designated 1330\textit{\text{Δ}ure}1\textit{K}\textit{Δure}2\textit{C}.

**Real-time PCR assays**

RNA was isolated from broth cultures of \textit{B. suis} strain 1330 by the procedure described previously [31]. After a 75% ethanol wash, the dried RNA pellet was resuspended in RNase- and DNase-free water (Sigma). The concentration of the RNA was be determined with the RiboGreen RNA Quantitation kit (Molecular Probes). Genomic DNA was digested with RNase-free DNase (Ambion), and precipitated with GlycoBlue (Ambion). RNA samples not treated with reverse transcriptase was also subjected to precipitation with GlycoBlue (Ambion). The concentration of protein in the extracts was determined using a Bradford-modified assay [34]. All assays (final volume of 2 mL) were performed in 31 mM Tris-HCl pH 8.0 buffer at 28°C, using a Beckman DU 800 UV/Vi spectrophotometer, with a stirred, temperature-controlled multi-cell holder. Urease activity was determined using a coupled assay with glutamate dehydrogenase [35]. Glutamate dehydrogenase 12 U/mL was incubated with 250 μM NADPH for 5 min. 2-Oxoglutarate (1 mM) and \textit{B. suis} extracts were added and the reaction was followed at 340 nm. The observed decrease in absorbance monitored during this period is due to nonspecific oxidation of NADPH by several enzymes in the extract. The absorption coefficient used for the internal standard of \textit{ure} genes were purchased from Sigma-Genosys. The target cDNA was normalized internally to the sigA cDNA level in the same sample [32,33].

**Growth rates of \textit{B. suis}**

Single colonies of strains 1330, 1330\textit{\text{Δ}ure}1\textit{K}, 1330\textit{\text{Δ}ure}2\textit{K}, 1330\textit{\text{Δ}ure}2\textit{C} and 1330\textit{\text{Δ}ure}1\textit{K}\textit{\text{Δ}ure}2\textit{C} were grown at 37°C for 72 h to stationary phase in 5 ml of TSB. These cultures were used to inoculate 25 ml of TSB in a Klett side-arm flask to 12 to 16 Klett units. Cultures were grown at 37°C at 200 rpm; Klett readings were recorded every 2 h in a Klett-Summerson colorimeter (New York, NY).

**Native polyacrylamide gel electrophoresis**

Extracts were prepared from \textit{B. suis} strains 1330,1330\textit{\text{Δ}ure}1\textit{K}, 1330\textit{\text{Δ}ure}2\textit{K}, 1330\textit{\text{Δ}ure}2\textit{C} and 1330\textit{\text{Δ}ure}1\textit{K}\textit{\text{Δ}ure}2\textit{C} using glass beads and vortex, in Tris-HCl 30 mM, pH 8.0). Buffer without SDS or mercaptoethanol were added to the extracts and the extracts were loaded into the gel that did not contain SDS. After running was complete, the gel was placed in 0.02% cresol red-0.1 EDTA, washed several times until it became yellow, and incubated with 1.5% urea at room temperature until pink bands appeared, i.e., the positive urease reaction.

**Urease enzyme activity of strains-qualitative**

Fifty μl of culture (grown for 72 h in TSB) volumes of strains 1330, 1330\textit{\text{Δ}ure}1\textit{K}, 1330\textit{\text{Δ}ure}2\textit{K}, 1330\textit{\text{Δ}ure}2\textit{C} and 1330\textit{\text{Δ}ure}1\textit{K}\textit{\text{Δ}ure}2\textit{C} were used to inoculate 5 ml volumes of urease test broth (Difco). The contents were incubated at 37°C with 200 rpm shaking. At 8, 24 and 48 hours after incubation, the cultures were centrifuged to remove the cells. The color change in the clarified urease test broth was measured using a Klett-Summerson colorimeter. The native color of the urease test broth was used as a blank.

**Urease enzyme activity of strains-quantitative**

The Coomassie Brilliant Blue G, TRIS, NADPH, 2-oxoglutarate and glutamate dehydrogenase (from bovine liver) were from Sigma-Aldrich. Urea was obtained from Qia-gen. All other reagents were of analytical grade.

The concentration of protein in the extracts was determined using a Bradford-modified assay [34]. All assays (final volume of 2 mL) were performed in 31 mM Tris-HCl pH 8.0 buffer at 28°C, using a Beckman DU 800 UV/Vis spectrophotometer, with a stirred, temperature-controlled multi-cell holder. Urease activity was determined using a coupled assay with glutamate dehydrogenase [35]. Glutamate dehydrogenase 12 U/mL was incubated with 250 μM NADPH for 5 min. 2-Oxoglutarate (1 mM) and \textit{B. suis} extracts were added and the reaction was followed at 340 nm. The observed decrease in absorbance monitored during this period is due to nonspecific oxidation of NADPH by several enzymes in the extract. When the absorbance was stable, urea (10 mM) was added and the decrease in absorbance at 340 due to urease activity was measured for 5 min. Initial rates were calculated from the linear portion of the curves, by linear regression using the least squares method. The absorption coefficient used for NADPH was 6.22 M\textsuperscript{-1}cm\textsuperscript{-1} [36]. The volume of extracts was varied and the specific activity of urease was calculated. One unit of urease activity was defined as the amount of enzyme that hydrolyzes 1 μmol of urea per min. Specific activities were calculated as units of urease per mg of protein in the extract.
**Preparation of B. suis infection inocula**

TSA plates were inoculated with single colonies of *B. suis* strains. After 4 days of incubation at 37°C with 5% CO₂, the cells were harvested from plates, washed with phosphate-buffered saline (PBS), resuspended in 20% glycerol, and frozen at -80°C. The number of viable cells or cfu was determined by counting after spreading of dilutions of the cell suspensions on TSA that were incubated at 37°C with 5% CO₂. The cultures from these were used to inoculate mice and macrophages as described below.

**Persistence of recombinant B. suis strains in macrophages**

The mouse macrophage-like cell lines J774 and H36.12a [Pixie 12a] were obtained from the American Type Culture Collection (Manassas, VA). The macrophage cells were seeded at a density of 5 × 10⁵/ml in Dulbecco’s modified essential medium (DMEM) (Sigma-Aldrich) into 24-well tissue culture dishes and cultured at 37°C with 5% CO₂ until confluent. The tissue culture medium was removed, 200 μl (10⁶ cells) of the bacterial suspension in PBS was added, and the cells were incubated at 37°C for 4 h. The suspension above the cell monolayer was removed, and the macrophages washed three times with PBS. One milliliter of DMEM containing 25 μg of gentamicin was added, and the cells were incubated for 48 h at 37°C. At various time points (0, 1, 4, 24, and 48 h of incubation), the growth medium was removed, the cells were washed with PBS, and 500 μl of 0.25% sodium deoxycholate was added to lyse the infected macrophages. After 5 min the lysate was diluted in PBS, and the number of *B. suis* cfu was determined after growth at 37°C with 5% CO₂ for 72 h on TSA. Triplicate samples were taken at all time points, and the assay was repeated two times.

**Survival of B. suis strains inoculated intraperitoneally**

The Animal Care Committee of the Virginia Polytechnic Institute and State University approved the procedures used in handling research animals. Six-week-old female BALB/c mice (Charles River Laboratories, Wilmington, MA) were allowed 1 week of acclimatization. Groups of seven or eight mice each were intraperitoneally injected with 4.1 to 5.2 log₁₀ cfu of *B. suis* strains 1330, 1330Δure1K, 1330Δure2K, 1330Δure2C and 1330Δure1KΔure2C, with or without supplementation of 10 mM urea in approximately 0.5 ml PBS. Mice were sacrificed 1 week after inoculation, and the *Brucella* cfu per spleen or per liver was determined as described previously [23].

**In-vitro pH sensitivity of strains**

The pH of the phosphate buffered saline (PBS) was adjusted to 2, 4 or 7 by adding 1N HCl. The urea concentration of the PBS was adjusted to 5, 10, or 20 mM by adding urea. The PBS with different pH and urea contents were inoculated with 8.0 log₁₀ cfu of strains 1330, 1330Δure1K, 1330Δure2K or 1330Δure2C, and incubated at 37°C. At the end of 90 min incubation, serial dilutions of each culture were plated on TSA. The number of cfu on plates was determined after 4 days of incubation at 37°C with 5% CO₂.

**Data analyses**

The Student t test was performed in the analysis of cfu data in the macrophage study. The cfu data from the splenic and liver clearance assays, and *in vitro* pH sensitivity study were analyzed by performing analysis of variance. The mean cfu counts among treatments were compared using the least-significance pair-wise comparison procedure [29].

**Authors’ contributions**

AB and VD constructed mutant strains, and carried out part of growth and splenic clearance assays. AC carried out part of splenic and hepatic clearance assays and acid tolerance assays. SP and PR carried out macrophage assays. GS, NS and SB conceived of the study and participated in its design and coordination. AC-R carried out native gel electrophoresis, and AM performed quantitative urease activity.

**Additional material**

### Additional File 1

**Identity of B. suis urease-1α and urease-2α sequences with the urease α subunits or urease proteins in GenBank.**

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### Additional File 2

**Sequences of primers (5’ to 3’) used to amplify ure genes.**

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