Two Plant Bacteria, *S. meliloti* and *Ca. Liberibacter asiaticus*, Share Functional *znuABC* Homologues That Encode for a High Affinity Zinc Uptake System

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Abstract

The Znu system, encoded for by *znuABC*, can be found in multiple genera of bacteria and has been shown to be responsible for the import of zinc under low zinc conditions. Although this high-affinity uptake system is known to be important for both growth and/or pathogenesis in bacteria, it has not been functionally characterized in a plant-associated bacterium. A single homologue of this system has been identified in the plant endosymbiont, *Sinorhizobium meliloti*, while two homologous systems were found in the destructive citrus pathogen, *Candidatus Liberibacter asiaticus*. To understand the role of these protein homologues, a complementation assay was devised allowing the individual genes that comprise the system to be assayed independently for their ability to reinstate a partially-inactivated Znu system. Results from the assays have demonstrated that although all of the genes from *S. meliloti* were able to restore activity, only one of the two *Ca. Liberibacter asiaticus* encoded gene clusters contained genes that were able to functionally complement the system. Additional analysis of the gene clusters reveals that distinct modes of regulation may also exist between the *Ca. Liberibacter asiaticus* and *S. meliloti* import systems despite the intracellular-plant niche common to both of these bacteria.

Introduction

The ability to import zinc is critical for many species of bacteria because of its use as an enzymatic cofactor or structural element in many cellular proteins [1]. Since zinc is both highly charged and hydrophilic, it cannot cross the bacterial membrane via passive diffusion [2] and must gain entrance into the bacterial cells through the actions of either non-specific or specific transport systems [3]. An example of each of the two types of systems for zinc import is that encoded by *pit* and *znu*, respectively [4]. The non-specific *pit* system is constitutively expressed and has a low affinity for zinc in contrast to the specific transport system encoded by the *znu* genes.

The Znu (zinc uptake) system has a high affinity for zinc and was discovered in *Escherichia coli* during a transposon screen with lacZ gene cluster fusions [5]. This system has since been classified as a member of the ATP-binding cassette (ABC) transporter family and is composed of three protein products: ZnuA, ZnuB, and ZnuC. ZnuA functions as a metallochaperone and is responsible for binding zinc within the periplasm. ZnuB acts as an integral membrane permease and works in concert with ZnuC. ZnuC is the ATPase subunit of the ABC transporter, which provides energy for the transport process through ATP hydrolysis. In *E. coli*, the zinc uptake regulator, or zur, appears to bind to and repress the promoters of both *znuA* and *znuBC*, thus providing a mechanism for zinc regulation within the cell [5]. Although the Znu system has been shown to be important for both growth and/or pathogenesis in several bacterium such as *Synechocystis* [6], *Pasteurella multocida* [7], and *Salmonella enterica* [8], it has not been characterized in plant-associated bacteria.

*Sinorhizobium meliloti* and *Candidatus Liberibacter asiaticus* are two closely related members of the Rhizobiaceae family with drastically different genome sizes [9] that can be found living in both similar and distinct ecological niches. *S. meliloti* is both a free-living soil microorganism and a plant-associated bacterium, which is known for producing nitrogen-fixing nodules on leguminous plants (for a review see [10]). The symbiosis takes place under nitrogen limiting conditions when the bacteria are internalized by the plant's roots and are subsequently able to convert atmospheric nitrogen into ammonia, which then acts as a source of nitrogen for the plant. In 2001, the completed sequence of both the chromosome and the two megaplasmids of *S. meliloti* became publicly available [11,12,13]. Since that time, tools allowing genome-wide screening have been developed [14,15,16] in an effort to understand the symbiotic relationship between the plant and the bacterium on a molecular level.

*Ca. Liberibacter asiaticus* is one of three different species of Liberibacter that cause the devastating disease of citrus plants known as huanglongbing (HLB) or citrus greening (for a review see [17,18]). This fastidious bacterium resides within the sieve tube elements [19] and is transmitted by Asian citrus psyllids (*Diaphorina citri*) that have fed on the phloem of infected plants. Symptoms
characteristic of HLB disease include blotchy mottle, vein corking, and leaf chlorosis in addition to other symptoms that resemble those of a zinc nutritional deficiency. Although no adequate control measures are currently available for this disease, the recent publication of the Ca. Liberibacter asiaticus genome sequence may provide insight into areas amenable to the development of novel control strategies for HLB disease [9].

In this report we have characterized the znu gene cluster(s) and their involvement in the high-affinity uptake of zinc in two closely related plant-associated intracellular bacteria, S. meliloti and Ca. Liberibacter asiaticus. The putative genes comprising the Znu system of S. meliloti along with that of Ca. Liberibacter asiaticus were tested individually for their ability to restore the high-affinity zinc uptake system in knock-out strains of both E. coli and S. meliloti. In addition, a bioinformatic analysis of the gene clusters was used to uncover putative modes of regulation for the import systems.

Materials and Methods

Bacterial Strains and Media

S. meliloti strain Sm1021, which was obtained from Brenda K. Schroeder at Washington State University, was grown at 28°C in either Luria-Bertani (LB) broth, or in tryptone-yeast (TY) media. The wild-type E.cherichella coli strain BW25113, along with its Δznu derivatives JW1847-1, JW1848-1, and JW5831-1 [20] were obtained from the E. coli Genetic Stock Center at Yale University (Table 1). E. coli Top10 cells (Invitrogen, Carlsbad, CA) were used as a host for plasmid construction. E. coli strains were grown at 37°C in either LB broth, on LB agar plates, or in TY media supplemented with the following concentrations of antibiotics: 50 μg/mL of ampicillin, and 50 μg/mL of kanamycin.

Sequence Analysis

The percent sequence identity was determined using the ClustalW method in the program AlignX (Vector NTI Advance 11.0). Signal sequences were predicted by both SignalP 3.0 (http://www.cbs.dtu.dk/services/SignalP/) and SIG-Pred (http://mbpvc36.leece.ac.uk/prot_analysis/Signal.html). The ProtParam tool on the ExPASy server was used to predict both the molecular weight and isoelectric point of the proteins [21]. The HMMTOP server (http://www.enzim.hu/hmmtop/index.html) was used to predict the transmembrane topology of the proteins.

Reverse Transcription PCR

The operon structure of the Znu system in S. meliloti 1021 was determined using reverse transcription PCR. Total RNA was extracted from the bacterial cultures grown at 25°C for 72 hours (hrs) in LB broth (Sigma, St. Louis, MO) using TRI Reagent (Sigma) according to the manufacture’s protocol. Subsequent to RNA isolation, each sample was treated with RQ1 DNase (Promega, Madison, WI) to remove any contaminating DNA and verified DNA free by PCR. Two step reverse transcription PCR (RT-PCR) analyses were performed with gene specific primers, which were designed to span the junction between the genes (Table S1 and Fig. 1), using the GoScript Reverse Transcription System (Promega) per manufacturer’s instructions. The PCR profile consisted of a 3 min. denaturation at 95°C, followed by 40 cycles of 95°C for 30 sec., 57°C for 30 sec., and 72°C for 1 min. The final PCR step was an incubation period of 5 min at 72°C before being held at 4°C. Genomic DNA extracted from S. meliloti 1021 was used as a positive control for the PCR reactions. Amplicon products were electrophoresed on 2% agarose gels and visualized with ethidium bromide staining. A Logic 200 Imaging System from Kodak with Kodak Molecular Imaging Software v. 4.0.5 (Eastman Kodak Company, Rochester, NY) was used for gel image capture.

DNA Manipulations

PCR amplification of the znu genes for cloning was performed with Platinum Taq High Fidelity Polymerase (Invitrogen), while all subsequent PCR amplifications were performed with Taq DNA Polymerase (New England Biolabs, Ipswich, MA). Resulting DNA products were purified from agarose gels using the QiAquick gel extraction kit (Qiagen, Valencia, CA), while restriction fragments were purified with the QiAquick PCR purification system (Qiagen). Genomic DNA was extracted from E. coli strain BW25113 by boiling a sample from an overnight culture, pelleting the cell debris, and using the soluble extract for PCR. Plasmid DNA was isolated from E. coli cultures using the QIAprep Spin Miniprep kit (Qiagen). DNA sequencing was done by the U.S. Horticulture Research Laboratory Core Facility using BigDye version 3.1 and the 3730×lDNA analyzer (Applied Biosystems, Carlsbad, CA).

Construction of Plasmids Containing the znuA, znuB or znuC Genes from Ca. Liberibacter Asiaticus, E. coli, and S. meliloti

Individual genes from the two znuABC gene clusters in Ca. Liberibacter asiaticus, S. meliloti, and E. coli were placed on a plasmid under the control of the tetracycline inducible promoter as follows: each znu gene was amplified from Ca. Liberibacter asiaticus genomic DNA isolated from Ca. Liberibacter asiaticus-infected psyllids [22] using primers specific for the gene of interest (Table S1), while the genes from E. coli were amplified from BW25113 genomic DNA and those from S. meliloti were amplified with gene specific primers from plasmids pESMc04243, pESMc04243, and pESMc04244, respectively [16]. The amplified products were purified on 1% agarose gels and both the PCR products and the vector, pASK-IBA3 (IBA Biotangnology, Goettingen, Germany), were subsequently digested with BsaI (New England Biolabs) using the engineered restriction sites. T4 DNA ligase (New England Biolabs) was used to ligate the resulting fragments to the vector producing pzmAu05, pzmAu12, pzmBu05, pzmBu12, and pzmCu12, pzyBaLe pzyBeL, pzcM, pCMV219, pCMV220, and pCMV221 (Table 1). The clones were verified via DNA sequencing and transformed into the corresponding E. coli znu knock-out strains.

Complementation Assay of Δznu E. coli Strains with Tetracycline-inducible Gene Expression Plasmids

Growth curves for the complementation assay of the Δznu E. coli strains were produced by using 5 μL of a culture grown overnight in TY to inoculate 10 mL of fresh TY or TY containing 0.4 mM EDTA. Media for complementation assays also contained 50 μg/mL ampicillin, 50 μg/mL kanamycin, and 50 μg/L anhydrotetracycline (AHT) for the induction of the znu genes on the complementing plasmids. Growth was measured at the time points indicated using a Biomate 3 spectrophotometer (Thermo Electron Corp.) at an absorbance of 600 nm (OD600).

Growth of Wild-type E. coli and S. meliloti in EDTA with and without Zinc Repletion

Growth inhibition studies were conducted on wild-type E. coli and S. meliloti cultures by growing them in TY media containing an increasing concentration of EDTA (pH 8.0), ranging from 0 mM to 3.2 mM. These cultures were inoculated with a 1:100 dilution of an overnight growth of the wild-type strains. After
approximately 20 hrs of growth, the OD_{600} of these strains were measured and recorded.

For zinc repletion experiments, wild-type *S. meliloti* 1021 cultures were grown in TY media containing 0, 0.1, 0.2, or 0.4 mM of EDTA (pH 8.0). Different concentrations of ZnSO_{4} wa... ranging from 0 mM to 0.5 mM, was added to the culture media in order to replete the zinc that was chelated by the EDTA. Cultures were inoculated and the growth recorded as above.

**Generation of *S. meliloti* znu Deletion Mutants**

Partial sequences of the three *znu* genes of *S. meliloti* were amplified from genomic DNA using the forwards (For) and reverse (Rev) primer pairs *ZnuA*.M, *ZunB*.M and *ZunC*.M, respectively. The resulting PCR products were ligated into pCR2.1 (Invitrogen), digested with *EcoRI*, and ligated into the mobilizable vector pK19mobGII [23]. The constructs were transformed into DH5α, and subsequently introduced into wild type *S. meliloti* 1021 by triparental mating with the helper pRK2073 for mutant isolation [24]. For complementation analysis, *znu* genes from *S. meliloti* and *Ca. Liberibacter asiaticus* were cloned into cosmids pBRR1MC-5 [25], and introduced into the corresponding mutant by triparental mating. The ability of each strain to transport zinc was tested as described above except the OD_{600} of the cultures were measured and recorded after 48 hrs of growth at 28°C.

**Results**

**Structure of the Znu Gene Cluster in *S. meliloti* and *Ca. Liberibacter asiaticus***

The annotation of the *S. meliloti* genome sequence reveals a putative gene cluster with some similarities to the high-affinity zinc uptake system (*znuABC*) in *E. coli*. The arrangement of the genes in the *S. meliloti* cluster resembles that of *E. coli* with the exception of the zinc uptake regulator, *zur*, which is located immediately

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**Table 1. Plasmids and strains used in this study.**

| Name             | Relevant Characteristic(s)                                                                 | Reference |
|------------------|------------------------------------------------------------------------------------------|-----------|
| BW25113          | Wild-type *E. coli* strain used in the construction of the Δznu strains                    | [20]      |
| DH5α             | Strain used for cloning of constructs                                                     | Invitrogen (Carlsbad, CA) |
| JW1847-1         | ΔznuC derivative of BW25113 strain                                                       | [20]      |
| JW1848-1         | ΔznuB derivative of BW25113 strain                                                       | [20]      |
| JW5831-1         | ΔznuA derivative of BW25113 strain                                                       | [20]      |
| Sm1021           | Wild-type *S. meliloti* 1021 strain                                                      | (Meade, 1982) |
| pASK-IBA3        | Expression plasmid containing the tetracycline promoter                                   | IBA (Gottingen, Germany) |
| pBRR1MC-5        | Broad host range cloning cosmid                                                           | [25]      |
| pBB.La           | *znuA* gene from *Ca. Liberibacter asiaticus* gene cluster 1 in the broad host range cloning cosmid | This study |
| pBB.Lb           | *znuB* gene from *Ca. Liberibacter asiaticus* gene cluster 1 in the broad host range cloning cosmid | This study |
| pBB.Lc           | *znuC* gene from *Ca. Liberibacter asiaticus* gene cluster 1 in the broad host range cloning cosmid | This study |
| pBB.Ra           | *znuA* gene from *S. meliloti* 1021 in the broad host range cloning cosmid                | This study |
| pBB.Rb           | *znuB* gene from *S. meliloti* 1021 in the broad host range cloning cosmid                | This study |
| pBB.Rc           | *znuC* gene from *S. meliloti* 1021 in the broad host range cloning cosmid                | This study |
| pCMV219          | *znuA* gene from *S. meliloti* 1021 under the inducible tetracycline promoter             | This study |
| pCMV220          | *znuB* gene from *S. meliloti* 1021 under the inducible tetracycline promoter             | This study |
| pCMV221          | *znuC* gene from *S. meliloti* 1021 under the inducible tetracycline promoter             | This study |
| pCR2.1           | TOPO TA cloning vector                                                                   | Invitrogen (Carlsbad, CA) |
| pESMo04243       | Plasmid containing the *znuB* gene from *S. meliloti* 1021                              | [16]      |
| pESMo04244       | Plasmid containing the *znuC* gene from *S. meliloti* 1021                              | [16]      |
| pESMo04245       | Plasmid containing the *znuA* gene from *S. meliloti* 1021                              | [16]      |
| pK19mobGII       | Mobilizable vector used for gene replacement in *S. meliloti* 1021                      | [23]      |
| pRK2073          | Triparental mating helper plasmid                                                         | [24]      |
| pyeb.l           | *znuB* gene from *E.coli* under the inducible tetracycline promoter                      | This study |
| pyeb.l           | *znuA* gene from *E.coli* under the inducible tetracycline promoter                      | This study |
| pyeb.m           | *znuC* gene from *E.coli* under the inducible tetracycline promoter                      | This study |
| pzuuA(05)        | *znuA* gene from *Ca. Liberibacter asiaticus* gene cluster 1 under the inducible tetracycline promoter | This study |
| pzuuA(12)        | *znuA* gene from *Ca. Liberibacter asiaticus* gene cluster 2 under the inducible tetracycline promoter | This study |
| pzuuB(05)        | *znuB* gene from *Ca. Liberibacter asiaticus* gene cluster 1 under the inducible tetracycline promoter | This study |
| pzuuB(12)        | *znuB* gene #1 from *Ca. Liberibacter asiaticus* gene cluster 2 under the inducible tetracycline promoter | This study |
| pzuuB(212)       | *znuB* gene #2 from *Ca. Liberibacter asiaticus* gene cluster 2 under the inducible tetracycline promoter | This study |
| pzuuC(05)        | *znuC* gene from *Ca. Liberibacter asiaticus* gene cluster 1 under the inducible tetracycline promoter | This study |
| pzuuC(12)        | *znuC* gene from *Ca. Liberibacter asiaticus* gene cluster 2 under the inducible tetracycline promoter | This study |

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E. coli regulon:  
\[ \text{znuB} \rightarrow \text{znuC} \rightarrow \text{znuA} \rightarrow \text{zur} \]

S. meliloti regulon:  
\[ \text{zur} \rightarrow \text{znuB} \rightarrow \text{znuC} \rightarrow \text{znuA} \]

Ca. L. asiaticus regulon #1:  
\[ \text{znuB}_{2} \rightarrow \text{znuB} \rightarrow \text{znuC} \rightarrow \text{znuA} \]

Ca. L. asiaticus regulon #2:  
\[ \text{znuB} \rightarrow \text{znuB} \rightarrow \text{znuC} \rightarrow \text{znuA} \]

Figure 1. Schematic comparison of the Znu gene cluster from E. coli, S. meliloti and Ca. Liberibacter asiaticus. Znu homologues have been identified in the two intracellular-plant bacterium, S. meliloti and Ca. Liberibacter asiaticus. The arrangement of znuA, znuB, and znuC genes in each bacterium are shown. A BLAST search of the completed genomes revealed that only the E. coli and S. meliloti gene clusters contain the zinc uptake regulator, zur. In E. coli, the zur gene is located 2.2 Mb downstream of znuA while in S. meliloti it is immediately upstream of znuB. Primers are depicted as small arrows and are colored coded to represent the genes in which they are located. Notations above the primers correspond to the final letters in the primer name and are colored according to the genes that they amplify. Names of primers that were used to cross the junction of a gene are listed in black.

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upstream of znuB in S. meliloti but is approximately 2.2 Mb downstream of znuB in E. coli (Fig. 1). To determine if the znuCB genes are co-transcribed in S. meliloti as they are in other bacteria [7], reverse transcription PCR was performed using RNA extracted from an axenic culture. A primer set was designed to span the junction between the genes of the putative operon (Fig. 1 and Table S1) so that a product would be seen only if a single transcript encoded the adjacent open reading frames (ORFs). In this assay, a 456 bp product was detected (Fig. 2), which corresponds to the distance between the primers on either side of the junction and indicates that znuC and znuB are indeed co-transcribed. Primer sets designed to amplify a region within each gene (583 bp for znuA, 526 bp for znuB, and 473 bp for znuC) were also used. Fragments corresponding to the individual genes were also detected, demonstrating the presence of transcript for all three genes under the conditions used.

Within the Ca. Liberibacter asiaticus genome, two putative gene clusters with similarity to znuABC of E. coli were identified. These gene clusters are located approximately 191 Kb apart on the chromosome and contain either three or four genes consisting of znuCB or znuCBB, respectively (Fig. 1). The gene arrangement of the first Ca. Liberibacter asiaticus gene cluster suggests that the znuA gene and the znuCB genes are divergently transcribed, whereas in the second Ca. Liberibacter asiaticus gene cluster, all of the znu genes appear to be co-transcribed. Because of the difficulty with its cultivation and the inability to obtain a substantial quantity of viable cells in culture [26], efforts were made to determine the operon structure of the Ca. Liberibacter asiaticus genes by extracting bacterial RNA from infected plant tissue. Despite several attempts, conclusive results could not be obtained (data not shown). Additional bioinformatic analysis using the basic local alignment search tool (BLAST) to query the complete genome of Ca. Liberibacter asiaticus did however reveal the lack of a gene within the Ca. Liberibacter asiaticus sequence that is homologous to the negative regulator, zur, from E. coli.

Sequence Analysis of Putative Znu Systems Supports Proper Structure and Localization of Protein Components

Sequence analysis of the individual components composing the Znu system supports the hypothesis that the Ca. Liberibacter asiaticus and S. meliloti proteins are functional even though the percent identity amongst species remains low (Table 2). For example, the size of the coding sequence for the znuA, znuB and znuC genes and the molecular weight of the corresponding protein products from both bacteria are comparable to that of the homologous genes/proteins in E. coli. Although the isoelectric point (pI) of most of the predicted proteins are similar to their E. coli homologues, the pI of one of the ZnuC proteins from Ca. Liberibacter asiaticus (pI of 5.8) and the ZnuC protein from S. meliloti (pI of 6.4), are vastly different from the pI of 9.4 predicted for the E. coli ZnuC. Nevertheless, all of the predicted ZnuC proteins contain both the Walker A motif (GXGXGK/ST) and Walker B motif (R/KXX_{7-8}KhhhhD) where X represents any amino acid and h represents any hydrophobic residue. These motifs are involved in ATP binding and are common to ATPases, which is indicative of their function in the ABC transporter. In addition, the HMMTOP server [27] predicts seven transmembrane helices for all of the ZnuB proteins except that of znuB from the second gene cluster in Ca. Liberibacter asiaticus, in which only six are predicted. The presence of these domains is in accordance with their predicted role as the membrane permease of the ABC transporter. Signal peptide prediction tools (SIG-Pred and SignalP 3.0) showed that ZnuA from both Ca. Liberibacter asiaticus and S. meliloti contain signal sequences, which suggests that the proteins are targeted to the periplasmic space and provides an additional line of evidence supporting the prediction that ZnuA acts as a metallochaperone in the system.

S. meliloti and Ca. Liberibacter Asiaticus znu Alleles Complement Δznu E. coli Strains

A previous study involving solute-binding, protein-dependent transporters in S. meliloti 1021 reported a 37-fold induction upon zinc limitation of the gene SMc04245 (referenced above as ZnuA) [28], providing indirect evidence that the Znu system is functional in this bacterium. However, no direct evidence pertaining to the function of ZnuA or the other Znu proteins has been reported for S. meliloti nor have any studies been published to date indicating that either of the two Ca. Liberibacter asiaticus systems is operational.

To understand the role of the Ca. Liberibacter asiaticus and S. meliloti Znu homologues, a complementation assay was devised to...
allow the individual genes that comprise the system to be assayed independently for their ability to re-establish the function of a partially-inactivated Znu system. 

**E. coli** was used as a heterologous host for these assays since a viable genetic system for **Ca. Liberibacter asiaticus** is not currently available in vitro. Here, three strains of **E. coli** containing an insertional inactivation of either **znuA**, **znuB**, or **znuC**, were transformed with a plasmid containing the respective **znu** allele from either **S. meliloti** or **Ca. Liberibacter asiaticus** (Table 1). The alleles were placed under the control of a tetracycline-inducible promoter in order to eliminate any effects that the native promoters might have on overall transcript levels. Each putative gene comprising the Znu system was then tested independently for its ability to restore the high-affinity zinc uptake system in the appropriate **E. coli** knock-out strains by placing it in an inducing media that was depleted for zinc through the addition of 0.4 mM ethylenediaminetetraacetic acid (EDTA) and measuring the growth of the culture (Fig. 3). The vector backbone (black lines/symbols) was transformed into the individual strains for use as a negative control as well as the respective native **E. coli** alleles (orange lines/symbols), which were used as positive controls. All strains reached an OD600 > 1.5 when grown in TY (Fig. 3, solid lines) although the lag phase was consistently longer for the

Table 2. Bioinformatic analysis of the components comprising the Znu gene clusters.

| Organism          | Gene  | GI Number* | Length (bp) | Molecular Weight (KDa) | Identity% (%) | pI    |
|-------------------|-------|------------|-------------|------------------------|----------------|-------|
| **E. coli** K-12  | **znuA** | 87081990   | 933         | 33.3                   | 100            | 5.6   |
|                   | **znuB** | 1788166    | 786         | 27.7                   | 100            | 8.7   |
|                   | **znuC** | 1788165    | 756         | 27.9                   | 100            | 9.4   |
| **Ca. Liberibacter asiaticus** | **znuA** #1 | 254040394 | 885         | 33.4                   | 29             | 6.7   |
|                   | **znuB** #1 | 254040396 | 783         | 28.7                   | 39             | 8.6   |
|                   | **znuC** #1 | 254040395 | 723         | 26.7                   | 42             | 9.4   |
|                   | **znuA** #2 | 254040214 | 885         | 33.2                   | 17             | 6.3   |
|                   | **znuB** #2 | 254040216 | 822         | 30.1                   | 18             | 8.9   |
|                   | **znuC** #2 | 254040217 | 834         | 30.3                   | 17             | 8.8   |
| **S. meliloti** 1021 | **znuA** | 15074849   | 1023        | 35.9                   | 34             | 4.9   |
|                   | **znuB** | 15074847   | 828         | 29.2                   | 47             | 6.7   |
|                   | **znuC** | 15074848   | 876         | 31.6                   | 41             | 6.4   |

*In accordance with the database at the National Center for Biotechnology Information.

Percentages are based upon a ClustalW alignment between the corresponding protein in **E. coli**.

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**Figure 2. Operon structure of the znu genes in S. meliloti 1021.** Reverse transcription PCR was performed on total RNA isolated from **S. meliloti** to determine if the **znuB** and **znuC** genes were co-transcribed. Primer sets were designed to amplify fragments either internal to **znuA** (lanes 2–4), **znuB** (lanes 5–7) and **znuC** (lanes 8–10) or a fragment that span the junction between **znuB-C** (lanes 11–13). Reverse transcriptase was added to the reaction in lanes 2, 5, 8, and 11 (RT+) but omitted from lanes 3, 6, 9, and 12 (RT-) to demonstrate that the RNA was free of any detectable DNA contaminations. Genomic DNA was also amplified with the corresponding primers for use as a positive control (lanes 4, 7, 10, and 13). Experiment shown is representative of data from three independent replicates.

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strains containing znuA from gene cluster #1 of Ca. Liberibacter asiaticus (Fig. 3A, solid red line) and E. coli (Fig. 3A, solid orange line). Even though the final OD\textsubscript{600} value was not as high when strains were grown in TY +0.4 mM EDTA, strains containing znuA, znuB, and znuC from S. meliloti (Fig. 3A–C, dotted green lines) appeared to complement the growth defect seen when only the vector control was present (Fig. 3A–C, dotted black lines). These strains reached an OD\textsubscript{600} that was similar to the E. coli controls (Fig. 3A–C, dotted orange lines). Furthermore, the strain containing znuC from gene cluster #1 of Ca. Liberibacter asiaticus (Fig. 3C, dotted red line) was also able to grow to an equivalent density. Interestingly, the OD\textsubscript{600} of the strain expressing znuB from gene cluster #1 of Ca. Liberibacter asiaticus (Fig. 3B, dotted red line) was less than cultures containing the E. coli or S. meliloti alleles but more than 2.5-fold greater than the vector control strain. This strain also showed dense growth on TY plates containing 0.4 mM EDTA compared to the vector control (data not shown), suggesting that it can complement the ΔznuB phenotype. Neither of the two znuC genes from Ca. Liberibacter asiaticus (Fig. 3A, dotted red and blue lines) appeared to complement the ΔznuA strain. In addition, neither the znuB (Fig. 3B, dotted blue and brown lines) nor the znuC genes from gene cluster #2 of Ca. Liberibacter asiaticus (Fig. 3C, dotted blue line) were able to complement the system and may in fact be detrimental to the growth since the OD\textsubscript{600} of these strains were below those of the vector control.

**Growth Inhibition of E. coli and S. meliloti by EDTA**

A concentration of EDTA as low as 0.2 mM was able to inhibit the growth of wild-type S. meliloti 1021 when added to the culture media although there were no negative effects on the overall growth of the wild-type E. coli strain at this concentration (Fig. 4). This could indicate that either the S. meliloti’s Znu system is less effective than the E. coli system under low zinc conditions or, perhaps, E. coli possesses additional machinery for zinc uptake such as ZinT [29,30], a protein involved in zinc uptake that is present in E. coli but not in S. meliloti [31]. However, when the znu genes of S. meliloti are expressed in a heterologous E. coli host those strains expressing the S. meliloti genes are able to grow in 0.4 mM EDTA (Fig. 3), which contradicts the hypothesis that the S. meliloti Znu system is less effective than the E. coli system.

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**Figure 3. Growth of E. coli complemented strains.** Strains of E. coli with A) znuA, B) znuB, or C) znuC insertionally inactivated were complemented with the corresponding znu genes from either S. meliloti (green), E. coli (orange), or Ca. Liberibacter asiaticus (red, blue, and brown) or the vector control (black). Strains were grown both in the absence (solid lines) and presence (dotted lines) of 0.4 mM EDTA. The absorbance at 600 nm (OD\textsubscript{600}) was measured at the time points indicated. Graphs are representative of data from three independent experiments.

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S. meliloti and Ca. Liberibacter Asiaticus znu Alleles Complement Δznu S. meliloti Strains

Since the inhibitory concentration of EDTA varied significantly between E. coli and S. meliloti, complementation experiments were performed (Fig. 5) in knock-out S. meliloti strains (ΔznuA, ΔznuB, and ΔznuC) using the corresponding genes from the Ca. Liberibacter asiaticus gene cluster 1 or the native S. meliloti genes (the two operons that contained genes which were functional in E. coli). The results of ΔznuC S. meliloti strains paralleled those of the ΔznuC E. coli strains, with the znuC genes from both S. meliloti and Ca. Liberibacter asiaticus demonstrating the ability to complement (Fig. 5C). Unique to the S. meliloti host strains was the fact that the Ca. Liberibacter asiaticus znuB gene appeared to restore the growth to relatively the same level as the S. meliloti znuB gene at the lower concentrations of EDTA (Fig. 5B) and the fact that the Ca. Liberibacter asiaticus znuA gene now appeared to partially complement the ΔznuA S. meliloti strain at 0.1 and 0.15 mM EDTA (Fig. 5A). All of the znu genes from S. meliloti were able to fully compliment, whereas the vector alone did not, as was predicted (Fig. 5, orange bars vs. grey bars).

Growth of S. meliloti in Zinc Repleted Media

Because EDTA is known to chelate other positively charged ions in addition to zinc, the growth of S. meliloti was observed in zinc repleted media to determine if the growth inhibition was the result of zinc chelation in the media (Fig. 6). When wild-type S. meliloti 1021 was grown in TY media with various concentrations of EDTA (0, 0.1, 0.2, or 0.4 mM) that had been supplemented with different amounts of zinc (0, 0.125, 0.25, or 0.5 mM), a pattern emerged in which growth was inhibited under both high concentrations of EDTA without the addition of zinc (Fig. 6-initial white bar) and under high concentrations of zinc without the addition of EDTA (Fig. 6-final black and grey bars). Given that growth could be restored in media containing 0.4 mM EDTA by the addition of zinc demonstrates the likelihood that the chelation of zinc by EDTA resulted in the diminished growth of S. meliloti. The growth of S. meliloti upon the addition of EDTA to media containing zinc concentrations that were previously inhibitive further supports this conclusion.

Modes of Regulation in S. meliloti and Ca. Liberibacter Asiaticus

Once the function of several znu genes was established, putative modes of gene regulation were subsequently examined. A protein homologous to the Zur protein of E. coli was found immediately upstream of znuB in S. meliloti, though no homologues could be identified in the completed genome sequence of Ca. Liberibacter asiaticus using BLAST. The palindromic sequence to which Zur binds (Zur box) [31], which is commonly found within the promoter region of znuA, was also present only in S. meliloti and not in the promoter region of either znuA genes in Ca. Liberibacter asiaticus (data not shown). In organisms such as P. multocida, transcriptional regulation of znuA has been shown to be under the control of the iron-uptake regulator, or Fur, instead of Zur [7]. However, a BLAST analysis of the Ca. Liberibacter asiaticus genome suggests that it does not contain a homologue of fur either.

These data imply that the mode of regulation for znuABC could differ between Ca. Liberibacter asiaticus and S. meliloti.
Complementation of the znuA, znuB, and znuC Alleles

Complementation experiments are an established method for assessing gene function in a biological system. Using this approach, we were able to define the functional role of several putative znu alleles from both *S. meliloti* and *Ca. Liberibacter asiaticus*. Despite only a modest degree of identity amongst the orthologous proteins and the difficulties involved in the execution of such studies because of the inability to reliably cultivate *Ca. Liberibacter asiaticus*, successful complementation assays were performed in two different host strains, both *E. coli* and *S. meliloti*. As a result of these studies, the role of the znuA, znuB, and znuC alleles in zinc uptake has now been demonstrated for the *S. meliloti* and *Ca. Liberibacter asiaticus* genes tested here.

The biophysical mechanisms involved in the binding and transportation of the zinc by the Znu system has been characterized previously. Using the crystal structure, Banerjee et al. suggested that three conserved His residues were crucial for the binding of zinc in ZnuA [6]. A sequence alignment of the amino acid composition of ZnuA from *E. coli*, *S. meliloti*, and *Ca. Liberibacter asiaticus* revealed that ZnuA from the second gene cluster of *Ca. Liberibacter asiaticus* contained only two of the three conserved His residues (Fig. 7), which may be a reason for its inability to complement (Fig. 3). This apparent lack of complementation was not unexpected considering that none of the other Znu proteins from the second gene cluster of *Ca. Liberibacter asiaticus* appeared functional in the complementation assay.

Conversely, it was surprising that the ZnuA from the first *Ca. Liberibacter asiaticus* gene cluster only partially complemented the Δznu *S. meliloti* strain (Fig. 5A) and did not complement the Δznu *E. coli* strain (Fig. 3A, dotted red line) even though the other
two genes present in that gene cluster were shown to be functional and all three His residues forming the zinc binding site were present (Fig. 7). The lack of full complementation could mean the existence of a protein that works in conjunction with ZnuA to deliver zinc to membrane permease (ZnuB) or, perhaps, it is the result of the inability of ZnuA from Ca. Liberibacter asiaticus to be expressed properly/interact fully with the ZnuBC complex within S. meliloti or E. coli. This hypothesis is plausible considering ZnuA from Ca. Liberibacter asiaticus shares only 29% identity (Table 2) with E. coli (where a complete lack of complementation was observed) but 34% identity with S. meliloti (where partial complementation was obtained). Similarly, when the plasmid expressing znuB from gene cluster #1 of Ca. Liberibacter asiaticus was placed in the ΔznuB E. coli strain (Fig. 3B, dotted red line), growth was only partially restored compared to the strain containing the S. meliloti znuB gene (Fig. 3B, dotted green line), although when the same genes were placed in the ΔznuB S. meliloti strain (Fig. 5B), both the Ca. Liberibacter asiaticus gene and the S. meliloti gene appeared to restore growth to levels similar to wild type at lower concentrations of EDTA. The higher percent identity that exists between the proteins from Ca. Liberibacter asiaticus and S. meliloti compared to Ca. Liberibacter asiaticus and E. coli implicates protein-protein interactions as a just cause for these discrepancies, however, when each of the two gene clusters from Ca. Liberibacter asiaticus was transformed into the ΔznuA E. coli strain in its entirety, the strains still did not outgrow the vector control in the presence of 0.4 mM EDTA (data not shown). Because the genes were under their native promoters in this construction, it is possible that not all of the genes were transcribed. It is also possible that the ZnuA gene from the second gene cluster interacts with the ZnuBC complex from the first Ca. Liberibacter asiaticus gene cluster but not with the ZnuBC complex from E. coli, which would also yield a negative result in this assay.

It is interesting to note that the structure and composition of both Ca. Liberibacter asiaticus gene clusters are highly conserved in a related species known as Candidatus Liberibacter solanacearum (Table 3) [32], the causal agent of zebra chip disease in potatoes. If these genes were completely non-functional, the high degree of conservation seen between the two would not be likely considering the greatly reduced genome sizes of these organisms and their different host ranges. Therefore, it is likely that these genes are functional, although how they function remains undefined. It is worth mentioning that the ZnuA protein from Ca. Liberibacter asiaticus gene cluster #2 is 33% identical to the known Mn²⁺ transporter, PsaA, from Streptococcus pneumoniae (Genebank accession # AAC24470.1) [33, 34]. Moreover, because the Ca. L. asiaticus ZnuA protein from gene cluster #2 contains a Glu (Fig. 7), a residue predicted to facilitate Mn²⁺ binding [35], instead of a His at a position implicated in the binding of a zinc ligand [6, 36], it is possible that this protein is involved in the uptake of metals other than zinc.

**Growth Inhibition from Znu Expression**

The expression of several znu genes from Ca. Liberibacter asiaticus, including znuA from the first gene cluster and both znuB and znuC genes from the second gene cluster, appears to be detrimental to the growth of the E. coli cultures in which the genes are expressed since the density of these strains when grown in TY + EDTA were below those of strains containing the vector alone (Fig. 3). There are several explanations for this phenomenon. For example, over-expression of these proteins may be eliciting the death of the cell although this is not likely because the growth

Figure 6. Growth of S. meliloti in zinc repleted media. Increasing concentrations of ZnSO₄, ranging from 0 to 0.5 mM (x-axis), was added to media containing different amounts of EDTA (light grey bars: 0 mM, black speckled bars: 0.1 mM, dark grey bars: 0.2 mM, and white bars: 0.4 mM). The growth of S. meliloti in each media was determined by measuring the absorbance of the culture at 600 nm (OD600) ~20 hrs post inoculation. Data shown is an average of three independent trials with three replicates grown per trial. Error bars shown represent the standard deviations. doi:10.1371/journal.pone.0037340.g006
defect is only seen when the cells are grown in TY + EDTA and not in TY alone. In these assays, the level of expression is determined by the amount of anhydrotetracycline in the media not by the depletion of zinc through the addition of EDTA because the genes were placed under an inducible promoter. This makes the expression levels identical under the two conditions, yet the growth phenotype is only seen under the one condition. Another possibility is that these genes, whose functions do not appear to be

Figure 7. Sequence alignment of ZnuA orthologues. Orthologues to ZnuA from *S. meliloti* and *Ca. Liberibacter asiaticus* were aligned to ZnuA from *E. coli* using ClustalW. Residues that are 100% conserved amongst all orthologues are shown in bold. The three conserved His residues involved in binding zinc are shaded gray.

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Table 3. Sequence comparison of the Znu proteins from *Ca. Liberibacter asiaticus* vs. *Ca. Liberibacter solanacearum*.

| Name         | Genbank Sequence Record Number | % Identity* |
|--------------|--------------------------------|-------------|
| ZnuA #1      | ACT57190.1                      | ADRS1883.1  | 63          |
| ZnuA #2      | ACT57010.1                      | ADRS2867.1  | 65          |
| ZnuB #1      | ACT57192.1                      | ADRS1885.1  | 82          |
| ZnuB #2      | ACT57013.1                      | ADRS2870.1  | 77          |
| ZnuC #1      | ACT57191.1                      | ADRS1884.1  | 84          |
| ZnuC #2      | ACT57011.1                      | ADRS2868.1  | 77          |

*aPercentages are based upon a ClustalW alignment between the corresponding proteins.

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related to the Znu system, may be producing proteins that are toxic to the cells once EDTA has chelated the metals in the system. These proteins could work by inhibiting one of the other zinc uptake systems in E. coli, such as ZupT [37], which may be responsible for the slight increase in culture density seen with the vector control. They could also be importing substances detrimental to growth under these conditions. Further characterization of these strains is necessary in order to determine the reason for their substantial lack of growth.

Regulation of Zinc Uptake

The possibility of Ca. Liberibacter asiaticus and S. meliloti having distinct modes of regulation for the Znu system is intriguing given the close phylogenetic relationship [9] and the intracellular-plant niche common to both of these bacteria. Although the Znu system in S. meliloti appears to be regulated by Zur, alternative metal-sensing transcriptional regulators found within the cell may be able to regulate the expression of the zinc transporter genes since Ca. Liberibacter asiaticus lacks both zur and jaw, the two currently known regulators of the Znu system [5,7,38]. It is also possible, given the obligate-intracellular nature of Ca. Liberibacter asiaticus, that the zinc level within host’s cells remain consistently low so the need for regulation has diminished and znu expression has become constitutive over time. Proteins involved in cation efflux, such as the metallothionein-like protein (GI: 254780343), could then be used to protect the bacterium against metal toxicity caused by small aberrances in intercellular zinc levels. Conversely, since an evolutionary trend towards constitutive expression of the Znu system would not be a viable mechanism in S. meliloti because of its existence as both a plant intracellular bacterium and an extracellular soil-borne bacterium, regulation via Zur was retained.

Use of Znu System as a Target for HLB Control

Because zinc deficiency is already one of the most widespread deficiencies in citrus [39], possession of the Znu system by Ca. Liberibacter asiaticus and the subsequent uptake of zinc from its host may be resulting in a localized zinc deficiency in the citrus plant, which can ultimately lead to the death of the host’s plant tissues. This hypothesis is corroborated by the fact that the symptoms of plants affected by HLB often mimic those of a plant deficient in zinc and that the latter stages of HLB are characterized by twig dieback. Since Znu mutants have been shown to be attenuated for virulence in a variety of pathogens such as Haemophilus ducreyi [40], Brucella abortus [41], uropathogenic E. coli [42], and Campylobacter jejuni [43], it could be expected that mutations in this system would also affect the virulence of Ca. Liberibacter asiaticus. Thus, disruption of this system may present a viable target to not only lessen HLB symptom severity by making more free zinc available to the plant but also by providing a means to decrease the overall virulence of the bacterium itself. In addition, its lack of redundancy greatly increases the viability of such an option.

In conclusion, our findings clearly indicate a role in zinc uptake for the ZnuA, ZnuB, and ZnuC proteins from S. meliloti as well as one homologous gene cluster in Ca. Liberibacter asiaticus, providing the first conclusive evidence of this system in these plant-associated intracellular bacteria. These data also indicates that two distinct modes of regulation may exist for this high-affinity zinc uptake system despite a common intracellular-plant niche and their close phylogenetic relationship. The genetic and functional characterization of the Znu system not only provides insight into a system important for the growth and survival of these organisms but also provides a potential mechanism by which they might be manipulated.

Notes

Mention of trade names or commercial products in this publication is solely for the purpose of providing specific information and does not imply recommendation or endorsement by the U.S. Department of Agriculture.

Supporting Information

Table S1 Primers used in this study.

(DOC)

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Author Contributions

Conceived and designed the experiments: YPD CMVA.Performed the experiments: CMVA HSZ. Contributed reagents/materials/analysis tools: CMVA HSZ. Wrote the paper: CMVA YPD.

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