Disulfide Bond Oxidoreductase DsbA2 of *Legionella pneumophila* Exhibits Protein Disulfide Isomerase Activity

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Running Title: DsbA2 in *Legionella*

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ABSTRACT

The extracytoplasmic assembly of the Dot/Icm type IVb secretion system (T4SS) of

*Legionella pneumophila* is dependent on correct disulfide bond (DSB) formation catalyzed by a novel and essential disulfide bond oxidoreductase DsbA2 and not by DsbA1, a second nonessential DSB oxidoreductase. DsbA2, which is widely distributed in the microbial world, is phylogenetically distinct from the canonical DsbA oxidase and the DsbC protein disulfide isomerase (PDI)/reductase of *Escherichia coli*. Here we show that the extended N-terminal amino acid sequence of DsbA2 (relative to DsbA proteins) contains a highly conserved amino acid dimerization domain enabling the protein to form a homodimer. Complementation tests with *E. coli* mutants established that *L. pneumophila dsbA1*, but not *dsbA2*, restored motility to a *dsbA* mutant. In a protein folding PDI detector assay strain, *dsbA2*, but not *dsbA1*, complemented a *dsbC* mutant of *E. coli*. Deletion of the dimerization domain sequences from DsbA2 produced the monomer (DsbA2N) which no longer exhibited PDI activity, but complemented the *E. coli dsbA* mutant. PDI activity was demonstrated in vitro for DsbA2, but not DsbA1 in a nitrocefin-based mutant TEM β-lactamase folding assay. In an insulin reduction assay, DsbA2N activity was intermediate between DsbA2 and DsbA1. In *L. pneumophila* DsbA2 is maintained as a mixture of thiol and disulfide forms while in *E. coli*, DsbA2 was present as the reduced thiol. Our studies suggest that DsbA2 is a naturally occurring bifunctional disulfide bond oxidoreductase that may be uniquely suited to the majority of intracellular bacterial pathogens expressing T4SSs as well as in many slow growing soil and aquatic bacteria.
INTRODUCTION

*Legionella pneumophila* and related species reside in aquatic environments as obligate intracellular parasites of amoebic hosts (1, 2). When transmitted to susceptible humans via aerosols, these bacteria can cause an acute pneumonia known as Legionnaires’ disease (LD) (3, 4). Since human to human transmission does not occur with LD, there is little opportunity for evolution of virulence or for development of antibiotic resistance. Like many aquatic microorganisms, *L. pneumophila* displays a dimorphic life cycle, alternating between vegetative replicative intracellular bacteria and planktonic terminally differentiated cysts that are highly infectious and resilient to environmental stresses (5, 6, 7, 8). Both ultrastructural and proteomic analyses of the transition to cyst forms indicate substantial remodeling of the cell envelope and the heightened-infectivity suggests the type IVb Dot/Icm secretion system (T4SS), the major virulence determinant, has become fully functional (7, 9, 10, 11). Since much of the cell remodeling occurs extracytoplasmically, we have been investigating proteins that participate in these processes. These studies led to identification of disulfide bond oxidoreductase DsbA2, that in addition to essential functions, seemed to be required for proper assembly or function of the Dot/Icm T4SS (10). Phylogenetic analysis revealed that DsbA2 formed a distinct clade from DsbA and was broadly distributed in the microbial world from environmental species to nearly all microbial pathogens expressing T4SSs (10).

The DSB oxidoreductase family is part of the thioredoxin (TRX) super family of proteins, which are defined by the presence of one or more thioredoxin folds (12). These enzymes participate in disulfide bond formation through a conserved Cys-X-X-Cys (CXXC) active site motif, and can either reduce or oxidize target substrates. DsbA is maintained in the oxidized (disulfide) form by membrane spanning DsbB partner which delivers reducing equivalents from DsbA into the electron transport chain via quinone reduction (13, 14). Similarly, DsbC and DsbG are maintained in the reduced (thiol) form by DsbD which transfers reducing equivalents from the cytoplasmic thioredoxin/thioredoxin reductase system across the cytoplasmic...
membrane to these periplasmic partners (15, 16, 17). Separation of the two systems is considered important to avoid futile cycling of reducing equivalents and this is accomplished through the high affinity of DsbD for homodimeric DsbC and DsbG, which are poor substrates of DsbB (15, 18, 19). While DsbA catalyzes the formation of consecutive disulfide bonds in nascent polypeptides entering the periplasm, DsbC, a protein disulfide isomerase (PDI), is thought to repair inappropriate disulfides or introduce nonconsecutive disulfide bonds through a process of reduction and re-formation of disulfide bonds to aid correct protein folding (15, 19, 20). It is generally believed that once correct disulfide bonds are formed, the rapid folding of proteins, their interactions with periplasmic chaperones or other assembly systems protects these bonds from further redox action (21).

As noted previously, the genes encoding DsbC and DsbG are absent from the genomes of DsbA2-expressing species and some like *Coxiella burnetii* also lack DsbA (10). In contrast to *dsbA* mutants in other bacteria, including pathogens (17, 22), we showed that *dsbA1* mutants of *L. pneumophila* were indistinguishable from wild type parental strains for infectivity (amoeba and HeLa cell models), motility and were without correlating phenotypes, suggesting that DsbA2 likely plays a greater role in managing disulfide bonding and protein folding (10). We also showed that expression of a mutant DsbA2 (P198T) protein in *L. pneumophila* produced a dominant negative effect on DsbA2 function, resulting in loss of motility and infectivity; the latter traced to a functional defect in the Dot/Icm secretion apparatus (10). Based on DsbA2 existing as a mixture of oxidized and reduced forms in the periplasm of *L. pneumophila* and together with the ability of DsbA2P198T mutant protein to form stable disulfide bonds with substrate proteins, including the cysteine containing components (DotG and DotC) of the core Dot/Icm structure, we concluded that DsbA2 was functionally similar to DsbA in delivering disulfide bonds to nascent polypeptides (10).

Here we show that the extended N-terminal amino acid sequence of DsbA2 (compared to DsbA) contains a highly conserved dimerization domain enabling DsbA2 to exist as a
homodimer in the periplasm of \textit{L. pneumophila}. Through complementation tests, we show that expression of \textit{dsbA2} in an \textit{E. coli dsbC} mutant restored protein disulfide isomerase activity to nearly wild type levels. While DsbC-DsbA chimeras have been shown to exist as a mixture of reduced thiol and disulfide moieties that enable oxidase and PDI activities in \textit{E. coli} (19), DsbA2 appears to be an example of a naturally occurring bifunctional DSB oxidoreductase that is widely distributed in the microbial world and especially in those intracellular parasites also expressing T4SSs.

**MATERIALS AND METHODS**

\textbf{Bacterial strains and growth conditions.} \textit{Legionella pneumophila} strains and plasmids are listed in Table 1. Strains were grown aerobically at 37°C on buffered charcoal yeast extract agar (BCYE) (23) or in buffered yeast extract (BYE) broth and supplemented with α-ketoglutaric acid (1 mg/ml), ferric pyrophosphate (250 μg/ml), L-cysteine (40 μg/ml), thymidine (100 μg/ml), and antibiotics where required. Starter cultures were prepared as previously described (24) and used to inoculate pre-warmed BYE to an optical density at 620 nm (OD\textsubscript{620}) of 0.2. For growth curve determinations, samples were taken every two hours (triplicate) and optical density was determined at 620 nm. \textit{E. coli} strains used in these studies are listed in Table 1. These strains were grown at 37°C on Luria-Bertani (LB) agar or in LB broth supplemented with the appropriate antibiotics. Antibiotics (Sigma-Aldrich Ltd.) were added to media at the following concentrations, when appropriate: streptomycin (100 μg/ml), kanamycin (40 μg/ml), gentamicin (10 μg/ml), chloramphenicol (20 μg/ml), and ampicillin (100 μg/ml). All strains were stored at -85°C in nutrient broth containing 10% dimethyl sulfoxide.

\textbf{Protein purification and estimation of mass.} An N-terminal Hexa-His (H\textsubscript{6}) tagged DsbA2 and DsbA1 were constructed in pET15b expression vector and cloned into \textit{E. coli} BL21(DE3) CodonPlus RIL (Stratagene) as previously described (10). N-terminal H\textsubscript{6} tagged
DsbA2LDN, minus the leader sequence and dimerization domain was created using primer pairs DsbA2LDNNdeI (5'-GGAATTCCATATGGCCGCAATTCAGGAAAAT) and DsbA2expBamHI (5'-TTAGATGGATCCTTAATTGCCAGCCG). The amplicon was cloned into pET15b and expressed in the BL21 strain. All genetic constructs were verified by DNA sequencing and protein expression was determined by SDS-PAGE and immunoblot with anti-DsbA2 serum (10). DsbA2LDN was purified by nickel affinity chromatography (Novagen (Madison, WI) as previously described (10). The molecular mass for DsbA2 and DsbA1 (ca 3 mg/ml protein) were estimated by gel filtration in 50 mM Tris-HCl, 100 mM NaCl at pH 8 on a calibrated HiLoad 16/60 Superdex-200 column (GE Healthcare).

**Isolation of native DsbA2 by osmotic shock.** Stationary phase bacteria (strain AA100) were collected by centrifugation at 13,000 x g for 15 min at 4°C and the pellet (1 g wet weight) was suspended in 40 ml of sucrose buffer (0.5 M sucrose, 30 mM Tris-HCl + 1 mM EDTA at pH 8) and gently shaken for 10 min at 25°C. Following centrifugation, the pellet was suspended in 20 ml of ice cold 30 mM Tris-HCl + 1 mM EDTA buffer, gently shaken for 10 min at 4°C and centrifuged. The shockate was decanted and concentrated to 5 ml and loaded onto the calibrated HiLoad 16/60 Superex-200 column and fractions corresponding to the homodimer and monomer were subjected to SDS-PAGE. The proteins were transferred to nitrocellulose and developed with DsbA2- specific antibody diluted 1:10 000 in PBS with 0.1% Tween 20 as previously described (10, 25).

**Complementation studies.** *E. coli* wild type (JCB570) and a *dsbA* mutant (JCB571) were kindly provided by Dr. J. C. Bardwell. The plasmid pBC containing either the coding sequence of *dsbA1*, *dsbA2*, or empty vector were transformed into the *E. coli* strains and selected by chloramphenicol resistance. To construct an in frame deletion of the dimerization domain of DsbA2 (DsbA2N), a vector-free strategy was employed using overlapping primers (DsbA2NF 5' TGGCAGCTGATCAGGAAAATGCTGAACAAG and DsbA2NR 5' GCATTTTCCTGATCAGCTGCCATTATTGCA) to join the 5’ leader sequence with sequences...
downstream of the dimerization domain and using flanking primers Com1FSD (5’
GGGAATTCTAAGGGGAATTACGTGAAATTTAC) and DsbA2BamHIR (5’
TAAGGATCCTTAATTGCCAGCCG) to amplify the joined construct (24). The resulting
amplicon was cloned into pBC and pMMB206 vectors. Soft agar LB plates (0.4% agar)
supplemented with 1 mM IPTG were prepared and 2 µl of cell suspension was inserted into the
middle of the plate. The LB plates were incubated overnight at 30°C and motility was assessed
by measuring the diameter of spreading bacterial growth and reported as percent of the WT
control. Motility assays were performed in triplicate and the results from representative plates
are presented.

**PDI Detector assay.** The PDI detector assay utilizes the TEM1 β-lactamase of pBR322
with an engineered non-consecutive disulfide bond (cysteine residues added S81C and T108C)
that requires disulfide bond isomerase activity to be properly folded in the *E. coli* periplasm (20).
Strains kindly provided by Dr. Bardwell and listed in Table 1 include: RGP209 (*dsbC*
mutant control); RGP663 + pPDI detector plasmid (wild type positive control); and RGP665 (RGP209 +
pPDI detector plasmid). Ampicillin resistance is evaluated at 0, 1, 2 or 3 g/l. For
complementation studies, *dsbA1*, *dsbA2*, *dsbA2LDN* and *dsbA2N* were cloned into pBC with
selection for chloramphenicol. Bacterial cells were grown overnight in appropriate antibiotic,
diluted 1:20, and grown for 2-3 hours to an OD600 of 0.7 with 1 mM IPTG induction. Cells were
then serially diluted and plated in triplicate on LB plates with 0 g/L Amp, 1 g/L Amp, 2 g/L Amp,
or 3 g/L Amp and CFU recorded at 24h. One set of representative plates depicting typical
results is presented.

**Insulin reduction assay.** Reductase activity was assessed by an insulin precipitation
assay with minor modifications (10, 13). Bovine insulin was dissolved in Tris/HCl to 10 mg/ml
(1.67 mM), and titrated to pH 7.5, creating a clear solution. Reactions (triplicate) were carried
out in 200 µl of 100 mM sodium phosphate buffer pH 7.0, 150 µM insulin, 0.33 mM DTT, and 2
mM EDTA and incubated in a 96 well plate format at room temperature in a VersaMax (Molecular Devices) plate reader and absorbance was measured at 650 nm. The insulin reduction assay was initiated by adding proteins at 5 μM of DsbA2, DsbA2LDN and DsbA1 protein purified as H6 tagged proteins following induction in E. coli strain BL21. Both time to start of insulin reduction and specific activity were determined in triplicate using the enzyme kinetics program as previously described (10). The results from a typical experiment are presented.

**Isomerization assay.** TEM1 β-lactamase (WT and PDI detector mutant enzyme Bla and MBla, respectively) were obtained from spent culture supernatants of strains RGP663 and RGP665, respectively, following overnight growth in LB medium. Supernatants were concentrated by spin columns (12 ml concentrated to 0.5 ml) (Amicon Ultracel 10k). Bla activity was tested by spotting 5 μl of concentrated supernatant onto nitrocefin impregnated paper disks (Becton Dickinson). The relative concentration of Bla in each fraction was determined by SDS-PAGE following staining with Coomassie brilliant blue. For the PDI assay, either purified leaderless H6-tagged DsbA1 (6.9 mg/ml) or H6-DsbA2 (3 mg/ml) was added to concentrated culture supernatant from strain RGP665 in ratios of 1:5, respectively, and incubated at 37°C for 30 min. Nitrocefin (Calbiochem, La Jolla, CA) was prepared according to manufacturer’s instructions for spectrophotometric assay and a working dilution of 500 μg/ml was prepared in 0.1 M sodium phosphate buffer at pH 7.0. The refolding assay was configured in a 96 well microplate (Costar, Corning, NY) assay (100 μl/well) containing sodium phosphate buffer, 15 μl of the MBla + DsbA1 or MBla + DsbA2 and the assay started by addition of 10 μl of nitrocefin stock. Controls contained no DsbA2 or DsbA1 protein or WT Bla as a positive control. The rates of hydrolysis of nitrocefin at 486 nm were obtained over the linear range of the reaction (30 min) at 30°C in a Molecular Dynamics plate reader and all reactions were run in triplicate and a representative recording of reaction kinetics is presented.
Periplasmic redox status of DsbA2. The in vivo redox status of DsbA2 was determined by alkylation of free thiol groups by 4-acetamido-4′-maleimidylstilbene-2,2′-disulphonic acid (AMS Molecular Probes, Eugene, OR) essentially as described (10, 26).

Briefly, *L. pneumophila* was grown to stationary phase (~24 h) and half of the culture was collected by centrifugation and washed once in sterile water and then suspended to an OD$_{660}$ of 0.5 in sterile filtered tap water. The water suspended bacteria were kept in the dark for 48 h at 22°C, a period of time previously shown to promote differentiation into cyst-like dormant forms (7). The stationary phase bacteria were collected by centrifugation and were divided into aliquots, one of which was first treated with 10 mM DTT, then TCA precipitated, and alkylated with 100 mM AMS; one TCA precipitated and alkylated with AMS; and one served as an untreated control. The water treated bacteria were TCA precipitated and treated with 100 mM AMS. MalPEG5000 (N-ethylmaleimide covalently bound to polyethylene glycol) was used to alkylate free thiols of DsbA2 and DsbA2N expressed in *E. coli*. Aliquots were similarly treated as described for AMS except that 2 mM DTT was used to reduce DsbA2 prior to alkylation with 5 mM MalPEG5000. In both treatments, samples were analyzed by SDS-PAGE and Immunoblot observed as band shifts.

RNA isolation. Total *L. pneumophila* RNA was extracted from cells at an OD$_{620}$ 0.9 by the RiboZol™ RNA Extraction Reagent (Amresco, OH). Total RNA was treated with DNase I and RNA concentrations were determined using a NanoDrop® ND-1000 spectrophotometer (Nanodrop Technologies, USA). First-strand cDNA was generated using SuperScript First-Strand synthesis system for RT-PCR (Invitrogen, CA). When the RT reaction was completed samples were adjusted to 200 μl with molecular biology water.

Real-time PCR analysis of gene expression. The expression of a *dsbA1* and *dsbA2* was measured in early exponential and stationery phases of growth. Real-time PCR assays
were performed using CFX96 Real-time PCR Detection System (Bio-Rad). The final reaction volume (25 μl) contained: 12.5 μl SYBR® Green PCR Master Mix (Applied Biosystems), 1 μl forward primer 10 μM, 1 μl reverse primer 10 μM, 5 μl cDNA template and 5.5 μl molecular biology water. The real time PCR assays were carried out using a standard program: 95 °C for 10 min, 35 cycles at 95 °C for 15 s, 55 °C for 15 sec and 60 °C for 30 sec. Gene expression was analyzed using the comparative threshold cycle (2^−ΔΔCT) method with rplJ expression as endogenous control (24). All qPCR reactions were carried out in triplicate and the mean and standard deviation computed.

**Phylogenetic analysis.** DsbA1 and DsbA2 amino acid sequences used in phylogenetic analysis were obtained from the Legiolist web server (http://genolist.pasteur.fr/LegioList/upload.cgi) and orthologous genes from GenBank by BLASTP search using DsbA1 and DsbA2 of *L. pneumophila* Philadelphia-1 and DsbA, DsbC and DsbG from *E. coli*. Phylogenetic trees were generated by multiple sequence alignment of unedited DsbA2 sequences using CLUSTALW. Analyses were further refined by BLASTP search using the highly conserved dimerization domain sequence (SLSDAQKKEIKVIHDYLINNPEVLLEASQA).

**RESULTS**

Previous phylogenetic analysis of DsbA2 clade indicated that this group has diverged from the DsbA lineage of disulfide bond oxidoreductases (10). Comparisons of the structures of DsbA2, DsbA and DsbC shows that DsbA2 contains an additional 56 amino acids (86 total) from the N-terminus to the CXXC motif compared with 31 for DsbA and 107 for DsbC. A refined BLASTP search using the dimerization domain region depicted in Figure 1A revealed a 27 aa sequence (bolded in Figure 1A) present in all the bacterial species expressing DsbA2 listed in Table 2. Moreover, this sequence is more discriminating than searches with the whole protein or the C-terminal DsbA-related sequences used previously (10). Table 2 includes representative
species based on a similarity cut off of $e^{-34}$ and is not meant to be comprehensive. We noticed that below this cut off, the DsbA2 clade begins to merge into the DsbA clade. The DsbA2 clade includes a group of intracellular parasites that commonly cause disease in humans, animals and plants and share in common a T4SS. Other genera that do not express T4SSs include the nitrogen fixing endosymbionts associated with leguminous plants that includes *Bradyrhizobium* and related genera (not listed) and many soil and aquatic free-living genera that include *Azospirillum, Rhodopseudomonas, Rhodospirillum* and *Caulobacter*. One exception is *Bordetella pertussis*, which has a T4SS (PTL) and expresses DsbA/DsbC system similar to the *E. coli* system.

In addition, the spacing between the CXXC motif and the resolving *cis*-proline motif is conserved between DsbA2 and DsbA compared with DsbC. Finally, as highlighted in Figure 1B, the first amino acid to the N-terminus of the *cis*-proline in DsbA2 (threonine) is similar to DsbC and not DsbA which substitutes valine. Further genomic analysis using *E. coli* DsbC and DsbG amino acid sequences in the BLASTP searches revealed that the genera expressing DsbA2 listed in Table 2 also lacked orthologues of DsbC and DsbG. Our previous studies showed that *L. pneumophila ΔdsbA1* was indistinguishable from the wild type strain for motility and infectivity for amoeba and HeLa cells, and in close relative *Coxiella burnetii, dsbA1* is completely absent from sequenced genomes (10). While there is plenty of precedent for interchangeability of DsbA and DsbC functions in *E. coli* (18, 19), we investigated whether DsbA2 represents a functional equivalent of DsbC or if equilibrium of monomers and homodimers is responsible for the apparent bifunctional phenotype and thus a variation of the *E. coli* DsbA/DsbC paradigm.

**DsbA2 exists as a homodimer in *E. coli* and *L. pneumophila**. To test whether DsbA2 exists as a mixture of monomers and dimers, size exclusion chromatography was employed. For this study, His-tagged DsbA1 and DsbA2 were treated with iodoacetamide to limit formation of spurious disulfides prior to nickel interaction chromatography. Each protein was applied to a calibrated gel filtration column and as seen in Figure 2A, DsbA1 eluted at 95
minutes, with an apparent mass of 27 kDa, consistent with a monomer. In Figure 2B, DsbA2 eluted as a single peak at 75 minutes, with an apparent mass of 60 kDa, consistent with the size of the homodimer. Subsequent purifications of DsbA2 indicated that inclusion of iodoacetamide was unnecessary (data not presented). To ensure that the homodimeric form was not an artifact generated in *E. coli,* we applied osmotic shockates from *L. pneumophila* similarly treated with iodoacetamide over the same calibrated column and DsbA2 eluted as the dimer (see Figure 2C). The elution of DsbA2 as a single peak suggests that if the monomer form is present in *L. pneumophila,* it is below the limit of detection in our assay.

**Legionella pneumophila dsbA1 and DsbA2N, but not dsbA2 restores motility to a dsbA mutant of *E. coli.*** In *E. coli,* DsbA is required for disulfide bond formation in the flagellar P-ring protein (FlgI), which is necessary for motility in soft agar (27). As seen in Figure 3A, wild type *E. coli* strain JCB570 is motile in 0.4% soft agar, whereas ∆*dsbA* mutant strain JCB571 is not. We introduced *L. pneumophila* dsbA1 and *dsbA2* expressed from pBC plasmids into JCB571 to test whether expression of these proteins could restore motility. As seen in Figure 3A, *dsbA1* complemented motility to ~60% of wild type (diameter of spreading growth), whereas *dsbA2* did not, consistent with the general view that dimeric proteins, such as DsbC, are poor substrates of the *E. coli* DsbB (18,19, 28). We next removed the dimerization domain of DsbA2 (DsbA2N) by an in frame deletion that retained the signal sequence as depicted in Figure 1A to create the monomer. The DsbA2N monomer was confirmed by gel filtration. Expression of *dsbA2N* in JCB571 restored motility to nearly wild type activity (>70%) of control (see Figure 3B). While growth rate differences might account for the partial complementation in our studies, these were not apparent and more likely; these differences reflect different efficiencies of DsbA1 and DsbA2N oxidation by the *E. coli* DsbB compared with DsbA. These results indicate that it is the dimerization domain of DsbA2 that prevents interaction of the homodimer with DsbB.
Legionella pneumophila DsbA2, but not DsbA1 or DsbA2N, exhibits PDI activity.

In *E. coli*, DsbA catalyzes disulfide bonding between consecutive cysteine residues as the nascent polypeptide enters the periplasm (26). For those proteins for which non-consecutive disulfide bonding is required for proper folding, consecutive disulfide bonds must be reduced and then reformed between non-consecutive cysteine residues. In *E. coli*, the protein disulfide isomerase DsbC catalyzes this reaction (20). In our previous studies, we noted that some of the proteins captured by DsbA2 contained an odd number of cysteine residues and prediction software analyses suggested that some of these might be non-consecutive (10). Since DsbA2 exists as a homodimer, we tested the possibility that DsbA2 might complement a *dsbC* mutant of *E. coli* by restoring PDI activity. To test this hypothesis, *dsbA2*, *dsbA2N* and *dsbA1* were cloned into an *E. coli* strain mutant for *dsbC* and containing a pPDI detector system (20). In this system, a TEM β-lactamase (Bla) which naturally contains two cysteine residues at positions 52 and 98 that are not required for biological activity was engineered to contain an additional two cysteine residues (81 and 108) positioned so that DsbA will introduce consecutive disulfide bonds and produce an inactive enzyme (20). In the presence of DsbC, biological activity is restored by PDI activity (20). In this PDI detector assay, only *dsbA2* expression restored ampicillin resistance to nearly wild type DsbC control levels at 2 or 3 g/l of drug (see Figure 4A and B). In contrast, DsbA1 (depicted in Figure 4A and 4B), DsbA2LDN (Figure 4A) and DsbA2N (Figure 4B) failed to complement in the PDI detector assay. The potent PDI activity displayed by DsbA2 in *E. coli* also suggests that DsbA2 is an efficient substrate of the DsbD reductase system, perhaps tipping the balance in favor of PDI/reductase activity over oxidase activity. To confirm the redox status of DsbA2 and DsbA2N in *E. coli*, we used the thiol alkylating agent MalPEG5000 to alkylate free thiols and those thiols reactive with MalPEG result in an increased mass of 5000 daltons per cysteine residue. As seen in Figure 5, whole *E. coli* bacteria expressing DsbA2 were first treated with DTT before alkylation with MalPEG (Lane 1) or in the absence of prior DTT treatment (Lane 3) and in both cases showed shifted high molecular...
weight bands resulting from complete alkylation by MalPEG relative to untreated control (Lane 2). In contrast, when DsbA2N is treated with reducing agent and then MalPEG, multiple shifted bands are observed (Figure 5, lane 6); whereas, treatment with MalPEG without reducing agent (Lane 8) produced no shifted bands and showed no differences from that of the untreated control (lane 7) indicating that DsbA2N is maintained as the disulfide in *E. coli*. These results indicate that DsbA2, but not DsbA2N is fully reduced to the free thiol in the periplasm of *E. coli*.

**In vitro PDI activity.** To confirm in vivo PDI activity for DsbA2, we developed an assay in which the PDI detector TEM β-lactamase serves as substrate. In this assay, gain of Bla activity as measured with chromogenic substrate nitrocefin, is dependent on PDI enzyme reducing and shuffling disulfide bonds until correct nonconsecutive disulfides are formed (29). As seen in Figure 6, the Bla activity of MBla in concentrated spent culture supernatants is below the level of detection over the course of the assay. Incubation of DsbA2, but not DsbA1 with MBla efficiently corrected disulfide bonding and catalyzed protein folding as noted by a gain in enzyme activity with nitrocefin. Compared with the WT Bla similarly concentrated from culture supernatants, near equivalent activity was obtained. These studies were repeated several times and a representative experiment presented. These studies confirm the biological assay for PDI activity for DsbA2.

**Insulin reduction assay of DsbA1, DsbA2 and DsbA2LDN.** An insulin reduction assay is often used to demonstrate thioredoxin like activity of DSBs (13, 30). This assay can distinguish between strong oxidases like DsbA and reductases such as thioredoxin by both the time to reduction and by the rate of insulin precipitation resulting from the reduction of its disulfide bond (31). In this regard, we previously demonstrated that DsbA2 was similar to thioredoxin in this assay (10). As seen in Figure 7, DsbA1 of *L. pneumophila* has a lag time of 20 minutes compared to DsbA2 which has a lag time of 5 minutes and more typical of reductases like DsbC and thioredoxin. However, as a monomer, DsbA2LDN is much more active in this assay than DsbA1, though the time to reduction is doubled for the monomer
compared with the dimer. These results suggest that monomeric DsbA2LDN, while structurally
more similar to DsbA than DsbC, favors the reductase function (10).

**Growth stage expression of dsbA1 and dsbA2.** To determine at what stage of growth
dsbA1 and dsbA2 are expressed, RT qPCR was performed on cDNA prepared from total RNA
obtained from exponential and stationary phase grown *L. pneumophila*. As seen in Figure 8,
dsbA1 was expressed two-fold higher in stationary phase than exponential phase, whereas
dsbA2 expression in stationary phase was increased more than five-fold. We had previously
reported that DsbA2 protein levels were more abundant in differentiated cyst forms than
stationary phase forms (7) and based on transcript levels; both *dsb* genes appear to be up-
regulated in stationary phase.

**Redox status of DsbA2 is unchanged through cyst phase.** In Gram negative
bacteria, DsbA is maintained fully oxidized by DsbB, while DsbC is maintained fully reduced by
the DsbD system. We previously reported that DsbA2 existed as a mixture of oxidized and
reduced forms in *L. pneumophila* during stationary phase (10). We examined bacteria harvested
from stationary phase as well as from water differentiated metabolically dormant forms (7).
Collected bacteria were TCA precipitated and subjected to AMS alkylation, which adds ~500
daltons to free cysteine thiol residues which can then be observed as a shift in apparent mass
by SDS-PAGE and immunoblot. As seen in Figure 9, even after suspension in tap water for
several days, DsbA2 exists as a mixture of thiols and disulfides in the cyst-like forms. Since
these forms are nearly dormant metabolically and exhibit no measurable respiration rate (7), it is
not know how the mixture is maintained under these conditions.
The results of our studies show that DsbA2 of *L. pneumophila*, which contains a 56 amino acid N-terminal extension relative to DsbA of *E. coli*, forms a homodimer in *L. pneumophila* and displays protein disulfide isomerase activity in both *in vivo* and *in vitro* assays. We previously reported that DsbA2 behaved similarly to DsbA by catalyzing the formation of disulfide bonds in secreted proteins of *L. pneumophila* (10). Consistent with DsbA-like activity, a DsbA2 cis-proline (P198T) mutant protein formed stable disulfide cross-linked complexes with substrate proteins (10). Only the disulfide is capable of capturing substrate proteins (26).

Previous studies found that *L. pneumophila* DsbA1 was dispensable for growth, motility, and infectivity in cell-based assays; and that this gene is naturally absent from the genome of *C. burnetii* which also expresses a *dsbB* (10). Taken together, these findings suggest that the DsbA2 lineage may be naturally bifunctional and therefore functionally unique from the DsbA/DsbC paradigm so well characterized in *E. coli*.

In support of this notion, we determined that DsbA2 was present as a mixture of reduced thiol and disulfide forms *in vivo*, which contrasts with DsbA and DsbC which are mostly oxidized and reduced, respectively (28). We determined by gel filtration that the mixture of oxidized and reduced forms was not due to equilibrium of monomer and dimer forms. Our studies do not rule out the possibility that each arm of the DsbA2 dimer might contribute to the mixture, one as the disulfide and the other as the free thiol. Our studies suggest that structural differences within the thioredoxin folds of DsbA2 might render them more accessible to the DsbB oxidases of *L. pneumophila*. In this regard, *L. pneumophila* expresses two alleles, DsbB1 and DsbB2 also known as LidJ (32) that show sequence divergence from *E. coli* DsbB. As previously reported, *L. pneumophila* also expresses two alleles of DsbD (10). We suggest that these competing oxidases and reductases interact with DsbA2 at differing efficiencies to maintain an equilibrium between oxidized and reduced forms. Interestingly, Segatori *et al.* showed that a chimera composed of the dimerization domain of DsbC fused to the catalytic domain of DsbA also
existed as a mixture in *E. coli* and displayed bifunctional activity including partial complementation of a *dsbA* mutant for motility (19). Since DsbA2 did not restore motility to a *dsbA* mutant, either DsbA2 is a poor substrate of DsbB or a good substrate of DsbD. Our studies found that DsbA2 is maintained as the free thiol in *E. coli* by DsbD and efficiently complemented a *dsbA* mutant in the PDI detector assay. By demonstrating that DsbA2N was able to complement a *dsbA* mutant of *E. coli*, we conclude that the lack of interaction between DsbA2 and DsbB is most likely the result of steric hindrance by the homodimer.

Our studies suggest that DsbA2 is not a homodimeric DsbA as demonstrated with the *E. coli* DsbC-DsbA chimera. We suggest that key amino acid differences in the cis-proline domain (TPA versus VPA in DsbA), similar to DsbC may favor PDI reductase activity. In the insulin reductase assay, DsbA2 and even the DsbA2N monomer exhibited reductase characteristics more similar to DsbC than to DsbA (see Figure 7). Thus, the DsbA2 lineage appears to be a unique adaptation which combines both oxidase and reductase functions into a single protein in situ. To explore the basis for this adaptation, we have replaced the *E. coli dsbB* and *dsbD* genes with alleles from *L. pneumophila* and future studies will likely show that structural divergence in the two DsbD alleles may affect reducing efficiency of DsbA2.

A number of refolding assays have been developed to demonstrate in vitro PDI activity (15, 33, 34). These assays often rely on refolding of scrambled enzymes like RNase I that contains consecutive as well as nonconsecutive disulfide bonds and a subsequent assay for gained RNase activity (15, 34). Since most DSB enzymes are technically bifunctional (oxidase and reductase), PDI assays can yield mixed results such as DsbA refolding RNase I (34). One advantage to use of the PDI detector MBla for these assays is that the protein does not require denaturation prior to assay and the consecutive disulfide bonds produced are the result of DsbA activity (20). In this folding assay, gain of enzyme activity is readily detected with the chromagenic penicillin substrate nitrocefin. DsbA2 and not DsbA1 efficiently restored the correct nonconsecutive disulfide bond pairing required for protein folding and Bla activity. The
adaptation of this assay to a micoplate format should enable development of rapid screens of proteins for PDI activity. In our hands, the deep red color of the cleaved β-lactam is easily distinguished by visual observation.

Remarkably, much of the microbial world apparently substitutes DsbA2 for DsbC/DsbG, including many soil and aquatic species such as *Caulobacter, Bradyrhizobium, Aeromonas*, and *Azospirillum*; and most human, animal, and plant pathogens that express T4SSs as major virulence factors (35). The latter group includes *Rickettsia, Legionella, Bartonella, Anaplasma, Brucella, Ehrlichia, Coxiella, Agrobacterium* and *Fransicella*. Most of these genera have doubling times of two hours or more which raises the possibility that in fast growing bacteria, DsbA/DsbC system is required to accelerate correct disulfide bond formation to enable protein folding to keep pace with cell division. This possibility seems to be supported by growth rate differences noted with *E. coli* strains carrying single mutations in *dsbA* and double mutations (*dsbA dsbC* and *dsbA dsbD*) (36). In this regard, DsbA accelerates disulfide bond formation between consecutive cysteines as most nascent polypeptides enter the periplasm (15, 37, 38). However, in other proteins, disulfide bonds may form after the protein has entered the periplasm (37) or require correction of mispaired disulfides by DsbC (21). We suspect that in those slow growing bacteria expressing DsbA2, such as *L. pneumophila*, disulfide bonds might be introduced after proteins enter the periplasm, which is supported by the efficiency and diversity of proteins captured by the DsbA2P198T mutant protein (10). It follows that the apparent essentiality of *dsbA2* is due to both its bifunctional nature as well as to an absence of any backup system to repair miss-oxidized proteins that would accumulate to toxic levels in the periplasm. We also noted that over expression of the DsbA2P198T mutant protein rendered *L. pneumophila* non-infectious by interfering with assembly and function of the T4SSs (10). Since *dsbA2* was also essential in strains with mutations in *dot/icm* structural genes, we concluded that DsbA2 must be associated with proper folding of other proteins whose functions might also be essential including OmpS porin and periplasmic Hsp60 (10, 39, 40, 41).
In summary, we have shown that the DsbA2 protein of *L. pneumophila* is a homodimer and is capable of delivering disulfide bonds to native unfolded proteins or through PDI activity to repair inappropriate disulfide bonds. While the DsbA2 lineage represents a subset of the DsbA family, the dimerization domain and threonine substitution for valine in the cis-proline domain are more similar to DsbC and likely contribute to PDI activity. We find it interesting that much of the microbial world relies on DsbA2 instead of the DsbA/DsbC system to manage extracytoplasmic disulfide bonding and protein folding. This genetically diverse group also includes most intracellular parasites that express T4SS, all of which grow very slowly when compared to *E. coli*. Perhaps DsbA2 represents an evolutionary adaptation that is more efficient in orchestrating disulfide bond formation and protein folding that suits the lifestyles of slow growing bacteria.

**Acknowledgements**

We thank J. C. Bardwell and G. Ren for helpful suggestions and providing the bacterial strains and other reagents used in these studies. This work was supported by NIH Grant R01 AI066058 to P.S.H.

**References**


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Table 1. Strains and plasmids.
This is a partial list from more than 60 genomes identified by BLASTP with DsbA2 and the 27 amino acid dimerization domain sequences depicted in Figure 1A. For DsbA1, DsbB, DsbD, DsbC, DsbG, and CcmG, the \textit{E. coli} protein sequences were used for the searches against each genus. The presence or absence of a type IV secretion system (T4SS) was determined from the literature. The question mark for \textit{Francisella tularensis} indicates that the system may be incomplete. CcmG is a thioredoxin like periplasmic enzyme associated with cytochrome c maturation.
Figure legends

**Figure 1.** Dimerization domain and Secondary structure comparison. A. DsbA2 amino acid sequence depicting the leader sequence and the detached dimerization domain sequence deleted to create DsbA2N. The underlined sequences (27 amino acids) of the dimerization domain are conserved and those in larger letters represent amino acids that are highly conserved among DsbA2 members listed in Table 2. The CXXC and cis-proline regions are in boldface. B. Secondary structure predictions for *L. pneumophila* DsbA1, which is similar to DsbA of *E. coli*, *E. coli* DsbC and *L. pneumophila* DsbA2 are depicted. The CXXC region and the cis-proline motifs are boxed. Note that the cis-P region of DsbA2 is similar to that of DsbC.

**Figure 2.** Size exclusion of native proteins from *L. pneumophila*. A. DsbA1 elutes as a single peak at 95 minutes consistent with a monomer and an estimated mass of 25 kDa. B. DsbA2 elutes at 75 minutes consistent with the homodimer and estimated mass of 60 kDa. The HiLoad 16/60 Superdex-200 column was calibrated with ribonuclease A (13.7 kDa), chymotrypsinogen A (25 kDa), ovalbumin (43 kDa), albumin (67 kDa), and blue dextran 2000 (void volume) are marked. C. Immunoblot of DsbA2 containing fractions prepared by osmotic shock from *L. pneumophila* and collected from the calibrated column. The DsbA2 protein from *L. pneumophila* eluted at 75 minutes, consistent with the homodimer. The Immunoblot was developed with antibody specific for DsbA2.

**Figure 3.** Complementation of motility defect in soft agar. Spreading motility by *E. coli* in soft agar is dependent on functional flagella and chemotaxis. A. Wild type *E. coli* (JCB570) is motile in soft agar at 30°C. Strain JCB571(*dsbA*) is mutant for *dsbA* and nonmotile in this medium. Strain JCB571 + pBCLpdsbA2 did not complement the motility defect whereas; JCB571 + pBCLpdsbA1 complemented (~60%) this strain for motility. B. Wild type *E. coli* strain...
JCB570 is motile in soft agar and the *dsbA* mutant strain JCB571 is not. Expression of
*pBCdsbA2N* (*pdsbA2N*) which contains an in frame deletion of the dimerization domain depicted
in Figure 1A restored motility to ~70% of wild type strain JCB570 activity. The results presented
in both panels are representative of typical results obtained from triplicate platings.

**Figure 4. Complementation tests in PDI detector system.** The PDI detector system uses a β-
lactamase engineered to form consecutive disulfide bonds which require DsbC PDI activity to
correct to nonconsecutive to facilitate protein folding (20). Growth on ampicillin indicates β-
lactamase is properly folded. Ampicillin concentrations are 0, 1, 2 or 3 g/L. Bacterial dilutions
are plated from top to bottom as detailed in the text. A. + = Wild type *E. coli* strain RGP663 +
pPDI detector plasmid; - = strain RGP665dsbC + pPDI detector plasmid; 1 = strain RGP665 +
pPDI detector plasmid + pBCdsbA1; 2 = strain RGP665 + pPDI detector plasmid + pBCdsbA2;
and LDN = strain RGP665 + pPDI detector plasmid + pBCdsbA2LDN (LDN = deleted leader
and dimerization domain sequences). B. Replicate PDI detector assay that includes RPG665 +
pPDI detector plasmid + pBCdsbA2N containing the leader sequence and an in frame deletion
of the dimerization domain sequences.

**Figure 5. DsbA2 and DsbA2N redox status in *E. coli.* A.** Redox status of DsbA2 in *E. coli.*
Lane 1, whole cells were reduced with DTT for 30 min and alkylated with MalPEG5000 (reduced
control); Lane 2, untreated control; and Lane 3, whole cells treated with MalPEG5000. Lanes 4
and 5 contain the molecular weight standards. B. Redox status of DsbA2N in *E. coli.* Lane 6,
whole cells treated with DTT for 30 min and then alkylated with MalPEG5000. Lane 7, untreated
control and Lane 8, whole cells treated with MalPEG5000. The molecular weight of DsbA2 is
~28,000 and for DsbA2N ~26,000. Anti-DsbA2 mouse serum was used to identify the DsbA2
proteins. The multiply shifted bands are indicative of DsbA2 or DsbA2N covalent complexes
with MalPEG5000 which adds 5000 daltons per cysteine. In *E. coli*, DsbA2 is maintained as the free thiol and for DsbA2N, as the disulfide.

**Figure 6. Nitrocefin PDI folding assay.** TEM1 β-lactamase (Bla) and pPDI mutant TEM1 β-lactamase (MBla) were obtained by concentrating culture supernatants from strains RGP663 and RGP665 respectively. MBla contains consecutive disulfide bonds and is inactive (nitrocefin negative). DsbA1 or DsbA2 was added to aliquots of MBla for 30 min at 37°C and 15 μl aliquots were tested in triplicate for Bla activity in microplates with nitrocefin as described in the text. Cleavage of the β-lactam of nitrocefin is measured as a change in absorbance at 486 nm over a 30 min assay period. DsbA2, but not DsbA1 exhibited PDI activity.

**Figure 7. Insulin reduction assay.** Reductase activity of equimolar concentrations of DsbA1, DsbA2 and DsbA2N were followed spectrophotometrically at 650 nm in a microplate assay as described in the text. The assays were performed in triplicate and a representative assay is presented. The time to reduction indicates that DsbA2N is intermediate between DsbA2 and DsbA1 (more reducing than oxidizing).

**Figure 8. Growth state transcript levels of dsbA1 and dsbA2.** *L. pneumophila dsbA1* and *dsbA2* mRNA transcript levels were determined with total RNA obtained during exponential (E) phase and stationary (S) phase as described in the text. *dsbA1* and *dsbA2* are up-regulated in stationary phase: *dsbA1* two-fold, and *dsbA2* five-fold. (DsbA1 E = 1 + 0.158; DsbA2 E = 1 + 0.182 DsbA1 S = 2.33 + 0.06; DsbA2 S = 5.68 + 0.415).
Figure 9. Redox status of DsbA2 in *L. pneumophila*. Lane 1, whole cells were treated with DTT for 30 min, then subjected to AMS alkylation (reduced control). Lane 2, whole cells untreated with AMS (oxidized control). Lane 3, stationary phase *L. pneumophila* treated with AMS and Lane 4, differentiated cyst like forms after suspension in tap water for 48 h. The cyst-like forms are resistant to antibiotics and are near dormant metabolically (7). The doublets depicted in lanes 3 and 4 indicate that DsbA2 is present in a mixture of thiol and disulfide forms.
**Figure 1**

**A**  
Dimerization Domain  
TASASLSDAQkKEIEKvehDYLINNPEVLLLeASQALOQQKQQNMQQQQAQA  
Leader sequence  
MKFTSLLTAGALASTLVSPAIMEAAAD QENAEQVFQGLTTVGPKNVTVLFDDYQGCIMCCKKMASTIENL VDKSGLRYKIEPFGKTSIALSRLAAGMGMQKYQAMNALTIDKRLDEKTVMADAASGLDMQKLX KOMDSOEVTDILDANRQLAEKLHLMGTPAFIGSTPDQQYKGSEISFIPGATSEQSLRELKKAGN

**B**  
Dsba1  
Dsbc  
Dsba2
Figure 2

(A) DsbA1

(B) DsbA2

(C) Dimer     Monomer

20.4
29.3
37.3

800 801 802 803 804 805 806 807 808 809 810 811 812 813 814 815 816 817 818 819 820 821 822 823
Figure 3.
Figure 4.

**A**

+ - 1 2 LDN

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**B**

+ - 2 N

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+= WT + pPDI
-= PDI + pPDI
1 = pPDI + dsbA1
2 = pPDI + dsbA2
N = pPDI + dsbA2N
LDN = pPDI + dsbA2LDN
Figure 5

Figure 6
Figure 7

Figure 8

34
Figure 9

1. Reduced
2. Oxidized
3. Stationary phase
4. Cyst