Insight into Disulfide Bond Catalysis in Chlamydia from the Structure and Function of DsbH, a Novel Oxidoreductase

The Chlamydia family of human pathogens uses outer envelope proteins that are highly cross-linked by disulfide bonds but nevertheless keeps an unusually high number of unpaired cysteines in its secreted proteins. To gain insight into chlamydial disulfide bond catalysis, the structure, function, and substrate interaction of a novel periplasmic oxidoreductase, termed DsbH, were determined. The structure of DsbH, its redox potential of −269 mV, and its functional properties are similar to thioredoxin and the C-terminal domain of DsbD, i.e. characteristic of a disulfide reductase. As compared with these proteins, the two central residues of the DsbH catalytic motif (CMWC) shield the catalytic disulfide bond and are selectively perturbed by a peptide ligand. This shows that these oxidoreductase family characteristic residues are not only important in determining the redox potential of the catalytic disulfide bond but also in influencing substrate interactions. For DsbH, three functional roles are conceivable; that is, reducing intermolecular disulfides between proteins and small molecules, keeping a specific subset of exported proteins reduced, or maintaining the periplasm of Chlamydia in a generally reducing state.

Chlamydia are obligate intracellular eubacteria that are phylogenetically distant from other bacterial divisions (1). Virtually every human being will be infected with Chlamydia pneumoniae in their lifetime, and this organism accounts for ~10% of pneumonia cases and 5% of bronchitis cases in the United States (2, 3). Acute infections are usually mild in immunocompetent hosts, but severe pneumonias are observed in immunocompromised patients. Chronic C. pneumoniae infections have been associated with arterial disease (4), where C. pneumoniae act to continuously stimulate inflammation and thereby contribute to stroke and coronary heart disease. Chlamydia have an unique developmental cycle in which the extracellular infectious form, the elementary body, is metabolically inactive and highly resistant to lysis. This is thought to be due in large part to a complex of outer envelope proteins that are highly cross-linked by disulfide bonds (5). Disulfide cross-linked envelope proteins appear to substitute for peptidoglycan in the elementary body (6). This is in contrast to the intracellular, non-infectious but metabolically active reticulate body which employs peptidoglycan (7), like other Gram-negative bacteria. The reliance of the elementary body on disulfide bond-linked proteins suggests that interference with chlamydial disulfide bond catalysis offers a potential avenue for treating chlamydial infections in addition to antibiotics administration. Despite the extensive use of disulfide cross-linked envelope proteins, Chlamydia exhibit an unusually high number of unpaired cysteines in their secreted proteins compared with Escherichia coli. Is the machinery for periplasmic disulfide bond catalysis in Chlamydia different from E. coli?

Functional gene assignment of the C. pneumoniae TW183 genome predicts 5 periplasmic oxidoreductases (8) of a total of 1113 predicted proteins. In comparison, E. coli K-12, for which the catalysis of periplasmic disulfide bond formation has been characterized best (9, 10), possesses 6 periplasmic oxidoreductases of a total of 4243 predicted proteins. The five central disulfide bond-forming proteins (Dsb) of E. coli catalyze the de novo formation of disulfide bonds as well as the isomerization of incorrectly formed disulfide bonds to the correct form. DsbB and DsbA form the functional pair that acts, respectively, to create disulfide bonds and transfer them to substrate proteins. The lack of DsbA, which is the primary periplasmic dithiol oxidant, greatly reduces the virulence of several bacterial species (11, 12), providing further evidence for the potential of treating chronic Chlamydia infections by interfering with disulfide bond catalysis. Reduced DsbA is reoxidized by the integral membrane protein DsbB (13, 14), which in turn regenerates its disulfide bonds by reducing ubiquinone within the membrane (15, 16). Disulfide bond isomerization is thought to be catalyzed by the DsbC-DsbD and DsbG-DsbD pairs. The

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The atomic coordinates and structure factors (code 2ju5) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http://www.rcsb.org/).

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4 The abbreviations used are: Dsb, disulfide bond-forming protein; BLAST, basic local alignment search tool; HPLC, high performance liquid chromatography; NOE, nuclear Overhauser effect.
soluble DsbC and DsbG proteins are kept in their active reduced state by the transmembrane protein DsbD (17, 18). In turn, DsbD gains its reducing equivalents from the cytosolic protein thioredoxin, which ultimately gains its reduced equivalents from NADPH (18). DsbE(CcmG), which is also kept reduced by the DsbD protein, is required for cytochrome c maturation. DsbE is thought to be responsible for keeping two cysteine residues in cytochromes reduced, a prerequisite for heme attachment (19).

The present study identifies two new subfamilies of periplasmic oxidoreductases in Chlamydia, termed DsbH and DsbJ, besides orthologs of DsbA, DsbB, and DsbD. As a first step in the characterization of the chlamydial disulfide bond catalysis, the solution structure of DsbH was determined to establish its structural relationship to the known oxidoreductases. To learn more about DsbH-substrate interactions, the interaction of DsbH with a peptide ligand was examined. The functional characterization of DsbH enzymatic properties was carried out, and the results suggest that DsbH plays the role of a reducing thiol disulfide oxidoreductase in Chlamydia. Compared with for example E. coli, the disulfide bond catalysis machinery of Chlamydia exhibits substantial differences.

**EXPERIMENTAL PROCEDURES**

**Molecular Biology—**The DsbH gene of C. pneumoniae TW-183 (CP80958) was assembled from overlapping oligonucleotides (Retrogen, Inc.). A C41G substitution was implemented based on sequence alignment with Chlamydia caviae GPIC (data not shown). For efficient periplasmic expression in E. coli, the DsbA signal sequence from E. coli was used instead of that from DsbH. A high sequence homology between the two E. coli GPIC TW-183 in ice-cold 50 mM Tris/HCl, pH 8.0, 300 mM NaCl, 10 mM imidazole, 8 M urea, 2 mM β-mercaptoethanol, and the cleared lysate was again applied on the immobilized metal affinity chromatography. After extensive washing in lysis buffer, urea was eliminated from the wash buffer, and subsequently, β-mercaptoethanol was also eliminated, allowing protein folding and disulfide bond formation before protein elution. The remainder of the purification was identical to that for periplasmic DsbH. The GB3-peptide construct was expressed and purified analogously to cytosolic DsbH. The monomeric GB3-peptide fraction from gel filtration containing only intramolecular disulfide bonds was cleaved with tobacco etch virus protease in the absence of any reducing agents at a molar ratio of 1:200 overnight at room temperature. Fusion protein and tobacco etch virus protease were removed from the peptide by immobilized metal affinity chromatography.

**NMR Sample Preparation—**Samples of 300-μl volumes were prepared in 50 mM NaH2PO4/Na2HPO4, pH 6.0, and 0.02% w/v NaCl buffer to contain 1.0 mM of DsbH (cytosolic η280 nm = 21,030 M−1·cm−1). Besides a sample in isotropic solution, the alignment of DsbH relative to the magnetic field was achieved in the presence of bicelles and stretched polyacrylamide gel. A 5% solution of bicelles consisting of 1,2-di-O-tridecanoylsn-glycero-3-phosphocholine (DTPC), 1,2-di-O-hexyl-sn-glycero-3-phosphocholine (DHPC), and cetyltrimethylammonium bromide (CTAB) at molar ratios of DTPC:DHPC = 3:1 and (DTPC+DHPC):CTAB = 30:1 was employed. Stretched, negatively charged polyacrylamide gels (21) were polymerized from a 5.1% w/v solution of acrylamide, 2-acrylamido-2-methyl-1-propanesulfonate (AMPS), and bisacrylamide with a monomer to cross-linker ratio of 39:1 (w/w) and a molar ratio of 98:2 of acrylamide to AMPS, employing the previously described conventions and procedures (22). The bicelle solution gave rise to a 2H splitting of 22.8 Hz at 29 °C, corresponding to a DsbH alignment tensor magnitude of 10.4 Hz. For the stretched gel a 2H splitting of 2.4 Hz was detected at 25 °C, corresponding to a tensor magnitude of 10.1 Hz. DsbH samples containing the employed peptide ligand (Fig. 5D) were prepared to contain 0.1 mM 15N-labeled DsbH and 0.05 mM peptide in the above phosphate buffer.

**NMR Spectroscopy—**All experiments were carried out on a Bruker Avance 700 spectrometer at 25 °C except for the bicelle solution, which was measured at 29 °C to maintain the liquid crystalline phase. Data were processed and analyzed with the nmrPipe package and CARA. H, N, Cα, Cβ, and Cγ assignments were made from HNCA, HNCACB, HNCO, and CBCA(CO)NH experiments. Subsequently, Hα and partial Hβ assignments were obtained from total correlation spectroscopy-heteronuclear single quantum correlation experiments (tmax = 35.9 ms). The side-chain assignments were completed using H(C)CH-1-correlation spectroscopy (COSY) and (H)CCH-COSY experiments (23). Partial assignments of the highly overlapped aromatic side chains were obtained from their constant time (CT)-heteronuclear single quantum correlation and NOE spectroscopy pat-
terns (24). 15N- and 13C-edited NOE spectroscopy spectra were recorded with a mixing time of 100 ms. NOE cross-peak assignments were performed manually. \(^1\)J_{C-H}, \(^1\)J_{C-C}, and \(^1\)J_{N-C} couplings for aromatic and aliphatic residues were obtained from quantitative \(^1\)J-correlation spectroscopy (25) with dephasing times of 50 and 100 ms, respectively. \(^1\)J_{H,N}, \(^3\)J_{C-C}, \(^3\)J_{C-N}, and \(^3\)J_{N-C} couplings were determined from mixed-constant time, H\(^N\)-coupled HNCO (26), and quantitative \(^1\)J-correlation HNCO experiments (27, 28) of isotropic and aligned samples, respectively. \(^1\)H-15N NOE measurements of Tyr and Phe were represented as a \(\text{intensity of methyl resonances, the upper distance bound for weak NOEs, respectively. To account for the higher apparent distance restraints were grouped into four ranges, 1.8–2.7 Å (1.8–2.9 Å for NOEs involving H\(\text{N}\) protons), 1.8–3.3 Å for 13C-edited NOE spectroscopy spectra to values found in \(\alpha\)-helical conformation. The distance restraints were grouped into four ranges, 1.8–2.7 Å (1.8–2.9 Å for NOEs involving H\(\text{N}\) protons), 1.8–3.3 Å (1.8–3.5 Å for NOEs involving H\(\text{N}\) protons), 1.8–5.0 Å, and 1.8–6.0 Å, corresponding to strong, medium, weak, and very weak NOEs, respectively. To account for the higher apparent intensity of methyl resonances, the upper distance bound for NOEs involving methyl groups was extended by 0.5 Å. Interproton distances involving non-stereospecifically assigned methylene protons, methyl groups, and H\(\delta\) and H\(\epsilon\) protons of Tyr and Phe were represented as a \(\Sigma(r^{-6})^{-1/6}\) sum (33). The alignment tensor magnitude and rhombicity for each residual dipolar coupling data set were estimated from molecular fragment replacement (34) and were kept fixed during structure calculations. Alignment tensor magnitudes of 10.4 and 10.1 Hz were obtained for the employed bicelle and gel media, respectively, eliminating the need for media-specific weighting factors.

DsBh structures were calculated by simulated annealing starting at 3000 K using the program XPLOR-NIH (35). Besides standard force field terms for covalent geometry (bonds, angles, and improper torsions) and nonbonded contacts (van der Waals repulsion), the following experimental and empirical potentials were included. Dihedral angle restraints and interproton distance restraints were implemented using quadratic square-well potentials. The difference between predicted and experimental residual dipolar couplings was described by a quadratic harmonic potential. A pseudopotential for the radius of gyration was used (36) as well as a backbone-backbone hydrogen-bonding potential (37) and torsion angle potentials of mean force (38). An ensemble of 20 structures was calculated. The structural statistics are summarized in supplemental Table 1. Atomic coordinates, structural restraints and chemical shift assignments for DsBh have been deposited in the Protein Data Bank and Biological Magnetic Resonance Bank under ID codes 2ju5 and 15431, respectively.

Redox Potential Measurement—Proteins whose redox potentials were to be measured were incubated with DsBh, a protein of known redox potential, \(-235\) mV (39). At equilibrium different protein species were separated by reverse-phase HPLC, and the equilibrium constant \(K_{eq}\) of the reaction and the standard redox potential \(E^{o}\) of the protein were determined as described (40). To verify that equilibrium was reached, it was approached from both directions. To do this DsBh was incubated separately with the reduced and the oxidized forms of the protein. Values obtained from these parallel experiments were within 1 mV of each other, showing that equilibrium had been closely approached. A value of \(-240\) mV was used for the standard redox potential of glutathione at pH 7.0 to calculate the standard redox potential of DsBh (41).

Insulin Turbidity Assay—The ability of DsBh to catalyze the reduction of bovine insulin was tested using an insulin turbidity assay. The reaction mixture, containing 131 \(\mu\)M insulin, 0.35 \(\mu\)M dithiothreitol, and 1, 5, or 8 \(\mu\)M reduced protein (thioredoxin and DsBh) and the insulin stock were prepared in 0.1 M sodium phosphate buffer containing 0.05 M potassium phosphate buffer, pH 7, 2 \(\mu\)M EDTA. The final volume was 1 ml, and the reaction was started by adding insulin.

The measurements were performed at 650 nm. The uncatalyzed reduction of insulin by dithiothreitol was monitored in a control reaction without adding reduced protein.

Measurement of Oxidase Activity by Stopped Flow—Stopped flow fluorescence measurements were performed on a KinTek SF-2004 instrument in single-mixing mode. The typical reaction contained 0.5 \(\mu\)M oxidized protein (DsBh, thioredoxin, and DsBh) and 2.5, 5, 7.5, and 10 \(\mu\)M reduced hirudin, respectively. The oxidized proteins and reduced hirudin were incubated in 100 \(\mu\)M sodium phosphate buffer containing 1 \(\mu\)M EDTA, pH 7.0, at 25 °C before mixing. An excitation wavelength of 295 nm and a bandpass filter were used to monitor the fluorescence change of the proteins.

Isomerization of Scrambled Hirudin—21 \(\mu\)M scrambled hirudin and 21 \(\mu\)M freshly reduced protein (DsBh, E. coli DsbC, and thioredoxin) were incubated in 20 \(\mu\)M sodium phosphate, 130 \(\mu\)M sodium chloride, 0.13% polyethylene glycol 8000 at 25 °C. Aliquots of 120 \(\mu\)l were removed after different reaction times and quenched with 15 \(\mu\)l of formic acid and 15 \(\mu\)l of acetonitrile. Hirudin folding intermediates were separated by reverse-phase HPLC on a C18 column at 55 °C in a 19−25% acetonitrile gradient in 0.1% (v/v) trifluoroacetic acid. The absorbance was recorded at 220 nm.

Spot Titer for Cadmium and Copper Resistance—Spot titers for cadmium resistance were performed to assay the ability of DsBh to complement DsBh. For this assay, the strain JP120 (supplemental Table 2) was transformed with the periplasmic DsBh expression vector (pET44-DsBh) by electroporation. Briefly, mid-log phase cells \((A_{600} = 1)\) were serially diluted in sterile 170 mM NaCl solution. 2 \(\mu\)l of each dilution was plated onto LB plates supplemented with CdCl\(_2\) in a concentration of 5, 6, and 7.5 \(\mu\)M. After 16 h at 37 °C, the growth of each strain at different cadmium concentrations was compared, and the cad-
 TABLE 1  

| Locus tag | CpB0685 | CpB0234 | CpB0233 | CpB0814 | CpB0958 | CpB0968 |
|-----------|---------|---------|---------|---------|---------|---------|
| GenBank accession no. | NP_876957 | NP_876510 | NP_876509 | NP_877086 | NP_877230 | NP_877239 |
| Length | 102 | 233 | 137 | 714 | 166 | 348 |
| Assignment | Thioredoxin | DsbA | DsbB | DsbD | DsbH | DsbJ |
| PSI-Blat E-value to *E. coli* orthologue (third iteration) | 3 × 10^-43 | 4 × 10^-12 | 2 × 10^-11 | 6 × 10^-70 | No clear orthologue detected | No clear orthologue detected |

* Locus tag refers to strain TW-183 of *C. pneumoniae*.

**RESULTS AND DISCUSSION**

Functional Gene Assignment of Chlamydial Dsb Oxidoreductase pathway proteins—The amino acid sequences of the five predicted periplasmic oxidoreductases of *Chlamydia* (8) were submitted to position-specific iterative BLAST searches to identify homologues to Dsb proteins. Chlamydial orthologs of DsbA, DsbC, and DsbD were identified this way (Table 1). However, the remaining two proteins, although distantly related to a number of thioredoxin family proteins, were not closely related to any described Dsb protein including DsbA, DsbB, DsbC, DsbD, DsbE, DsbG, or any other thioredoxin-related protein of known function. Thus, they appear to represent new subfamilies of thioredoxin related proteins. Because the common function of thioredoxin-related proteins is thiol-disulfide exchange, the two proteins are termed DsbH and DsbJ, respectively (Table 1), to highlight their putative involvement in periplasmic disulfide bond manipulation. It is further noted that for both proteins signal peptides are predicted with very high probabilities of 0.987 and 0.999 using the program SignalP 3.0 (supplemental Fig. 1) (20), clearly indicating the periplasmic localization of DsbH and DsbJ.

Significant similarities to DsbH can be detected using position-specific iterative BLAST searches for a wide variety of thioredoxin-related proteins, including DUF255 (a thioredoxin-related protein family of unknown function), the C-terminal domain of DsbD (DsbDy), and somewhat more distantly, to thioredoxin itself (Fig. 1). It is noted that the term DsbH has previously been used in GenBankTM for proteins found in *Pseudomonas* (e.g.YP_001171076) and *Chromobacter*, but these proteins are clearly DsbB homologues and, thus, do not represent a new Dsb protein family. To learn more about the new DsbH family of thioredoxin-related periplasmic oxidoreductases, DsbH from *C. pneumoniae* is structurally and functionally characterized.

**DsbH Structure Determination**—The solution structure of oxidized DsbH was determined by multidimensional heteronuclear NMR spectroscopy, making use of uniformly 13C,15N-labeled protein. As reflected by its rotational correlation time of 10.0 ns and gel filtration profile (supplemental Fig. 2), DsbH is a monomer in solution. A wealth of local structural information on interproton distances and dihedral angles was provided by NMR observables such as NOEs, coupling constants, and chemical shifts. In addition, an extensive set of residual dipolar couplings was used to improve the spatial relationship of more remote protein segments by relating the respective internuclear bond vector orientations to a common molecular alignment frame. The final ensemble of structures was calculated by simulated annealing on the basis of a total of 3410 experimental NMR restraints. Fig. 2 shows a superposition of the final ensemble of 20 simulated annealing structures for which a coordinate precision of 0.69 Å was obtained for all heavy atoms (in the well-folded DsbH domain). A summary of the structural statistics is provided in supplemental Table 1.

**Description of the Structure**—In accordance with its putative disulfide oxidoreductase role, residues Gln-20—Arg-144 of...
DsbH adopt a thioredoxin fold (Fig. 2, C and D). Residues Ala-1 to Ala-19, which follow the predicted signal peptide cleavage site, are unfolded (Fig. 2B). Overall, five $\alpha$-helices surround five central $\beta$-strands in the DsbH domain. The first $\beta$-strand ($\beta1$) is almost perpendicular to the other $\beta$-sheets and consists of only two residues, Trp-23—Glu-24 (Fig. 2D); however, Leu-21—Asn-22 are also in extended conformations, and Leu-21 is intimately connected to the hydrophobic protein interior as shown by methyl-methyl NOEs to Leu-61 of the $\alpha2$—$\alpha2'$ helix connector and Met-76 of strand $\beta3$. The active site disulfide bond, Cys-50—Cys-53, is located at the beginning of helix $\alpha3$, which is typical for thioredoxin-like oxidoreductases (43). Likewise, the orientation of the longest helix, $\alpha3$, is striking in the sense that it is close to perpendicular to the central $\beta$-sheet as well as all other $\alpha$-helices. $\beta$-Sheets $\beta2$ to $\beta4$ are moderately twisted relative to one another, but the $\beta4$—$\beta5$ pair shows a conspicuous twist (Fig. 2D). Strand $\beta5$, comprising Gln-119—Met-123, runs along the $\alpha4$ helix axis accompanied by mainly hydrophobic side-chain contacts, as evidenced by, for example, methyl-methyl NOEs between Met-123 and Val-137 and Ala-121 and Leu-141. The twist at the end of $\beta5$ leads to a suboptimal hydrogen bond geometries for the four interstrand hydrogen bonds between Leu-110 and Met-123 and Phe-112 and Ala-121. The intervening Arg-122 is fully solvent-exposed with the availability of water molecules as hydrogen-bond partners. Backbone dynamics on a fast timescale (pico- to nanosecond), which is indicative of secondary structure stability, identifies distinctive dynamics for the unfolded first 19 residues (as expected) and slightly elevated dynamics for Lys-84—Gln-87 and Gly-106 (Fig. 2B). These residues lie in the $\alpha3$—$\beta3$ and $\alpha3$—$\beta4$ connectors, respectively.

Similarities to Other Oxidoreductases—When comparing the structure of DsbH to other structures in the Protein Data Bank code, DsbH is most similar to the thioredoxin domain present in the C-terminal domain of DsbD (DsbD$_{\gamma}$; Z-score = 12.9). Structurally, DsbH is also highly similar to thioredoxin itself (Z-score = 11.2 for E. coli thioredoxin). In fact, the topology of DsbH is identical to thioredoxin, whereas DsbD$_{\gamma}$ contains an additional helix in exchange for strand $\beta5$ (Fig. 3). Interestingly, unlike in thioredoxin, strand $\beta5$ of DsbH is not well connected to its central $\beta$-sheet, as discussed above. The change in strand $\beta5$ may, therefore, represent a meaningful evolutionary change over the original thioredoxin conformation. The higher structural similarity of DsbH to DsbD$_{\gamma}$ arises from the orientation and length of the $\alpha$-helices in DsbH, which is a second notable change of both proteins compared with thioredoxin (Fig. 3, B and C). The increase in length from E. coli thioredoxin (109 residues) to the (folded) DsbH domain (125 residues) has been mainly incorporated into the $\alpha$-helices, which now have a minimum of three turns. The “break” of the kinked thioredoxin helix $\alpha2$ into two shorter helices, $\alpha2$ and $\alpha2'$, in DsbH and DsbD$_{\gamma}$ is also noteworthy (Fig. 3B). It allows helix $\alpha2'$ to change its orientation relative to the central $\beta$-sheet and allows helix $\alpha2$ to adopt a longer linear segment (44).

The high structural homology between DsbH, DsbD$_{\gamma}$, and thioredoxin exists despite the fact that the DsbH-thioredoxin and DsbH-DsbD$_{\gamma}$ pairs only share sequence identities of 18.9 and 16.5%, respectively. The levels of structural and sequence identity between DsbH, DsbD$_{\gamma}$, and thioredoxin are similar to that found between DsbA and thioredoxin but much lower than the 43.5% sequence identity found between E. coli thioredoxin and its chlamydial orthologue (Table 1). Thus, although DsbH, DsbD$_{\gamma}$, and thioredoxin are clearly structurally related, they are...
Chlamydia does possess a full-length DsbD orthologue (Table 1), so DsbH is not orthologous to DsbD. In addition, DsbH lacks the membrane-embedded β domain and the periplasmic α domain found in DsbD. Thus, it is unlikely to perform the function of DsbD, namely, the transduction of reducing equivalents from the cytoplasm-located thioredoxin to a periplasmic thioredoxin-related protein such as DsbC. Rather, DsbH similarity to DsbDγ may reflect the evolutionary origin of DsbH. DsbDγ may have provided a convenient evolutionary progenitor for DsbH. The close structural similarity of DsbH, DsbDγ, and thioredoxin does not necessarily imply their functional similarity since the redox potential is critically dependent on the two central residues of the CXXC motif (45), which differ substantially among the three proteins (Fig. 1).

Functional Characterization of DsbH—To learn more about the functional characteristics of DsbH, its redox potential was measured, and its catalytic properties were compared with the oxidase DsbA, the reductase thioredoxin, and the isomerase DsbC. Despite the clear difference in catalytic motifs between DsbH (CMWC) and thioredoxin (CGPC), DsbH exhibits a redox potential of $-269.7 \pm 0.8 \text{ mV}$, which is virtually identical to thioredoxin, at $-270 \text{ mV}$ (46). DsbH is also indistinguishable from thioredoxin in its ability to reduce insulin 	extit{in vitro} (Fig. 4A), suggesting that DsbH, like thioredoxin, functions as a reducing thiol disulfide oxidoreductase 	extit{in vitro}. To test for the ability of DsbH to function as an oxidizing thiol-disulfide oxidoreductase, the rate of hirudin oxidation by DsbH was measured, and its ability to complement the disulfide oxidase DsbA for motility and cadmium resistance was evaluated. DsbH is essentially unable to oxidize hirudin in comparison to DsbA (Fig. 4B) and fails to complement DsbA in cadmium resistance and motility assays (supplemental Tables 2–4). The isomerase activity of DsbH was tested by measuring its ability to isomerize the disulfides in misoxidized hirudin. The activity of DsbH in this assay was identical to the very weak isomerase activity exhibited by thioredoxin (data not shown), which is much weaker than the activity of the known isomerase DsbC (42). In addition, DsbH was unable to complement for DsbC in copper resistance assays (Table 2), showing the absence of isomerase functionality also 	extit{in vivo}. In addition, dimerization is known to be very important for disulfide isomerase activity (47, 48), but DsbH is a monomer in solution. Thus, in accordance with its redox potential, DsbH appears to be a periplasmic reductase.
reductases thioredoxin and DsbDy followed by the evaluation of a DsbH-peptide ligand interaction.

**DsbH Active Site Characteristics**—The catalytic disulfide bond of DsbH is surrounded by six key residues: Tyr-49, Met-51, Trp-52, Ile-54, Phe-107, and Pro-108 (Fig. 5, A and B). The functional relevance of some of these residues is reflected by the presence of the residue in DsbH with a disulfide bond (Fig. 2), which makes it the only aromatic residue in DsbH with a side-chain angle close to +60°, and Pro-108 is in cis conformation. Moreover, the geometry of the catalytic Cys-50—Cys-53 disulfide falls in the +/−RHHook group, which classifies it as “catalytic,” like the disulfide of thioredoxin and DsbDy (49). The active site backbone geometries of DsbH, DsbDy, and thioredoxin are highly homologous (Fig. 5, A and B, and supplemental Fig. 3) and, therefore, allow a comparison of these key residues among the three proteins. The structurally homologous active site residues for DsbDy are Trp-460, Val-462, Ala-463, Lys-465, Leu-510, and Pro-511, and for thioredoxin they are Trp-31, Gly-33, Pro-34, Lys-36, Ile-75, and Pro-76 (Fig. 5C). Again, both prolines are cis conformation, and the Cys of Trp-460 and Trp-31 are +36° and +41°, respectively, similar to Trp-49 of DsbH.

The largest difference among the active site residues is seen in the two central residues of the catalytic motifs, which reside on the solvent-exposed side of helix α2 (Fig. 5, A and B). Compared with Met-51—Trp-52 of DsbH, Val-481—Ala-482 of DsbDy as well as Gly-33—Pro-34 of thioredoxin are small and pose less steric hurdles upon approaching the catalytic disulfide (Fig. 5, A and B, and supplemental Fig. 3). In place of the charged Kys-484 and Lys-36 residues of DsbDy and thioredoxin, the hydrophobic Ile-54 residue is inserted in DsbH (Fig. 5C), making its active site residues exclusively hydrophobic and bulky. Phe-107 of DsbH is also slightly more protective of the catalytic disulfide than Leu-510 and Ile-75 of DsbDy and thioredoxin, respectively (Fig. 5, A and B, and supplemental Fig. 3). Taken together, it appears that the readily accessible active site disulfide of thioredoxin is most interesting in productive substrate encounters, whereas substrate approach to DsbH is sterically more hindered and more geared toward hydrophobic substrates. DsbDy appears intermediate between thioredoxin and DsbH. Finally, there is one more residue in the vicinity of the active site that is noteworthy; that is, Gly-106 of DsbH, which is also preserved in DsbDy, thioredoxin, and even DUF255 (Figs. 5C and 1). In addition to being the smallest and conformationally most flexible residue, Gly-106 shows elevated backbone dynamics in DsbH (Fig. 2B). Conformational flexibility at Gly-106 may be of importance during catalysis when the catalytic disulfide engages a substrate and Phe-107 and Pro-108 may have to move out of the way (Fig. 5, A and B).

**DsbH Interaction with a Peptide Ligand**—The large nature of the prominent Met-51 and Trp-52 side chains of DsbH (Fig. 5, A and B) sug-

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**TABLE 2**

| Rank order copper resistance |
|----------------------------|
| Strain | Rank order |
| SQ416 | 2 |
| SQ416 pET44-DsbH | 2 |
| SQ451 | 4 |
| SQ451 pET44-DsbH | 4 |
| SQ417 | 6 |
| SQ453 | 7 |

* Strain descriptions are given in supplemental Table 2.
* Copper resistance was ranked 1–7, with higher values indicating increased resistance.

**FIGURE 5.** Comparison of DsbH and thioredoxin active site geometries and DsbH interaction with a peptide ligand. A and B, superposition of the active sites of DsbH (green) and *E. coli* thioredoxin (Trx; cyan; Protein Data Bank code 2TRX, chain A). Key amino acid residues are shown in ball-and-stick representation and are labeled for DsbH. Lys-36 of thioredoxin is omitted for clarity. C, sequence alignment of the active site residues. D, interaction of DsbH with the depicted peptide ligand encompassing two disulfide bond motifs. The DsbH backbone N-H heteronuclear single quantum coherence signal intensities, $I_{NH}$, are labeled for DsbH. The unfolded DsbH residues (Ala-1—Ala-19), whose motions are uncoupled from the DsbH backbone, are shown in green; the DsbH backbone is shown in blue. The large nature of the prominent Met-51 and Trp-52 side chains of DsbH (Fig. 5, A and B) sug-

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**FIGURE 6.** Comparison of DsbH and thioredoxin active site geometries and DsbH interaction with a peptide ligand. A and B, superposition of the active sites of DsbH (green) and *E. coli* thioredoxin (Trx; cyan; Protein Data Bank code 2TRX, chain A). Key amino acid residues are shown in ball-and-stick representation and are labeled for DsbH. Lys-36 of thioredoxin is omitted for clarity. C, sequence alignment of the active site residues. D, interaction of DsbH with the depicted peptide ligand encompassing two disulfide bond motifs. The DsbH backbone N-H heteronuclear single quantum coherence signal intensities, $I_{NH}$, are labeled for DsbH. The unfolded DsbH residues (Ala-1—Ala-19), whose motions are uncoupled from the DsbH backbone, are shown in green; the DsbH backbone is shown in blue. The large nature of the prominent Met-51 and Trp-52 side chains of DsbH (Fig. 5, A and B) sug-

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**FIGURE 7.** Comparison of DsbH and thioredoxin active site geometries and DsbH interaction with a peptide ligand. A and B, superposition of the active sites of DsbH (green) and *E. coli* thioredoxin (Trx; cyan; Protein Data Bank code 2TRX, chain A). Key amino acid residues are shown in ball-and-stick representation and are labeled for DsbH. Lys-36 of thioredoxin is omitted for clarity. C, sequence alignment of the active site residues. D, interaction of DsbH with the depicted peptide ligand encompassing two disulfide bond motifs. The DsbH backbone N-H heteronuclear single quantum coherence signal intensities, $I_{NH}$, are labeled for DsbH. The unfolded DsbH residues (Ala-1—Ala-19), whose motions are uncoupled from the DsbH backbone, are shown in green; the DsbH backbone is shown in blue. The large nature of the prominent Met-51 and Trp-52 side chains of DsbH (Fig. 5, A and B) sug-

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**FIGURE 8.** Comparison of DsbH and thioredoxin active site geometries and DsbH interaction with a peptide ligand. A and B, superposition of the active sites of DsbH (green) and *E. coli* thioredoxin (Trx; cyan; Protein Data Bank code 2TRX, chain A). Key amino acid residues are shown in ball-and-stick representation and are labeled for DsbH. Lys-36 of thioredoxin is omitted for clarity. C, sequence alignment of the active site residues. D, interaction of DsbH with the depicted peptide ligand encompassing two disulfide bond motifs. The DsbH backbone N-H heteronuclear single quantum coherence signal intensities, $I_{NH}$, are labeled for DsbH. The unfolded DsbH residues (Ala-1—Ala-19), whose motions are uncoupled from the DsbH backbone, are shown in green; the DsbH backbone is shown in blue. The large nature of the prominent Met-51 and Trp-52 side chains of DsbH (Fig. 5, A and B) sug-
gests that any ligand approaching the catalytic disulfide would have to contact these two residues. A peptide that encompasses two disulfide bond motifs (Fig. 5D) was used to test this hypothesis. The peptide was not designed to specifically complement the DsbH active site, which is expected to reduce any (accessible) disulfide bond with reaction kinetics that are modulated by the specific disulfide bond environment of the ligand. Pursuant to our hypothesis, the peptide selectively perturbs the Met-51 and Trp-52 resonances of DsbH (Fig. 5D), demonstrating that these two residues are important for enzyme–substrate interactions. In the absence of an actual catalytic turnover, the interaction between the two oxidized partners is overall weak as judged from the modest effects on the NMR spectrum (Fig. 5D).

For thioredoxin, three structures of ligand–protein complexes are available (50–52). In all three cases the complexes are stabilized by an intermolecular disulfide bond and, therefore, represent a catalytic intermediate rather than the ligand approach to and departure from the active site observed here. Nevertheless, the structures suggest that the two rather small central residues of the thioredoxin catalytic motif, CGPC, do not substantially contribute to thioredoxin–substrate interactions. Thus, the central two residues of the dithiol oxidoreductase catalytic CXXC motif are not only crucial for determining the redox potential and reactivity of an oxidoreductase (45) but can also exert control over enzyme–substrate interactions, i.e. influence substrate selectivity and, thereby, reaction kinetics (Fig. 4). These two factors may explain the rich variety of these residues in the different oxidoreductase families (43, 52) of which each is adapted to its own set of substrates and regeneration partner.

Functional Role of DsbH in Chlamydia—Three functions appear conceivable for a periplasmic reductase in Chlamydia. DsbH may reduce mixed disulfides between proteins and small molecules in the periplasm. Another possible function is to keep a specific subset of periplasmic proteins in the reduced state. Third, DsbH may be responsible for keeping the thiol disulfide status of the periplasm generally reducing. Which of the three outlined scenarios is most probable?

Although the small molecule thiol content of the Chlamydia periplasm is unknown, a wide variety of thiol-containing compounds is present in the periplasm of E. coli (53). Recently, an ATP binding cassette-type transporter has been identified in E. coli that can mediate GSH and cysteine transport into the periplasm (54, 55). This means that, at least in the E. coli periplasm, there is the potential to form mixed disulfides with small molecules.

Cysteine residues can be involved in the ligation of a number of cofactors including hemes, iron sulfur clusters, and metal ions. To be functional in these ligation reactions, the cysteines need to be in the reduced form. Consequently, DsbH may be involved in the specific reduction of cysteine residues in cofactor-containing periplasmic proteins and membrane proteins with periplasmic domains. A reducing function of this type has already been suggested for the DsbE/TlpA/HeIX family of inner membrane proteins (56). DsbH is only distantly related to DsbE/TlpA/HeIX family members, and DsbH is also different from them in that it lacks a membrane-anchoring domain.

Finally, DsbH may be responsible for keeping a generally reducing periplasm. Such a scenario would be reflected in the ratio of even to odd cysteine residues for all chlamydial periplasmic proteins. This can be attributed to the tendency of unpaired cysteines to form incorrect disulfide bonds in an oxidizing milieu and the resulting expectation that, for example, E. coli with an oxidizing periplasm favors periplasmic proteins with even numbers of cysteines. E. coli has a ratio of even to odd numbers of cysteines of 1.58 (Table 3) fulfilling the expectation. In Chlamydia, however, there are more secreted proteins with an odd number of cysteines than an even number, resulting in an even to odd ratio of 0.75 (Table 3). This reversal of the balance of even to odd cysteines correlates with the emergence of DsbH in C. pneumoniae and suggests that Chlamydia do not oxidize periplasmic cysteines to disulfides. Instead, Chlamydia may keep their cysteine-containing periplasmic proteins in the reduced state.

It should be noted that Chlamydia do contain DsbA and DsbB orthologues (Table 1) that are expected to function to oxidize protein cysteines. However, these DsbAB homologues may function in a specialized role in oxidizing outer envelope cysteine rich periplasmic proteins to form a peptidoglycan equivalent. A generally reducing periplasm may benefit Chlamydia in their specific environment, which is intracellular. As an obligate organism, a reducing periplasm may represent an adaptation to the reducing cytosol of the host. In conclusion, although there is no direct evidence for the role of DsbH nor for why the periplasm of Chlamydia has a higher number of proteins with an even number of cysteines than many other bacteria, an attractive possibility is that DsbH might be responsible for maintaining a reducing periplasm.

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TABLE 3
Cysteine distribution in secreted proteins of E. coli and C. pneumoniae (comprising predicted transmembrane and secreted proteins)
The reported numbers are for E. coli strain K12 and C. pneumoniae strain TW-183. Other strains yield similar numbers.

| Protein          | No. proteins (%) with an even number of cysteines | No. proteins (%) with an odd number of cysteines | Ratio even/odd |
|------------------|-----------------------------------------------|-----------------------------------------------|---------------|
| E. coli          | 720 (61%)                                      | 454 (39%)                                      | 1.58          |
| C. pneumoniae    | 125 (43%)                                      | 167 (57%)                                      | 0.75          |

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