Crystal Structure and Biophysical Properties of *Bacillus subtilis* BdbD

**AN OXIDIZING THIOL:DISULFIDE OXIDOREDUCTASE CONTAINING A NOVEL METAL SITE**

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Allister Crow, Allison Lewin, Oliver Hecht, Mirja Carlsson Møller, Geoffrey R. Moore, Lars Hederstedt, and Nick E. Le Brun

From the Centre for Molecular and Structural Biochemistry, School of Chemical Sciences and Pharmacy, University of East Anglia, Norwich NR4 7TJ, United Kingdom and the Department of Cell and Organism Biology, University of Lund, Lund SE-22362, Sweden

BdbD is a thiol-disulfide oxidoreductase (TDOR) from *Bacillus subtilis* that functions to introduce disulfide bonds in substrate proteins/peptides on the outside of the cytoplasmic membrane and, as such, plays a key role in disulfide bond management. Here we demonstrate that the protein is membrane-associated in *B. subtilis* and present the crystal structure of the soluble part of the protein lacking its membrane anchor. This reveals that BdbD is similar in structure to *Escherichia coli* DsbA, with a thioredoxin-like domain with an inserted helical domain. A major difference, however, is the presence in BdbD of a metallocsite, fully occupied by Ca\(^{2+}\), some 14 Å away from the CXXC active site. The midpoint reduction potential of soluble BdbD was determined as −75 mV versus normal hydrogen electrode, and the active site N-terminal cysteine thiol was shown to have a low pK\(_a\) consistent with BdbD being an oxidizing TDOR. Equilibrium unfolding studies revealed that the oxidizing power of the protein is based on the instability introduced by the disulfide bond in the oxidized form. The crystal structure of Ca\(^{2+}\)-depleted BdbD showed that the protein remained folded, with only minor conformational changes. However, the reduced form of Ca\(^{2+}\)-depleted BdbD was significantly less stable than reduced Ca\(^{2+}\)-containing protein, and the midpoint reduction potential was shifted by approximately −20 mV, suggesting that Ca\(^{2+}\) functions to boost the oxidizing power of the protein. Finally, we demonstrate that electron exchange does not occur between BdbD and *B. subtilis* ResA, a low potential extra-cytoplasmic TDOR.

Disulfide bonds, formed upon oxidation of two cysteine residues side chain thiols, are key for the stability and/or function of many secreted and membrane-bound peptides and proteins in bacteria, and the failure to insert these correctly has wide ranging effects (1–4). To regulate the redox state of cysteine residues on the outside of the cytoplasmic membrane, intricate disulfide bond regulatory systems have evolved. These involve enzymes of the thiol-disulfide oxidoreductase (TDOR) family, which contain cysteine residues often arranged in a Cys-Xaa-Cys thioredoxin motif (5, 6). These enzymes function in pathways that lead to the formation of disulfide bonds, rearrangement of incorrectly positioned disulfide bonds, or the removal of unwanted disulfide bonds, and the redox properties of the enzymes appear to correlate closely with function.

The paradigm system for disulfide bond formation is the DsbA-DsbB system of *Escherichia coli*, which has been characterized in great detail (7–11). DsbA is a soluble periplasmic TDOR, which has a thioredoxin-like fold with an additional helical domain (8). The protein oxidizes the di-thiol motifs of a range of substrates, generating in each a disulfide bond. Reduced DsbA is rapidly re-oxidized by DsbB, a membrane-bound TDOR (12) that channels the resulting electrons into the membrane quinol pool (13, 14). Dsb-like homologues appear to be extremely widespread in Gram-negative bacteria.

Gram-positive bacteria, unlike Gram-negatives, do not have an outer membrane and so have no spatially defined periplasmic compartment. This places different functional and structural demands on extra-cytoplasmic proteins, and one consequence of this is that extra-cytoplasmic TDORs are generally membrane-anchored in Gram-positive bacteria. Systems for the introduction of disulfide bonds appear to be variable in these organisms (15, 16). As an example, *Mycobacterium tuberculosis* does not contain close homologues of DsbA/B but contains another thioredoxin-like TDOR, DsbE, which has been shown to have redox properties similar to those of DsbA (17). Some Gram-positive bacteria, including *Staphylococcus aureus*, contain a DsbA homologue but no homologue of DsbB (18). The recent structural and biochemical characterization of *S. aureus* DsbA revealed major similarities with the *E. coli* pro-

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9 [Author’s Choice—Final version full access.]

The atomic coordinates and structure factors (codes 3EU3, 3EU4, 3GHA, and 3GHA) have been deposited in the Protein Data Bank,Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http://www.rcsb.org/).

3 The abbreviations used are: TDOR, thiol:disulfide oxidoreductase; badan, 6-bromooctyl-2-dimethylaminothiophenol; Chex, N-cyclohexyl-2-ami-nothanesulfonic acid; DTT, dithiothreitol; ICP-AE, inductively coupled plasma-atomic emission; Mes, 2-(N-morpholino)ethanesulfonate; MALDI-TOF, Matrix-assisted laser desorption/ionization-time of flight; Mops, 3-(morpholino)propanesulfonate; PEG, polyethylene glycol; HSQC, heteronuclear single quantum coherence; NHE, normal hydrogen electrode.
tein but was also consistent with a distinct mechanism of re-oxidation (16).

Some Gram-positive bacteria, however, contain clear homologues of both DsbA and DsbB. In the model organism Bacillus subtilis, BdB and BdbC/BdB have been identified as homologues of DsbA and DsbB, respectively, and demonstrated to be involved in processes such as natural competence development, which requires the insertion of disulfide bonds (19). BdB and BdbC are also involved in a number of other pathways that do not require the insertion of a disulfide bond; B. subtilis contains several extra-lysosomal TDORs, for example RsA and StoA (required for cytochrome c maturation and endosome biogenesis, respectively), which function to specifically reduce disulfide bonds introduced by BdB (20, 21). To understand disulfide bond management systems in Gram-positive bacteria, detailed information on each of the different systems found is required.

Here we present the crystal structure of the catalytic domain of B. subtilis BdB and in both reduced and oxidized states. The structure is broadly similar to that of E. coli DsbA, but it also reveals the presence of a novel Ca\(^{2+}\)-binding site remote from the CXXC active site. Using two-dimensional NMR methods and fluorescence kinetic studies of thiolate alklylation, we report the reduction potential and pK\(_a\) properties of the soluble protein, which are entirely consistent with an oxidizing function for the protein in vivo. Conformational stability studies and NMR studies showed that the occupancy of the metal site by Ca\(^{2+}\) ion is not required for folding/stability but leads to a significant increase of the midpoint reduction potential. The possibility that the principal function of the metal site is to boost the oxidizing power of the protein is discussed.

**EXPERIMENTAL PROCEDURES**

**Bacterial Strains and Plasmids—**B. subtilis strains 1A1 (trpC2) and LUL10 (trpC2 bdbD/Tn10) (20) were used for production of extracts for immunoblot analysis. E. coli TOP10 (F\(^-\) mcrA Δ[mrr-hsdRMS-merC] F80lacZΔM15 ΔlacX74 recA1 araD139 Δ[ara-leu]7697 galU galK rpsL endA1 nupG) (Invitrogen) and B834(DE3) (F\(^-\) ompT hsdSB (r\(_{B}\) m\(_{B}\) gal dcm met) (DE3)) (Novagen) containing pLLE34 (22) or pACN2 was used for production of sBdB. pACN2 was constructed by PCR amplifying part of the bdbD gene using B. subtilis 1A1 genomic DNA as template and two primers (forward, 5′-A CAT ATG AAT AAC AAA ACG GAA CAA GGC 3′; reverse, 5′-CTC GAG CTT CCC TTT CAG CTC TTT TCC 3′; engineered Ndel and Xhol restriction sites are underlined). The resulting PCR product was blunt-end ligated into pUC18, and the Ndel and Xhol fragment was subsequently ligated into pET21a (Novagen) digested with the same enzymes and treated with shrimp alkaline phosphatase, and the construct was verified by DNA sequencing (MWG Biotech). Growth of B. subtilis strains and preparation of cell extracts for immunoblot analysis were done as recently described for StoA (23).

**BdB Antiserum and Immunoblot—**sBdB was produced in E. coli TOP10/pLLE34 and purified as described before (22). BdB antiserum was obtained by immunizing New Zealand White rabbits with purified protein. For immunoblot, proteins were fractionated by SDS-PAGE (24) and transferred from the gel to a polyvinylidene fluoride blotting membrane (Immobilon P, Millipore) using a wet blot and 20 mM Tris, 150 mM glycine buffer containing 20% (v/v) methanol. Immunodetection was carried out by chemiluminescence using horseradish peroxidase-labeled secondary antibodies against rabbit (GE Healthcare) and Super Signal\textsuperscript{®} West Pico chemiluminescent substrate (Pierce) and a Kodak image station 440 CF for signal recording.

**Production and Purification of sBdB for Crystallization and NMR—**A 23-kDa truncated form of the BdBd protein (sBdB) consisting of residues 30–222 of the full-length BdB protein (lacking the N-terminal transmembrane helix) by an N-terminal methionine residue and a C-terminal His tag with the sequence LEHHHHHH was expressed from pACN2 in E. coli B834 (DE3). Cultures were grown at 37 °C on minimal glucose medium supplemented with either 60 μg liter\(^{-1}\) methionine or 60 μg liter\(^{-1}\) selenomethionine to an A\(_{600}\) of 0.6 and centrifuged at 6,000 × g for 10 min at 4 °C. Cells were resuspended in fresh minimal glucose medium, and expression of the bdbD gene was induced by the addition of 0.5 mM isopropyl β-D-thiogalactoside. Cells were grown for a further 12 h at 37 °C before harvesting at 6,000 × g for 10 min at 4 °C. Cells were broken by sonication, and the cell lysate was centrifuged to remove unbroken cells and debris. The supernatant was applied to a nickel affinity column equilibrated in 100 mM Tris, 500 mM NaN\(_{3}\), pH 8.0, and eluted with 100 mM Tris, pH 8.0, 500 mM NaCl, 250 mM imidazole, pH 8.0. Fractions containing sBdB were then applied to a 5-ml anion exchange column (HiTrap Q, GE Healthcare) equilibrated in 100 mM Tris, 0.5 mM DTT, pH 8.2, and subsequently eluted with a 0–1 M gradient of NaCl in the same buffer. DTT was subsequently removed using a Sephadex G-25 column (PD10, GE Healthcare). MALDI-TOF mass spectrometry was used to confirm that the purified protein matched the predicted mass expected from the sequence. 15\textsuperscript{N}-Labelled sBdB was produced using the same method, but with 15\textsuperscript{NH}_4\text{Cl} (1 g/liter) as the sole nitrogen source.

The UV-visible absorption spectrum of sBdB contained an intense band at 280 nm with an unusually pronounced peak at 290 nm, presumably due to tryptophan. \(\varepsilon_{280}\,\text{nm}\) was determined using the method of Pace et al. (25) as 21,033 M\(^{-1}\) cm\(^{-1}\), and this was used subsequently to determine the protein concentration. Metal ion content of the protein was determined by ICP-AE spectroscopy (Varian Vista); Ca\(^{2+}\) was removed from sBdB by incubating the protein with 10 mM EDTA for 1 h at 25 °C. EDTA was subsequently removed using a PD10 desalting column. EDTA-treated samples were analyzed by ICP-AE and found to contain <0.1 Ca\(^{2+}\) per protein. Oxidized and reduced sBdB samples were prepared by incubation in the dark for 2 h in the presence of either 1 mM diame to 1 mM DTT before removing reduc buffer using a PD10 desalting column.

**Crystallization and Structure Determinations—**Purified sBdB (native or selenomethionine-labeled) was concentrated to 12 mg/mL in 10 mM Mops, pH 7.0, and crystallized using the sitting drop vapor diffusion method. Equal volumes (1 μl) of protein and crystallization reagent (25–32.5% PEG 2000 (w/v), 0.1 mM ammonium acetate, 0.1 mM Mes, pH 6.5) were mixed and equilibrated over an 800-μl reservoir of the crystallization reagent alone. Low quality crystals grown at high concentrations of PEG (e.g. 30–32.5% (w/v)) were broken using a pipette tip and
used for streak-seeding of drops containing lower concentrations of PEG (25–27% (w/v)). Crystals deemed suitable for diffraction were then cryo-protected in a solution composed of 30% (w/v) PEG 2000, 20% (v/v) ethylene glycol, 0.1 M ammonium acetate, 0.1 M Mops, pH 7.0 (with or without 1 mM DTT as required), before being flash-frozen in preparation for data collection.

X-ray data sets for both native and selenomethionine-labeled protein crystals were collected on beam line ID23-1 of the European synchrotron radiation facility. Diffraction data were indexed and integrated with MOSFLM (26) and then scaled with SCALA (27). Phases were determined using BP3 of the CCP4 suite (28), having already identified the selenium sites by manual inspection of Patterson maps. Further phase improvement was accomplished by solvent flattening and histogram matching with DM, and the vast majority of the structure was built automatically using ARP/WARP (29).

Further model building and refinement (including refinement of individual anisotropic b-factors) utilized COOT (30) and REFMAC (31) as part of the CCP4 suite (28). Structure validation was aided by PROCHECK (32), and the final models were submitted to the Protein Data Bank with accession codes 3EU3 (reduced), 3EU4 (oxidized), 3GH9 (EDTA-treated/oxidized), and 3GHA (EDTA-treated/reduced).

Analysis of the metal site was conducted by systematically refining different metals and examining electron density difference maps (i.e., $|F_o| - |F_c|$ maps) for indications of the quality of fit. To quantify the degree of difference density associated with each candidate metal, MAPMAN (33, 34) from the RAVE suite was used to integrate the difference density within a 3.5-Å radius of the metal center. The anomalous scattering characteristics of the metal site were also compared with those of the selenium atoms in our selenomethionine structure. FFT (CCP4 suite) was used to generate an anomalous difference map using data obtained from the “peak” data set of the selenomethionine MAD experiment (with phases from the refined model), and MAPMAN was used to integrate the anomalous density around the unknown metal site and the selenium atom of SeMet$^{187}$. The experimentally determined $f$ value for selenium at this wavelength was 5.55 electrons, and the integrated anomalous density associated with the metal site was 24% of that of the ordered selenium of SeMet$^{187}$, suggesting that the $f$ value of the unknown metal is ~1.33 electrons. The $f$ values of Ca$^{2+}$, K$^+$, and Fe$^{2+}$ were all roughly consistent with the anomalous scattering at the unknown metal site, although other metals such as Na$^+$, Mg$^{2+}$, Ni$^{2+}$, Cu$^{2+}$, and Zn$^{2+}$ were not.

Assignment of the metal site geometry as “capped octahedral” was based on a consideration of all possibilities; angles between each ligand and the central metal were measured from the refined structure and compared with ideal angles calculated for each potential geometry. The root mean square deviation from the ideal angles (root mean square angle) was then used to find the geometry that best represents the site. The root mean square angle values for each possible 7-ligand geometry were as follows: pentagonal bipyramid (18.57°); square face-capped trigonal prism (20.23°); triangle face-capped trigonal prism (32.21°), and capped octahedron (12.15°).

**Reduction Potential Determinations—$^{15}$N-Labeled sBdbD**

(100 μM final protein concentration) in 50 mM potassium phosphate, pH 7, was treated with 10 mM GSSG or with 1 mM diamide (reacted overnight in the dark) to generate the fully oxidized protein or with 10 mM GSH or 5 mM DTT to generate the fully reduced protein. $^1$H $^{15}$N-HSQC spectra of oxidized and reduced sBdbD showed that the exchange between oxidized and reduced proteins was slow on the NMR time scale, and so spectra of mixtures corresponded to a superposition of spectra arising from the individual oxidized and reduced proteins. Several resonances that were dependent on the redox state of the protein were selected for analysis. sBdbD was oxidized at 4 °C overnight with 1 mM diamide, and excess oxidant was removed using a PD10 column and the protein equilibrated in 50 mM potassium phosphate buffer, pH 7. Fully oxidized sBdbD (100 μM final concentration) in the same buffer was then prepared as above, and a final concentration of 10 mM GSSG was added. Increasing concentrations of GSH (from a ratio of GSH:GSSG of 1:50 to 1:1.5) were added, and the samples were incubated at 25 °C. Redox equilibrium was established after 1 h, and $^1$H $^{15}$N-HSQC spectra were recorded for each, and the intensity changes at the selected chemical shift values were monitored. The resonance at $^1$H/$^{15}$N = 9.516/115.587 was used as an intensity standard to minimize variation between samples. Spectra of fully reduced and oxidized sBdbD, which provided the end points of the titration, were recorded following the removal of reductant/oxidant by gel filtration. GSH and GSSG concentrations were calibrated, respectively, by reaction with 5,5′-dithiobis-(2-nitrobenzoic acid), and by absorbance measurements at 248 nm using $\varepsilon_{248 \text{ nm}} = 382$ M$^{-1}$ cm$^{-1}$ (35). The fraction of sBdbD reduced, $f_s$, at each point in the titration and for each selected resonance in the NMR spectrum was calculated, and the midpoint reduction potential of sBdbD was determined as described previously (23, 36); further details are given in the supplemental material.

NMR spectra were acquired using a Bruker Avance III 800 MHz spectrometer equipped with a triple resonance, pulsed field gradient probe, operating at $^1$H frequency of 800.23 MHz and $^{15}$N frequencies of 81.09 MHz, using pulse sequences incorporated into the Bruker Topspin 2.1 software. Two-dimensional $^1$H $^{15}$N-HSQC spectra were recorded at 25 °C. The $^1$H carrier frequency was positioned at the resonance of the water during the experiments. The $^{15}$N carrier frequency was at 115 ppm. Spectra were processed using NMRPipe (37) and peak intensities analyzed in NMRDraw (37). Before Fourier transformation, a cosine-bell window function was applied to each dimension for apodization. The indirect dimensions were first linearly predicted to double the number of data points and then zero-filled to round up the number of data points to the nearest power of 2.

**Determination of pH Stability Range and Cysteine Thiol pK$^a$ Values—For pK$^a$ determination experiments, a mixed buffer system (PCTC buffer: K$_2$HPO$_4$, sodium citrate, Tris, and Ches, at 50 mM) was used to maintain constant pH. To determine the range of pH over which sBdbD was stable (and therefore the range within which pK$^a$ values could be measured), changes in the intrinsic tryptophan fluorescence of sBdbD (0.25 μM final concentration) as a function of pH were followed upon excita-
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at 280 nm. For pKₐ measurements, pre-reduced sBdbD was reacted with 6-bromoacetyl-2-dimethylaminonaphthalene (badan) under pseudo-first order conditions, and analyzed as described previously (38). Further details are given in the supplemental material.

Protein Conformational Stability Measurements—sBdbD (both with and without Ca²⁺) was pre-reduced or oxidized by incubation in the dark for 2.5 h at 4 °C in 2 mM DTT or diamide (39), respectively. Excess reductant/oxidant was removed using a PD10 column equilibrated in 50 mM potassium phosphate, pH 7.0. Equilibrium unfolding experiments were conducted by incubating sBdbD in various quantities of urea in 50 mM phosphate buffer, pH 7.0 (reduced samples included 1 mM DTT to prevent re-oxidation), and monitoring intrinsic tryptophan fluorescence as a probe of the degree of unfolding. Refolding experiments were performed in a similar manner starting with protein that was incubated overnight in 6 M urea. In all cases, sBdbD (0.25 μM final concentration) was allowed to equilibrate at 25 °C for 4 h before measurements were made. Tryptophan fluorescence was measured at 25 °C using an LS55 luminescence spectrophotometer (PerkinElmer Life Sciences) with an excitation wavelength of 290 nm. Fluorescence intensity at 353 nm was used to determine Gibbs free energies of stabilization (ΔGstab) and midpoints of unfolding, as described previously (36); further details are given in the supplemental material.

Other Methods—For experiments involving substrate analogue peptides, pre-oxidized ¹⁵N-labeled sBdbD (300 μM in 50 mM potassium phosphate buffer, pH 7) was mixed with 600 μM reduced peptide and incubated at 25 °C for 1 h before measuring ¹H-¹⁵N-HSQC spectra. Peptides were purchased from Severn Biotech, Kidderminster, UK) and contained the sequences IYKANCIACHGENYE (referred to as CXXCH peptide) and IYKANSIASHGENYE (SXXSH peptide), which correspond to the wild-type sequence and a double Cys to Ser mutation at 280 nm. For MALDI-TOF experiments, 10 mg/ml sinapinic acid in 50% acetonitrile and 0.05% trifluoroacetic acid was used as the matrix. Spectra were recorded using a Shimadzu Biotech AXIMA-CFR instrument operating in linear mode, calibrated using cytochrome c (equine) 12,361.96 Da, apo-myoglobin (equine) 16,952.27 Da, and aldolase (rabbit muscle) 39,212.28 Da (Sigma). The soluble domain of ResA (sResA) was produced in E. coli and purified as described before (21).

RESULTS AND DISCUSSION

BdbD Is Membrane-localized—The B. subtilis bdbD gene encodes a protein of 222 amino acid residues with a putative N-terminal signal peptide for export to the outside of the cytoplasmic membrane. Although BdbD was predicted to have a type I signal peptidase cleavage site between residue 36 and 37 (40), such predictions are unreliable. To establish if BdbD is a membrane-bound protein, B. subtilis strain 1A1 was grown in NSMP medium (nutrient sporulation medium with phosphate). Because BdbD plays roles in both growing and stationary phase cells, samples were taken from the culture at various points in the growth curve. Soluble and membrane fractions of cell lysates were analyzed for BdbD by using immunoblot with polyclonal antiserum directed against BdbD without the putative membrane anchor. BdbD was not detected in the soluble fraction but was present in the membrane fraction, at a level that was largely independent of the growth phase (Fig. 1). Thus, we conclude that the TDOR domain of BdbD is attached to the membrane by the N-terminal transmembrane segment.

High Resolution Crystal Structures of BdbD in Reduced and Oxidized Forms—The soluble domain of BdbD (residues 37–222, sBdbD) was produced in E. coli, purified, and crystallized. The crystal structure of sBdbD was solved using selenomethionine MAD with subsequent refinement against high resolution native data sets obtained for crystals of both oxidized and reduced forms of the protein. Crystals of sBdbD belonged to space group P2₁ with a single molecule in the asymmetric unit. Full data collection and refinement statistics for the oxidized and reduced structures are given in Table 1.

The overall crystal structure of BdbD is shown in Fig. 2A. The fold of BdbD is clearly related to the DsbA family of TDORs, for which a number of proteins have been structurally characterized. Like all DsbA-like proteins, BdbD is composed of two domains, a classical thioredoxin-like domain and an α-helical domain (Fig. 2A, green and teal, respectively). The thioredoxin-like domain consists of residues 59–103 (forming the βαβ component of the fold) and 182–222 (forming the αββα component of the fold). The helical domain consists of residues 104–181 that form five α-helices, the fifth of which is an extension of the first helix of the thioredoxin αββα fold, therefore providing a structurally defined connection between the two domains. Preceding the βαβ component of the thioredoxin fold at the N terminus is an additional β-strand, which forms a β-sheet with the other β-strands of the thioredoxin fold. The arrangement of secondary structure elements is shown in Fig. 2C.

BdbD is most similar to S. aureus DsbA (SaDsbA), sharing a number of structural features with this protein that differ from E. coli DsbA (EcDsbA) (16). For example, BdbD possesses a loop between helices α3 and α4 of the all-helical domain that is considerably longer than the equivalent loop observed in EcDsbA and also lacks the hydrophobic groove observed on the surface of EcDsbA, which was recently shown to be important for interaction of DsbA with DsbB (41). BdbD can be superposed with SaDsbA and EcDsbA with root mean square devia-
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In the structure of reduced sBdbD, the cysteine sulfur atoms are separated by a distance of 3.19 Å (Fig. 3C), with clear indications from the electron density that, despite their close proximity, the cysteine sulfurs are in the free thiol/thiolate forms ("unbiased" omit maps for the oxidized and reduced structures are shown in supplemental Fig. S1A). In contrast, the oxidized structure showed continuous electron density between the sulfur atoms of Cys69 and Cys72 demonstrating the formation of a disulfide bond (Fig. 3B and supplemental Fig. S1B). Although crystallized as a fully oxidized sample, refinement of a disulfide bond in the oxidized structure did show some residual electron density indicating a small fraction of sBdbD was in the reduced state, probably due to a degree of photo-reduction in the x-ray beam. We thus refined the "oxidized" structure as a mixture of oxidized (70%) and reduced states (30%), with the active site dominated by the oxidized form (a more completely oxidized disulfide structure was obtained for a sample treated with EDTA, see below and supplemental Fig. S1C). The structural data show that there are very few redox-linked conformational changes between the oxidized and reduced forms of the protein, such that superposition of oxidized and reduced BdbD gave a root mean square deviation of 0.268 Å (calculated for all atoms of residues 37–221).

Thiolate Form of Cys69 Is Significantly Stabilized Relative to the Thiol Form—The reduction potentials of thioredoxin-like TDORs are intimately linked with the pKₐ values of their active site cysteines: in general, the more extensive the stabilization of the thiolate form of the active site cysteine, the lower their corresponding pKₐ value and the higher the active site reduction potential (43–45). Furthermore, within the thioredoxin-like family of TDORs, it is the N-terminal active site cysteine that exhibits the greatest variation and is thus likely to be the key determinant of reduction potential. In the structure of reduced sBdbD, the Cys69 sulfur atom is hydrogen-bonded to the backbone amide groups of both Ser71 (3.4 Å) and Cys72 (3.2 Å) as well as to the side chain hydroxyl group of Thr192. The extensive hydrogen bonding network around the Cys69 sulfur is clearly an important factor in maintaining a low cysteine pKₐ for Cys69 and is therefore likely to be a key structural determinant of the reduction potential of the protein.

### Table 1

Data collection and refinement statistics

Values in parentheses represent the highest resolution shell. r.m.s. indicates root mean square. Multiplicity and completeness are "anomalous" for seleno-methionine data but not for the native datasets. Rmerge refers to cross-dataset scaling using the reduced native dataset as reference. Free c.c. is the correlation coefficient calculated with the free reflection set.

|                      | Selenomethionine sBdbD | Reduced native | Oxidized native |
|----------------------|------------------------|----------------|-----------------|
| Space group          | P2₁                    |                |                 |
| Unit cell            |                        |                |                 |
| Unique reflections   | 28,058                 | 27,977         | 27,961          |
| Energy               | 12,654 eV               | 12,653 eV      | 12,661 eV       |
| f                    | −9.18                  | −10.41         | −5.52           |
| f'                   | 5.55                   | 3.61           | 4.53            |
| Resolution           | 27.51 to 1.50 Å (1.58 to 1.50 Å) | 28.21 to 1.50 Å (1.58 to 1.50 Å) | 52.2 to 2.3 Å (2.42 to 2.3 Å) |
| Rmerge               | 0.072 (0.444)          | 0.052 (0.267) | 0.086 (0.119)   |
| Rfree                | 1.11 (1.8)             | 15.0 (3.4)     | 15.7 (7.3)      |
| I/σ(I)               | 97.7% (97.8%)          | 97.8% (89.5%)  | 94.8% (81.2%)   |
| Completeness         | 1.8 (1.8)              | 1.9 (1.9)      | 2.3 (2.2)       |
| Multiplicity         | 0.078                  | 0.068          | 0.077           |
| r.m.s. bond          | 0.2051                 | 0.2480         | 0.1937          |
| r.m.s. angle         | 1.225°                 | 1.262°         | 1.232           |
| Free c.c.            | 0.0098 Å               | 0.0090 Å       | 0.8918          |

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| r.m.s. bond          | 0.2051                 | 0.2480         | 0.1937          |
| r.m.s. angle         | 1.225°                 | 1.262°         | 1.232           |
| Free c.c.            | 0.0098 Å               | 0.0090 Å       | 0.8918          |
Structural Implications for the Interaction of BdbD with Substrates

—The loop region directly opposite the active site (composed of Thr^{192} and cis-Pro^{193}) is very likely to be involved in the interaction of BdbD with its substrates (46). This cis-proline residue is conserved among all known thioredoxin-like proteins and is likely to be important in presenting the carboxyl oxygen of the preceding residue for hydrogen bond interactions with substrate cysteine residues as observed in various TDOR-substrate complexes (47–49). Substitution of the DsbA equivalent of BdbD Pro^{193} has been shown previously to lead to an accumulation of covalent DsbA complexes in vivo (50). Furthermore, the crystal structure of EcDsbA in a covalent complex with DsbB has shown that this loop is important in mediating contacts between DsbA and DsbB (41). We propose that similar interactions are likely to be important for the interaction of BdbD with its substrates. However, we note that although BdbC is clearly a homologue of DsbB, the manner by which BdbD is re-oxidized by BdbC cannot be the same as for DsbA/DsbB, because BdbC lacks the external loop domain with which DsbA interacts (20). Furthermore, as noted above, the peptide-binding groove identified in EcDsbA (which appears to interact with a well defined portion of DsbB) does not exist in the structure of BdbD (Fig. 2B). Indeed, the electrostatic surface view of BdbD (Fig. 2B) shows that little of the surface shape or the electrostatic properties are conserved between BdbD and DsbA-like proteins (8, 16).

Role of Glu^{63}—Our structures show that Glu^{63} is buried in a position very close to the Cys^{72} sulfur atom (Fig. 3A). Buried polar or charged residues, which destabilize folding, are usually functionally important, and in support of this, a negative charge at the equivalent position of Glu^{63} is highly conserved among BdbD and DsbA homologues although, interestingly, it is not

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**FIGURE 2. Structure of the soluble domain of B. subtilis BdbD.** A, three-dimensional structure of sBdbD showing that the protein exhibits a thioredoxin-like fold with an inserted helical domain. Secondary structure elements are labeled from the N terminus with the predicted transmembrane helix of the full-length protein (covered by residues 1–36 that were not present in the protein used for crystallization and therefore not indicated on the figure) being 0. The active site and Ca^{2+}-binding sites are indicated. The thioredoxin-like part of BdbD is colored green, and the all-helical domain is colored teal. Note that the same color scheme is used in Figs. 3 and 4. B, surface representation of sBdbD. Regions colored red indicate areas of high negative electrostatic potential, and blue areas indicate areas of high positive potential. Neutral regions are in white. Electrostatic potentials of surfaces were calculated using PyMOL (73). All structural figures were prepared with PyMOL and annotated with GIMP. C, amino acid residue sequence comparison of B. subtilis BdbD (BsBdbD) with other DsbA-like proteins from Bacillus anthracis (BaBdbD), Bacillus cereus (BcBdbD), S. aureus (SaDsbA), and E. coli (EcDsbA). Invariant residues are marked in black and highly conserved residues in gray. Diamonds indicate the active site residues Cys^{69} and Cys^{72}, and asterisks indicate the residues acting as ligands to the Ca^{2+} ion. A black circle indicates the near-active site glutamate residue (Glu^{63}), and an inverted triangle indicates the beginning of the soluble domain of B. subtilis BdbD studied here. Secondary structural elements, derived from the structure of BdbD, are indicated above the sequence. Elements belonging to the thioredoxin-fold are indicated in dark gray; those belonging to the helical domain are indicated in light gray, and those not belonging to either are indicated in white. Residue numbers are indicated on the right; these refer to the non-processed predicted translated sequence. The alignment was obtained using ClustalW (74) and annotated in Genedoc (75).

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**FIGURE 3. Active site region of BdbD.** A, detailed view of the N terminus of helix α1 of sBdbD showing the Cys-Pro-Ser-Cys active site of sBdbD and the closely lying cis-proline (Pro^{193}), which is invariant in all thioredoxin-like proteins. B and C, electron density (contoured at 1.0σ) of the active site region of sBdbD in oxidized and reduced states, respectively.

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Glu63 of BdbD, was proposed to act in just such a way during oredoxin, which is located spatially in a similar position to anomalous scattering of the metal site (at a wavelength of through the investigation of Glu-to-Gln mutants (23, 38). StoA have both been confirmed as mechanistically important TDORs. Furthermore, these glutamate residues of ResA and to the C-terminal active site cysteine is similar in all of these primary sequence, but the location of the negative charge close –The metal ion site located between the thioredoxin, which is located spatially in a similar position to Glu63 of BdbD, was proposed to act in just such a way during interaction with its substrates (51). We note that buried glutamate residues also occur in ResA (Glu80) and StoA (Glu77) which, like BdbD, are extra- cytoplasmic membrane-bound TDORs from B. subtilis (23, 52). The buried glutamates of StoA and ResA are not in equivalent positions to Glu63 in terms of primary sequence, but the location of the negative charge close to the C-terminal active site cysteine is similar in all of these TDORs. Furthermore, these glutamate residues of ResA and StoA have both been confirmed as mechanistically important through the investigation of Glu-to-Gln mutants (23, 38).

BdbD Contains a Ca2+ Ion in a Capped Octahedral Geometry—The metal ion site located between the thioredoxin and helical domains is 14 Å from the nearest active site cysteine sulfur and is coordinated by three protein residues as follows: Glu49, Glu115, and Asp180. In addition to these three residues, the central ion also forms coordinate bonds with three low b-factor solvent molecules (Fig. 4A). The metal therefore has a total of seven ligands (the carboxylate group of Asp180 is bidentate) arranged with a capped octahedral geometry (Fig. 4B). We identified the metal as Ca2+ on the basis of several lines of evidence. First, ICP-AE spectroscopy of purified sBdbD demonstrated the presence of 0.9 ± 0.1 Ca2+ (and no other metal ions) per protein. Second, only Ca2+ and K+ refined well against the x-ray data. Smaller candidate ions such as Na+ or Mg2+ refined to unrealistically low b-factors and gave positive peaks in difference maps ([I - |Fb|], suggesting they are not electron dense enough to fit the observed data. Larger ions such as Zn2+ or Fe2+ also refined poorly, with strong negative density in difference maps suggestive of a smaller metal or a partial occupancy. The latter possibility can be ruled out because the anomalous scattering of the metal site (at a wavelength of 0.9798 Å) was very small; a low occupancy of Zn2+ (or another transition metal) would be expected to give much stronger anomalous scattering. Third, the metal-to-water bond lengths binding site remote from the CXXC active site, raising the question of what the functional significance of the metal site is. Multiple sequence alignments of BdbD/DsbA proteins show that the three residues involved in coordinating the metal site are strictly conserved among all Bacillus species, but not in Gram-negative bacteria nor in other Gram-positive organisms such as S. aureus (Fig. 2C). This suggests that the site is likely to be relevant to the function of BdbD. A second question raised by the structure is whether or not Ca2+ is the native metal ion in B. subtilis. The protein was heterologously produced in the cytoplasm of E. coli cells, and it is possible that Ca2+ was the only ion available in sufficient quantities to fill the site in the absence of an alternative. However, Ca2+ concentrations in bacterial cells, as in eukaryotic cells, are tightly regulated and are estimated to be between 100 and 300 nM (54). Therefore, Ca2+ is not normally found at sites usually occupied by other metal ions. Furthermore, we can rule out that Ca2+ was added during or after purification as it was not used at any stage during the purification or as a component in the crystallization buffer. Therefore, the data indicate that Ca2+ is the natural metal ion in BdbD.

Ca2+ is extremely important for a range of processes in eukaryotic cells. Its role in bacterial cells is much less clear, but there are a number of examples of bacterial Ca2+-binding proteins, which are proposed to be involved in a range of processes, including the maintenance of cell structure and motility, and in cell differentiation processes such as sporulation (55). Many of these are related to eukaryotic EF-hand proteins that contain a conserved helix-loop-helix structural motif in which Ca2+ is coordinated by the side chains of three residues from the loop, a backbone atom from another loop residue, a water molecule, and the side chain of an acidic residue in the second helix (56, 57). The structures of several bacterial EF-hand-like proteins have been determined, revealing that although they share the conserved Ca2+-binding motif DX(D/N)XDG within the central loop of the EF-hand domain, they exhibit significant diversity in the structural elements surrounding the motif. Ca2+ is also found in other types of bacterial proteins, including those belonging to the RTX family of bacterial cytolysins (58) that are
normally secreted, which includes toxins (59) and lipases (60), as well as a lectin (61) and a collagenase (62), which both contain unusual binuclear Ca\(^{2+}\) sites. In all of these structures, Ca\(^{2+}\) is bound in a loop that provides the majority of the ligands, and therefore all are distinct from the site described here in sBdB, in which the three residue side chains acting as ligands are derived from three different parts of the primary sequence (Fig. 2C). A greater degree of similarity is observed for the Ca\(^{2+}\) sites of peroxidases and, in particular, the di-heme cytochrome \(c\) peroxidase, in which Ca\(^{2+}\) is coordinated by three residues and four water molecules and lies at the interface between the two domains of the protein (63).

Ca\(^{2+}\) Site Is Not Required for Folding of BdB—To investigate the importance of the Ca\(^{2+}\) site for the structure of sBdB, the crystal structure of a metal chelator-treated sample was solved in both oxidized and reduced states (data collection and refinement statistics are given in supplemental Table S1). The sample used for crystallization contained <10% Ca\(^{2+}\) according to metal analysis, and the chelator, EDTA, was maintained at 100 \(\mu\)M in all crystallization conditions and cryo-protectants used during preparation of the crystals. Despite this, some residual electron density was present within the metal site of both the oxidized and reduced EDTA-treated sBdB structures, although this was substantially less than in the as-isolated sBdB. Overall structures of reduced and oxidized chelator-treated sBdB were very similar to those of the as-isolated protein (see supplemental Fig. S2). There were several minor changes to the local structure of residues at the site, including Glu\(^{115}\), which adopted multiple distinct side chain conformations consistent with heterogeneity at the site. Furthermore, there was essentially no anomalous scattering at the position of the metal. Best fits to the data were obtained using either a low occupancy Ca\(^{2+}\) ion or a full occupancy Na\(^{+}\) ion. Given that EDTA-treated sBdB contained some Ca\(^{2+}\), at least some of the observed intensity must arise from Ca\(^{2+}\) occupation, but it is possible that Na\(^{+}\) replaces Ca\(^{2+}\) in some BdB molecules. We note that other metals (such as K\(^{+}\)) can be ruled out on the basis of the metal-to-water coordination distances that are consistent with occupation by Na\(^{+}\) or Ca\(^{2+}\). Although it has not yet been possible to completely remove all traces of metal(s) from the observed site, it is nonetheless clear that occupation by Ca\(^{2+}\) is drastically reduced in the EDTA-treated structures. This indicates that Ca\(^{2+}\) is not required to maintain the native three-dimensional fold of the protein.

Two-dimensional NMR methods were also used to investigate conformational changes upon removal of Ca\(^{2+}\). The \(^{1}H\) \(^{15}N\)-HSQC spectrum of reduced \(^{15}N\)-labeled sBdB was recorded before and after treatment with EDTA (Fig. 5). A significant proportion of resonances was observed to have shifted upon depletion of Ca\(^{2+}\), indicating that the presence of the ion does have structural consequences in solution. Similar results were obtained for the oxidized form of the protein (data not shown). In light of the crystal structure of Ca\(^{2+}\)-depleted sBdB, we conclude that these changes must be subtle (for example, similar to those caused by an oxidation state change, which do not cause major structural rearrangement but result in shifts in many resonance positions (see below)). Titration of EDTA-treated sBdB with Ca\(^{2+}\) ions resulted in a \(^{1}H\) \(^{15}N\)-HSQC spectrum identical to that recorded for the native protein (data not shown), whereas titration with Mg\(^{2+}\) ions resulted in an HSQC spectrum distinct from those of Ca\(^{2+}\)-bound and Ca\(^{2+}\)-depleted sBdB (Fig. 5). These data indicate that the structural effects of Ca\(^{2+}\) are specific to this metal ion.

Reduction Potential Determination for BdB—The sensitivity of tryptophan fluorescence intensity to the redox state of TDOR proteins has provided a convenient means to measure their redox potential (23, 36). BdB lacks the near-active site tryptophan, and so fluorescence methods could not be used to monitor redox state. Therefore, an alternative method was sought. Two-dimensional NMR methods have been used previously to determine the reduction potential of human thioredoxin 1 (64). NMR has the potential to provide much more detailed information about changes occurring in the protein during a redox transition. For example, the effect of redox buffers can readily be monitored through the \(^{1}H\) \(^{15}N\)-HSQC spectrum of a \(^{15}N\)-labeled protein, and the formation of adducts (e.g. mixed disulfides between the protein and glutathione) can be detected. \(^{1}H\) \(^{15}N\)-HSQC spectra of fully oxidized and reduced sBdB (Fig. 6, A and B) revealed differences in a number of resonances that, although not assigned to specific residues, enabled the redox composition of sBdB samples to be followed as a function of the potential generated by the GSSG/GSH couple (Fig. 6C). Under the conditions of concentration required for the NMR experiments, the highest potential that could be reliably generated by the GSSG/GSH couple was limited to approximately −80 mV. Normally, this would be more than sufficient to cover the entire transition for a thioredoxin-like protein. In this case, the protein was not fully oxidized at −80 mV, and so a complete redox titration data set was not obtained. Nevertheless, the data could be fitted to the Nernst equation, giving a midpoint reduction potential (\(E_m\)) of −75 ± 5 mV versus NHE at pH 7 and 25 °C with \(n\), the number of electrons involved in the reduction, at 1.85 ± 0.26. This value of \(E_m\) is similar to those reported for DsbA from \(E.\) \(coli\) and \(S.\) \(aureus\) and is consistent with an oxidative function for BdB on the outside of the cytoplasmic membrane in \(B.\) \(subtilis.\)
Oxidized BdB Contains a Destabilizing Disulfide Bond—The conformational stabilities of sBdB in both reduced and oxidized states were investigated through equilibrium unfolding (and refolding) experiments using urea, in which the intrinsic tryptophan fluorescence of sBdB was used to monitor the state of folding (Fig. 7, A and B, respectively). Urea-induced unfolding of sBdB was found to be completely reversible and was analyzed in terms of a two-state model, yielding a free energy of stabilization of $\Delta G = -11.8 \pm 1.6$ kJ mol$^{-1}$ for the oxidized protein. The greater stability of the reduced state compared with the oxidized state is a characteristic of oxidizing TDORs. For sBdB, $\Delta G_{\text{ox/red}} = 18.6 \pm 4.4$ kJ mol$^{-1}$, a value similar to that measured for EcDsbA ($14.8 \pm 4.4$ kJ mol$^{-1}$) (65). The associated $m$ values for the folding/unfolding transition were similar for both redox states (Table 2), indicating that the protein undergoes a similar degree of unfolding in each state. This is consistent with there being few structural changes between the reduced and oxidized states and that the disulfide bond is between two cysteines residues located close to one another and, therefore, has little effect on the degree of unfolding.

With information about the folding stabilities of the reduced and oxidized protein, the thermodynamic cycle (10) (illustrated in Fig. 6D) can be used to estimate the reduction potential of the folded protein. From the cycle, the difference in folding stabilities between the oxidized and reduced states ($\Delta G_{\text{ox/red}}$) is equivalent to the difference in $\Delta G$ between oxidized and reduced states in folded and unfolded states. In other words $\Delta G_{\text{ox/red}}$ is equal to the difference in reduction potentials of the protein in the folded and unfolded states. Using the expres-
TABLE 2

Free energies of stabilization of oxidized and reduced sBdbD in the presence and absence of Ca^{2+} ions at 25 °C

| sBdbD       | Midpoint of transition (D_{H2O}, N) | Cooperativity | ΔG_{stab} | ΔG_{ox/red} |
|-------------|-------------------------------------|---------------|-----------|-------------|
|             | Gdn-HCl                             | m, kJ mol^{-1} | kJ mol^{-1} | kJ mol^{-1} |
| With Ca^{2+}/reduced | 3.60                          | 8.5 ± 0.7  | -30.4 ± 2.5 | 18.6         |
| With Ca^{2+}/oxidized   | 1.37                          | 8.6 ± 1.1  | -11.8 ± 3.5 |             |
| Without Ca^{2+}/reduced | 3.28                         | 6.4 ± 0.6  | -20.9 ± 2.1 | 6.5          |
| Without Ca^{2+}/oxidized| 1.35                          | 10.7 ± 0.8 | -14.4 ± 1.1 |             |

*ΔG_{ox/red} indicates the difference between the conformational stabilities of the oxidized and reduced proteins, i.e. ΔG_{ox} - ΔG_{red}.*

**Solution K_j/K_1 = K_2/K_3**, in which K_1 and K_2 are equilibrium constants for the unfolding of reduced and oxidized sBdbD, respectively, and K_3 and K_4 are redox equilibrium constants for the unfolded and native proteins, respectively, the value of K_4 (which gives the reduction potential of the folded protein) can be obtained. To do this, the reduction potential of the unfolded protein is required to calculate K_4. It has been shown previously that reduction potentials of a range of unfolded thioxedoxin-like proteins, including *E. coli* glutaredoxin 1 (Grx1) and glutaredoxin 3 (Grx3), DsbA, and DsbC, exhibit remarkably little variation, with values of approximately -215 mV (10, 44, 66, 67). Furthermore, the reduction potential of the CXXC motifs of a range of peptides that contain the active site motifs of TDOR proteins also exhibits little variation, with values of approximately -205 mV (68). Therefore, it is reasonable to assume that the reduction potential of unfolded sBdbD is approximately ~-210 mV. Using this, a reduction potential of approximately -110 mV (versus NHE at pH 7) was calculated for folded sBdbD. This is somewhat lower than that determined by redox titration (Fig. 6) but is entirely consistent with an oxidative role of BdbD.

The removal of Ca^{2+} from sBdbD resulted in some conformational changes (see above); therefore, it was of interest to determine the relative stabilities of reduced and oxidized sBdbD in the absence of Ca^{2+}. Equilibrium unfolding experiments were carried out as above (Fig. 7, C and D, and Table 2). Reversible unfolding was observed in each case, and the data could be fitted to a two-state model. For both reduced and oxidized states, the data demonstrated that Ca^{2+} is not essential for stability; folded sBdbD is stable in both oxidation states, consistent with the structural data above. However, reduced sBdbD had a folding stability of -20.9 ± 2.0 kJ mol^{-1}, ~10 kJ mol^{-1} lower than in the presence of Ca^{2+}, and so the metal ion significantly stabilizes the reduced protein. The oxidized protein, on the other hand, in the absence of Ca^{2+} exhibited a folding stability similar to and even somewhat greater than that of Ca^{2+}-replete sBdbD (-14.4 ± 1.0 kJ mol^{-1} compared with -11.8 ± 1.6 kJ mol^{-1} for the as-isolated protein). Therefore, Ca^{2+} does not contribute to the stability of the oxidized state and may even destabilize it relative to the Ca^{2+}-free form. It is also of note that the m values for Ca^{2+}-depleted sBdbD are significantly different for the reduced and oxidized states, implying that the extent of unfolding is different, consistent with a difference in structure between the oxidized and reduced states in their native and/or denatured forms.

The data indicate clearly that the loss of Ca^{2+} leads to a drop in the value of ΔG_{ox/red} and therefore a decrease in the difference between reduction potentials in folded and unfolded states. From the cycle in Fig. 7D, the reduction potential of sBdbD in the absence of Ca^{2+} was calculated to be ~60 mV lower than when Ca^{2+} is present, suggesting that a major role of Ca^{2+} is to increase the oxidizing power of BdbD.

*Ca^{2+} Site of BdbD Serves to Increase the Active Site Reduction Potential*—To determine directly the effect of the presence of Ca^{2+} on the redox properties of sBdbD, the mid-point reduction potential of the Ca^{2+}-free protein was determined by NMR methods, as described above for the as-isolated Ca^{2+}-containing protein (Fig. 6C). At the highest potential that could be generated by the GSSG/GSH couple (-80 mV), the extent of reduction was much more significant than in the Ca^{2+}-containing protein, and fitting of the data to the Nernst equation, giving a midpoint reduction potential (E_{m}) of -95 ± 5 mV versus NHE at pH 7 and 25 °C, with n, the number of electrons involved in the reduction, at 1.98 ± 0.2. This value of E_{m} is
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significantly lower than that determined for the Ca$^{2+}$-containing protein, demonstrating that the Ca$^{2+}$ site, although 14 Å away from the nearest active site cysteine, exerts an important effect on the redox properties of the protein, such that in its presence the potential is $\sim 20$ mV more positive. This difference, although less than that predicted from unfolding studies, is qualitatively in good agreement.

$pK_a$ Values of BdbD Active Site Cysteines—To define the range over which the $pK_a$ values of the sBdbD active site cysteines can be measured, the pH stability of folded sBdbD was investigated using the intrinsic fluorescence of tryptophan. A plot of emission wavelength maximum versus pH (Fig. 8A) showed that pH-induced unfolding of sBdbD led to a sizeable shift in emission wavelength from around 353 nm at pH 7 to much longer wavelengths at extremes of pH. The apparent stability range of BdbD lies between pH 4.5 and pH 10 and is similar to both ResA and StoA from B. subtilis (23, 36). Within the stable range itself, there was little evidence for titration of other residues affecting the fluorescence of tryptophan residues. The data indicate that $pK_a$ values lower than 4.5 cannot be measured. This is particularly important for sBdbD because, as a DsbA-like protein, it is likely to have a highly nucleophilic N-terminal cysteine in the CXXC motif (11, 44).

The acid-base properties of the active site cysteines of sBdbD were investigated by measuring rates of reaction with the fluorescent probe badan, as described under “Experimental Procedures” (Fig. 8B). During the badan modification experiment, a single fluorescence peak with a maximum emission at 510 nm was observed for badan-treated sBdbD, corresponding to covalent badan-adduct(s) of protein cysteine residue(s). A plot of the pseudo-first order rate constant versus pH revealed a significant rate constant at low pH and an apparent protonation/deprotonation event at pH 8.6 (Fig. 8C). The rate constant at low pH is roughly constant ($\sim 0.045$ min$^{-1}$), whereas that measured after the transition is estimated to be $\sim 0.102$ min$^{-1}$ on the basis of the fitted Henderson-Hasselbalch curve. sBdbD contains only two cysteines (Cys$^{69}$ and Cys$^{72}$) and, although only a single titration event is detected, the high rate of badan modification at low pH strongly suggests that a cysteine residue must be reacting with badan at acidic pH values. We therefore conclude that the apparent $pK_a$ event is likely to be due to titration of Cys$^{72}$, whereas the high rate at low pH is because of the thiolate form of Cys$^{69}$, which is likely to have a $pK_a$ value below the minimum pH at which sBdbD is stable. This is consistent with $pK_a$ values reported for other DsbA-like proteins where the N-terminal cysteine usually has a very low $pK_a$ value (11, 16, 44).

The pH stability studies reported here highlight that the determination of accurate cysteine $pK_a$ values requires knowledge of the pH-induced unfolding, because the $pK_a$ values of cysteines in the unfolded protein are likely to be significantly higher than in the folded state. This raises the question of how the effects of protein unfolding at low pH have been taken into account in previous determinations of cysteine $pK_a$ values in EcDsbA and other DsbA-like proteins. We note that for SaDsbA, a $pK_a$ of 3.37 was determined (16), but the pH stability of the protein was not reported, although Charbonnier et al. (69) concluded that the $pK_a$ value of wild-type EcDsbA could not be accurately determined because of protein unfolding at low pH. Nelson and Creighton (11) directly addressed this issue by monitoring thiolate formation in EcDsbA using the absorbance change at 240 nm to report on thiol/thiolate status, while simultaneously measuring the 288 nm signal as indicative of folding/unfolding transitions. From these data, a $pK_a$ of $\sim 3.5$ for the N-terminal cysteine was determined, but it was also shown that the protein unfolded below pH 3. Thus, deprotona-
tion and unfolding events overlap, raising the question of whether the pK_a values reported are really because of protonation/deprotonation of the active site cysteine in the native state or, in fact, measure protonation/deprotonation because of the relaxation of structural factors that depress the cysteine pK_a value in the folded state. Thus, although it is clear that the N-terminal cysteine pK_a values are considerably lower in DsbA-like proteins (including BdbD) than in reducing thioredoxin-like proteins (which have pK_a values of >6.5), or for free cysteine in solution (pK_a value ~8.5), it is not clear that the pK_a value for any of these proteins has been accurately determined.

Probing Redox Interactions of BdbD—To investigate the ability of sBdbD to oxidize a substrate, a 15-residue peptide containing the active site sequence CXXCH of B. subtilis apo-cytochrome c_50 (CccA) was used. A significant body of evidence indicates that BdbD oxidizes apo-cytochrome c substrates upon their export to the outside of the cytoplasmic membrane before they are subsequently re-reduced by the reducing TDOR ResA immediately prior to covalent heme attachment (21, 38, 52). sBdbD was oxidized overnight by the addition of 10 mm GSSG, and a 2:1 excess of reduced peptide was added. The resulting ^1H ^15N-HSQC spectrum showed no evidence for the presence of a stable interaction between sBdbD and the peptide, consistent with the transient nature of the interaction of DsbA-like proteins with substrates, but it did reveal that sBdbD was present entirely in the reduced form (data not shown). An identical experiment with an equivalent peptide containing an SXXSH active site resulted in the observation of sBdbD only in the oxidized state. These experiments clearly demonstrate that oxidized sBdbD is able to transfer its disulfide bond to the apo-cytochrome c_50 active site mimic peptide.

The data presented here (and previously) show that BdbD is an oxidizing extra-cytoplasmic TDOR. B. subtilis contains a number of other extra-cytoplasmic TDORs, two of which, ResA and StoA, have been shown to be reducing TDORs involved in cytochrome c maturation and endospore biogenesis, respectively (21–23). The existence of both oxidizing and reducing TDORs on the outside of the membrane raises questions about how unproductive redox cycling of these proteins and their substrates is avoided. We therefore sought to determine whether or not BdbD can directly oxidize reduced ResA in vitro. The fluorescence emission intensity of sResA at 353 nm is highly sensitive to the redox state of the protein, becoming significantly quenched upon oxidation (21). As noted above, the fluorescence intensity of sBdbD, on the other hand, is insensitive to the redox state of the protein. Therefore, fluorescence spectroscopy was used to investigate whether sBdbD and sResA can undergo a redox reaction with one another. Oxidized sBdbD was mixed with an equimolar concentration of reduced sResA and fluorescence spectra measured over a period of 1 h (Fig. 9). Initially, a small loss of intensity occurred, but overall significant changes in intensity were not observed. A similar slow loss of intensity was observed upon addition of a redox-inactive SXXSH cytochrome c_50 active site mimic peptide, indicating that small intensity changes are observed even when electron transfer cannot take place (data not shown). Intensity changes are shown more clearly in the plot of fluorescence intensity at 345 nm as a function of time (Fig. 9, inset). As a control, sBdbD was also mixed in 1:1 ratio with oxidized sResA, verifying that oxidation of sResA results in a large decrease in intensity. The data indicate that, despite a significant thermodynamic driving force, the proteins do not undergo a redox reaction, at least in vitro.

Concluding Remarks—The requirement for disulfide bonds in many proteins located on the outside of the cytoplasmic membrane has led to the evolution of a range of thiol-disulfide exchange systems in many bacteria (15). Although most of these involve proteins closely related to the DsbA/DsbB system of E. coli, there are significant variations. M. tuberculosis, for example, utilizes a protein, DsbE, more closely related to thioredoxin than DsbA to generate extra-cytoplasmic disulfide bonds, and S. aureus does not contain a DsbB homologue and therefore must re-oxidize its DsbA protein by another pathway. Members of the genus Bacillus, with few exceptions, contain BdbD/BdbC, which are homologues of DsbA/DsbB. Here our studies of BdbD from B. subtilis have revealed further evidence of variation, in that the protein possesses an intrinsic metal site not present in other DsbA-like proteins. Upon isolation, the site is fully occupied by a Ca^{2+} ion, which commonly plays a structural role in proteins. In this case, the function of the site appears to be more subtle as removal of the metal does not result in unfolding, or indeed any major structural changes. It does have an effect on conformational stability, however, and, intriguingly, this is specific to the reduced state of the protein, such that presence of Ca^{2+} at the site significantly affects the redox properties of the protein, despite being some 14 Å away from the enzyme active site. Electrostatics may be a significant contributor to the observed effect. In simple terms, binding of a positively charged calcium ion would be expected to stabilize the reduced form of the active site, in which Cys^{19} is negatively charged at physiological pH. On the other hand, no stabilization would be expected to result from Ca^{2+} binding to the neutral oxidized (disulfide-bonded) state. This provides a qualitative explanation for the selective stabilization by Ca^{2+} of the reduced state of BdbD, which would be predicted to result in an increase in reduction potential, as observed.
Some TDORs, such as human glutaredoxin 2, are iron-sulfur cluster-containing proteins (70, 71). However, in such cases, the cluster cofactor is coordinated by at least one of the active site cysteines, and therefore TDOR activity and cluster coordination are mutually exclusive properties. To the best of our knowledge, B. subtilis BdbD is the first example of a TDOR protein that contains a metal cofactor remote from the CXXC active site, and it represents an important example in which a non-redox active metal affects the redox properties of a protein. This is somewhat reminiscent of peroxidase enzymes in which Ca²⁺ has an important effect on the properties of the active site heme (72). For BdbD, the effect of Ca²⁺ is to increase the reduction potential, and the magnitude of the effect is of the same order as that observed for XX dipeptide substitutions at a TDOR CXXC active site (38). This leads us to propose that the role of the metal site is to provide a boost to the oxidizing power of BdbD. A further possibility that remains to be explored is that modulation of the active site reduction potential is important for the re-oxidation of the BdbD active site by its integral membrane partner BdbC.

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