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Subdivision of the bacterioferritin comigratory protein (BCP) family of bacterial peroxiredoxins based on catalytic activity^{\dagger}

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Running title: Characterization of 1-Cys peroxiredoxin catalytic activity

Abbreviations used: BCP, bacterioferritin comigratory protein; *Ec*BCP, BCP of *Escherichia coli*; *Bc*BCP, BCP of *Burkholderia cenocepacia*; *Xc*BCP, BCP of *Xanthomonas campestris*; Prx, peroxiredoxin; ROS, reactive oxygen species; CF, cystic fibrosis; Bcc, *Burkholderia cepacia* complex; WT, wildtype; Trx, thioredoxin; TrxR, thioredoxin reductase; GSH, reduced glutathione; Grx, Glutaredoxin; IAM, iodoacetamide; NEM, N-ethyl maleimide; TCEP, Tris(2-carboxyethyl)phosphine hydrochloride; FT-ICR-MS, Fourier transform ion cyclotron resonance mass spectrometry; MS, mass spectrometry; ECD, electron capture dissociation; CID, collision-induced dissociation.

ABSTRACT

Peroxiredoxins are ubiquitous proteins that catalyze the reduction of hydroperoxides, thus conferring resistance to oxidative stress. Using high-resolution mass spectrometry, we recently re-classified one such peroxiredoxin, bacterioferritin comigratory protein (BCP) of Escherichia coli, as an atypical 2-Cys peroxiredoxin that functions through the formation of an intramolecular disulfide bond between the active and resolving cysteine. An engineered E. *coli* BCP, which lacked the resolving cysteine, retained enzyme activity through a novel catalytic pathway. Unlike the active cysteine, the resolving cysteine of BCP peroxiredoxins is not conserved across all members of the family. To clarify the catalytic mechanism of native BCP enzymes that lack the resolving cysteine, we have investigated the BCP homologue of Burkholderia cenocepacia. We demonstrate that the B. cenocepacia BCP (BcBCP) homologue functions through a 1-Cys catalytic pathway. During catalysis, BcBCP can utilize thioredoxin as a reductant for the sulfenic acid intermediate. However, significantly higher peroxidase activity is observed utilizing glutathione as a resolving cysteine and glutaredoxin as a redox partner. Introduction of a resolving cysteine into BcBCP changes the activity from a 1-Cys pathway to an atypical 2-Cys pathway, analogous to the E. coli enzyme. In contrast to the native B. cenocepacia enzyme, thioredoxin is the preferred redox partner for this atypical 2-Cys variant. BCP-deficient B. cenocepacia exhibit a growth-phase dependent hypersensitivity to oxidative killing. Based on sequence alignments, we believe that BcBCP described herein is representative of the major class of bacterial BCP peroxiredoxins. To our knowledge, this is the first detailed characterization of their catalytic activity. These studies support the subdivision of the BCP family of peroxiredoxins into two classes based on their catalytic activity.

Reactive oxygen species (ROS) arising from the incomplete reduction of oxygen are an inevitable consequence of aerobic metabolism, and result in wide-ranging oxidative damage to nucleic acids, proteins and lipids. Organisms occupying aerobic environments have evolved numerous reductive defense mechanisms against ROS. Peroxiredoxins (Prxs), which undertake the thiol-dependent reduction of peroxide substrates, constitute one such defense mechanism. The catalytic activity of Prxs can be divided into 1-Cys and 2-Cys pathways based on the number of cysteines required for catalysis. The 2-Cys Prxs are further divided into typical and atypical 2-Cys peroxidases; typical 2-Cys Prxs being obligate dimers, whilst atypical 2-Cys Prxs are monomeric. During the reduction of hydroperoxides, all Prxs share a common first step, which involves an active (peroxidatic) cysteine thiolate attacking the peroxide substrate resulting in the formation of a sulfenic acid (Cys-SOH) intermediate on the active cysteine. Thereafter, different pathways may be employed in the resolution of the sulfenic acid intermediate, and it is variation in this process which leads to the delineation of the Prxs into 1-Cys and 2-Cys peroxidases (1). 2-Cys Prxs contain a second cysteine, termed the resolving cysteine, which attacks the sulfenic acid intermediate of the active cysteine, forming a disulfide bond that is subsequently reduced by a cell-specific oxidoreductase. The disulfide bond between the active and resolving cysteine can be either intramolecular (monomeric atypical 2-Cys pathway) or intermolecular (dimeric typical 2-Cys). In contrast, 1-Cys Prxs do not harbour a resolving cysteine. In this case, the sulfenic acid intermediate is thought to be resolved by thiol-containing reductants such as glutathione or thioredoxin.

The bacterioferritin comigratory protein (BCP) family of Prxs was first characterized in *Escherichia coli* (2). BCP is a monomeric protein with thiol-dependent peroxidase activity, which until recently was assumed to function via a 1-Cys mechanism, based on observations made of the *E. coli* enzyme. However, by applying high-resolution mass spectrometry, we recently reclassified *E. coli* BCP (*Ec*BCP) as an atypical 2-Cys enzyme, functioning via the

formation of an intramolecular disulfide bond between an active and resolving cysteine (3). Sequence alignments of prokaryotic BCP homologues revealed that the majority of homologues lack the resolving cysteine, and as such would be expected to function via a 1-Cys catalytic process. However, an engineered *Ec*BCP, in which the resolving cysteine was replaced with a serine (C50S), retained peroxidatic activity through a novel catalytic mechanism distinct from the anticipated 1-Cys pathway (3).

BCP homologues lacking a resolving cysteine have been assumed to function via the 1-Cys Prx pathway. However, the mechanistic details of such enzymes have never been fully characterized. In light of the novel pathway adopted by the engineered *Ec*BCP C50S enzyme (*3*), we sought to define the catalytic mechanism employed by native BCP homologues that lack a resolving cysteine. We herein characterize one such member of the BCP family; the BCP Prx of *Burkholderia cenocepacia*, an aerobic Gram-negative organism that belongs to a group of closely-related bacterial species referred to as the *B. cepacia* complex (Bcc) (*4*). *B. cenocepacia* is a significant pathogen of the cystic fibrosis (CF) lung, an environment that is considered highly oxidative as a consequence of the profound influx of neutrophils and the resulting release of ROS. In addition, *B. cenocepacia* is capable of surviving the oxidative burst within activated macrophages (*5*, *6*). Consequently, anti-oxidant activities are thought to play a role in the ability of this organism to establish chronic infection of the CF airways.

We previously reported that the *bcp* homologue of *B. cenocepacia* is coexpressed with the *arn* locus (7). The *arn* locus encodes the biosynthetic enzymes required for the modification of lipopolysaccharide with aminoarabinose, a positively-charged sugar that lowers the binding affinity of cationic antimicrobial peptides, thus conferring resistance (8). Such antimicrobial peptides are the dominant mechanism of non-oxidative killing by phagocytic cells. Thus, the genetic linkage of *bcp* with the *arn* locus within *Burkholderia* species combines putative resistance mechanisms against both oxidative and non-oxidative killing by phagocytes. In the

present study, we demonstrate that the BCP of *B. cenocepacia* displays the anticipated thioldependent peroxidase activity, which protects against oxidative stress in a growth-phase dependent manner. The native enzyme, which lacks a resolving cysteine, functions via a 1-Cys catalytic pathway, whilst introduction of a resolving cysteine switches the catalytic pathway to an atypical 2-Cys mechanism that is functionally indistinguishable from that of the *E. coli* enzyme. We conclude that the BCP family of bacterial Prxs can be subdivided on the basis of catalytic activity, and that the 1-Cys peroxidatic mechanism, typified by the *Burkholderia* enzyme, represents the dominant class.

EXPERIMENTAL PROCEDURES

Bacterial strains, macrophages and culture conditions. B. cenocepacia strain K56-2 was used in this study. Based on multilocus sequence typing (MLST), *B. cenocepacia* K56-2 belongs to the same clonal complex as the genome sequence strain *B. cenocepacia* J2315, a representative of the highly transmissible ET12 epidemic lineage. *B. cenocepacia* XOA14 was created by insertional inactivation of the *B. cenocepacia* K56-2 *bcp* homologue (corresponding to BCAL1936 of *B. cenocepacia* J2315) using the pGP Ω Tp suicide plasmid (*9*). *E. coli* strains DH5 α and BL21(DE3) were used for cloning and protein expression respectively. All bacterial strains were routinely grown at 37 °C in Luria Bertani (LB) medium. *Burkholderia* cultures were supplemented as required with 100 µg/ml of trimethoprim (Tp), 100 µg/ml tetracycline (Tet) and/or 50 µg/ml of gentamicin (Gm). *E. coli* cultures were supplemented with Tp (50 µg/ml), Tet (20 µg/ml) or kanamycin (30 µg/ml). Bacterial strains are listed in Table 1.

The murine macrophage cell line RAW264.7 was obtained from the American Type Culture Collection. ANA1 macrophages were a gift of Danuta Radzioch, McGill University, Quebec,

Canada. Cells were maintained in DMEM with 10% fetal bovine serum and grown at 37°C with 5% carbon dioxide.

Cloning and site-directed mutagenesis of BCP. The *bcp* gene of *B. cenocepacia* K56-2 was PCR amplified (PfuUltra II Fusion HS DNA Polymerase, Stratagene), incorporating *Nde*I and *BamH*I restriction sites into the forward and reverse primers respectively, enabling ligation into *NdeI/BamH*I-digested pET-28a vector (Novagen). This resulting vector, pARB1, encodes wildtype (WT) *Bc*BCP with an N-terminal 6-His tag. His-tagged WT *Ec*BCP was expressed from a comparable vector (pARB5) following amplification of the open-reading frame from *E. coli* K12. Using pARB1 vector as template and the QuikChange II Site-Directed Mutagenesis Kit (Stratagene), three mutant open-reading frames of *Bc*BCP were created: C44S (pARB2), C98S (pARB3) and L49C (pARB4). The resulting plasmids (pARB1 – pARB5) were used to transform *E. coli* strain BL21 (DE3). Primer sequences used for cloning and site-directed mutagenesis are available upon request.

Expression and purification of BCP enzymes. E. coli BL21 (DE3) cells transformed with the appropriate BCP-encoding plasmid were cultured overnight at 37 °C in 50 ml LB medium supplemented with kanamycin (30 μ g/ml). Overnight cultures were used to inoculate 1 litre of fresh medium, which was cultured to an optical density at 600 nm of approximately 0.6. Isopropyl-1-thio- β -D-galactopyranoside (IPTG) was added to a final concentration of 0.25 mM and cultures incubated for 3 h. Cells were harvested by centrifugation and resuspended in 20 ml Tris (50 mM), NaCl (500 mM), imidazole (20 mM), pH 7.5. Cells were disrupted by sonication and lysates centrifuged. Clarified supernatant was filtered (0.45 μ M) prior to loading onto a HisTrap FF 5 ml column (GE Healthcare). His-tagged BCP protein was eluted from the nickel column with increasing imidazole concentration and de-salted using a PD Miditrap G-25 column (GE Healthcare), eluting in Hepes-NaOH (pH 7.0).

Determination of thioredoxin-linked peroxidase activity of BCP. Thioredoxin-linked peroxidase activity was assessed essentially as described previously (2). In brief, reactions were performed at room temperature in 500 μ L volumes, comprising 0.36 μ M thioredoxin reductase (TrxR; Sigma-Aldrich), 1.7 μ M thioredoxin (Trx; Sigma Aldrich), 0.1 mM hydrogen peroxide and 0.2 mM NADPH (Roche) in 50 mM Hepes-NaOH (pH 7). BCP concentrations were variable, ranging from 0 to 16 μ M. Peroxide was added to the reaction mixture last, and the absorbance at 340 nm was measured continuously for 5 min. The assay was also performed with 1 mM GSH in the reaction mixture.

Determination of glutaredoxin-linked peroxidase activity of BCP. Reactions were performed at room temperature in 500 μL volumes, comprising 1 unit glutathione reductase (Calbiochem), 12 μM glutaredoxin (Grx, Calbiochem), 1 mM reduced glutathione (GSH, Sigma-Aldrich), 0.1 mM hydrogen peroxide and 0.15 mM NADPH (Roche) in 50 mM Hepes-NaOH (pH 7). BCP concentrations were variable, ranging from 0 to 16 μM. Peroxide was added to the reaction mixture last, and the absorbance at 340 nm was measured continuously for 5 min.

 H_2O_2 oxidation. Typically, wildtype and mutant BCP proteins were at a concentration of 50 μ M in ammonium acetate (100 mM, pH 7.2), GSH was added to a final concentration of 1mM before reaction with H_2O_2 . Typically, oxidation was allowed to proceed for a few seconds before the addition of 4-fold H_2O :MeOH:HCOOH (50:48:2) (v/v) quenched the reaction.

Preparation of reduced thioredoxin and glutaredoxin. Thioredoxin and glutaredoxin were resuspended in 100 mM ammonium acetate (pH 7.2) before reduction with 2 mM TCEP. Excess TCEP was removed by buffer exchange using a PD-Miditrap desalting column (GE Healthcare) and the sample was eluted in 50 mM ammonium acetate, pH 5.5. Disulfide bond

reduction was verified by FT-ICR-MS and the reduced protein was stored at 4 °C. Protein concentration was measured using the BCA assay (Pierce, Rockford, IL).

Reaction of oxidized BCP species with reduced thioredoxin or glutaredoxin. Oxidized wildtype and L49C BCP species were produced by treatment of 50 μ M protein with 100 μ M hydrogen peroxide, in the presence of 1 mM GSH. The resulting oxidized species were purified by PD-Miditrap desalting column (GE Healthcare) and eluted with 200mM ammonium acetate (pH 7.2). The samples (10 μ M) were then treated with 10 μ M reduced thioredoxin or glutaredoxin and incubated at room temperature for 5 min. The reaction was quenched with 2-fold H₂O:MeOH:HCOOH (v/v/v 50:48:2), and analyzed by FT-ICR-MS.

FT-ICR mass spectrometry. Protein samples were buffer exchanged into 100 mM ammonium acetate, pH 7.4 before addition of hydrogen peroxide and mass spectrometry analysis. Reactions were quenched with 2-fold H₂O:MeOH:HCOOH (v/v/v 50:48:2), and analyzed by FT-ICR-MS. Mass spectrometry data was acquired on an Apex Ultra Qh-FT-ICR mass spectrometer equipped with a 12 Tesla superconducting magnet and an electrospray ion source (Bruker Daltonics, Billerica, MA). Nano-ESI was performed using a TriVersa Nanomate (Advion BioSciences, Ithaca, NY) running in infusion mode. Broadband data was typically acquired between m/z 600 and 4000 for 0.5 s and each spectrum was the sum of 32 scans. The mass spectra were externally calibrated using ES tuning mix (Agilent) and analyzed using DataAnalysis software (Bruker Daltonics). Isotope distributions of specific charge states were predicted using IsotopePattern software (Bruker Daltonics) from theoretical empirical formulae. These were overlaid upon the recorded experimental data as scatter plots, with the theoretical apex of each isotope peak designated by a circle.

Top-Down FT-ICR-tandem mass spectrometry. Top-down fragmentation was performed on the 12T Qh-FT-ICR. First, a specific ion species was isolated with the instrument's mass

resolving quadrupole and MS/MS was performed using collision-induced dissociation (CID) or electron capture dissociation (ECD). For CID, the collision voltage was typically set between 20-35V. For ECD, 1.8 amperes was applied to the dispenser cathode filament (Heatwave Technologies), 20V to the lens and a pulse of 4-9 ms was employed. Fragmentation data was the sum of 250-750 scans and data analyses were performed using DataAnalysis (Bruker Daltonics). The SNAP algorithm was used for automated peak picking and the resulting top-down fragment mass lists were searched against the primary sequence of BCP using Prosight-PTM software (*10, 11*). Mass error tolerances were set for all Prosight searches at 10 ppm.

Complementation of BCP function. The N-terminal 6-His tagged WT *bcp* ORF was PCR amplified from pARB1, incorporating *EcoR*I and *Xba*I restriction sites into the forward and reverse primers respectively. The ORF was subsequently ligated into *EcoR*I- and *Xba*I-digested pDA17 vector (Tet^R), generating pARB10. pARB10 was introduced into XOA14 by triparental mating, resulting in strain ARB201. Expression of 6-His-tagged *Bc*BCP was confirmed by Western analysis of whole-cell lysates using anti-His antibody (GE Healthcare) and HRP-linked anti-mouse IgG (Cell Signaling Technology).

Expression analysis of BCP. B. cenocepacia K56-2 was grown to logarithmic phase in LB prior to isolation of RNA (RNeasy Protect Bacteria Mini, Qiagen), DNAse treatment (RNase-free DNase set, Qiagen) and subsequent cDNA synthesis (SuperScript III Reverse Transcriptase, Invitrogen). To assess expression of *bcp* and representative genes from the upstream *arn* locus (*arnT* and *arnB*), quantitative real-time PCR (QRT-PCR) was performed using the Platinum SYBR Green qPCR SuperMix-UDG (Invitrogen) in a 25 μ L reaction volume containing 10 μ L of 1 in 100 diluted cDNA, 400 nM primers and 4 % (v/v) DMSO. All primer pairs exhibited PCR efficiencies between 90 and 104 %, with R² values greater than 0.99 over at least a 10³ range of template concentrations. Primer sequences are available

upon request. Thermal cycling was performed on a Stratagene Mx3000P thermal cycler and data analyzed using the Comparative Quantitation algorithm. Dissociation curve analysis and electrophoresis of representative samples confirmed identity of PCR products.

Northern blot analysis was performed according to standard procedures, with ³²P-labelled probes prepared using the RadPrime DNA Labeling System (Invitrogen).

Assessment of sensitivity to hydrogen peroxide. Strains were grown in LB without supplements. At appropriate time points throughout the growth phase, culture aliquots were removed to fresh culture vessels and challenged with either hydrogen peroxide (to a final concentration of 1 mM) or an equivalent volume of PBS (control). After 30 min incubation at 37 °C, appropriate serial dilutions were plated, allowing viable counts and percentage survival to be determined.

Macrophage infection assays. Macrophage infection assays were performed to assess intracellular survival of *B. cenocepacia*. To quantify intracellular survival, bacteria were added to macrophages at a multiplicity of infection (MOI) of 50 bacteria per macrophage, and internalization of bacteria was allowed to proceed for 2 h. Following internalization, cell culture medium was supplemented with ceftazidime (1 mg/ml) and kanamycin (500 µg/ml) to kill remaining extracellular bacteria. The number of surviving intracellular bacteria at 2 and 4 h post-internalization was determined by triplicate plating of serially diluted macrophage cell lysate.

Localization of *Burkholderia* within macrophages was assessed using tetramethyl-rhodaminedextran (TMR-dextran) to label lysosomes, and LysoTracker Red to label acidic compartments. For lysosome labeling prior to bacterial infection, ANA-1 macrophages were incubated with 250 µg/mL TMR-dextran (Molecular Probes) for 2 h, followed by a 1 h chase in dye-free medium to ensure full delivery of TMR-dextran to lysosomes. Macrophages were

infected at MOI 50 and incubated for a further 2 or 4 h, prior to visualization using differential interference contrast (DIC) and fluorescence microscopy to quantify the percentage of *Burkholderia cepacia*-containing vacuoles (BcCVs) colocalizing with TMR-dextran.

For labeling of acidic compartments with LysoTracker Red, RAW264.7 macrophages were infected with bacteria at an MOI of 50. At 4 h postinfection (and immediately prior to visualization by DIC and fluorescence microscopy), acidic compartments within infected macrophages were labeled by incubation with LysoTracker Red DND-99 (Molecular Probes) at a final concentration of 0.25 μ M for 1 min.

RESULTS

Burkholderia BCP displays thiol-dependent peroxidase activity. Based on homology with known BCP peroxiredoxins, it was predicted that *Bc*BCP would exhibit thiol-dependent peroxidase activity. N-terminal 6His-tagged recombinant WT *Bc*BCP was expressed and purified to homogeneity using a nickel affinity column. Gel filtration of the protein showed it to behave as a monomer with a mass of approximately 19 kDa, which was subsequently confirmed by ESI mass spectrometry (average mass 18992 Da).

The peroxidase activity of *Bc*BCP linked to oxidation of NADPH via the *E. coli* thioredoxin/thioredoxin reductase (Trx/TrxR) system was monitored spectrophotometrically by measuring the decrease in A_{340} . In the absence of *Bc*BCP, a background level of NADPH oxidation occurred at a steady rate of 0.10 μ Mmin⁻¹. In the presence of *Bc*BCP, the rate of NADPH oxidation increased in a BCP concentration-dependent manner and was recorded as $1.04 \pm 0.08 \ 0.1 \ \mu$ Mmin⁻¹ per μ M of BCP (μ Mmin⁻¹ μ M⁻¹). Comparable BCP enzyme activity was observed in the presence of all peroxide substrates tested (hydrogen peroxide, cumene hydroperoxide and tert-butyl hydroperoxide). Pre-treatment of *Bc*BCP with iodoacetamide

(IAM; 10 mM final concentration), which covalently modifies free thiol groups, abolished enzyme activity (0.10 μ Mmin⁻¹ μ M⁻¹), confirming that the activity of *Bc*BCP is thiol-dependent.

Cys-44 is the active site of Burkholderia BCP. Site-directed mutants of *Bc*BCP were generated, in which each of the cysteine residues was individually replaced with a serine (C44S and C98S *Bc*BCP). The enzymatic activity of these mutants was studied as before. Activity was abolished in the C44S mutant (Figure 1). In contrast, C98S *Bc*BCP retained full enzyme activity. Together, these findings are consistent with the Cys-44 residue being the active site of the thiol-dependent *Bc*BCP, whilst the Cys-98 residue is a bystander, not involved in the catalytic process. This is consistent with previous observations made of the equivalent cysteine residues (Cys-45 and Cys-99) of *Ec*BCP (2).

The Cys-44 residue of *Bc*BCP was confirmed as the active (peroxidatic) cysteine by mass spectrometry (Figure 2). Treatment of WT *Bc*BCP with 10 molar equivalents of H₂O₂ resulted in an increased mass of 32 Da, consistent with one free thiol group (-SH) being oxidized to the sulfinic acid derivative (-SOOH, Figure 2B). Following treatment with H₂O₂, the mass of C98S *Bc*BCP similarly increased whilst the mass of C44S *Bc*BCP remained unchanged, suggesting that it is the Cys-44 residue that is oxidized during the catalytic process (data not shown). Top-down fragmentation of the oxidized wildtype enzyme, using ECD, subsequently mapped the sulfinic acid modification to Cys-44, confirming that residue as the peroxidatic cysteine (Figure 2C and Supporting Information Figure S1).

Burkholderia BCP is a 1-Cys peroxiredoxin. E. coli BCP functions via an atypical 2-Cys pathway, employing a peroxidatic cysteine at position 45 and a resolving cysteine at position 50 (*3*). However, *Bc*BCP lacks the equivalent resolving cysteine (Figure 3). We previously demonstrated that in the absence of the resolving cysteine, *Ec*BCP (C50S) adopts a novel

mechanistic pathway, progressing via a dimeric intermediate, which contains an intermolecular disulfide bridge between the Cys-45 residues of two monomers (*3*). The native *Bc*BCP was interrogated by high-resolution mass spectrometry to investigate if it too adopts this novel reaction pathway.

As stated above, in excess H_2O_2 and in the absence of any redox partner, *Bc*BCP forms a stable sulfinic acid (Cys-SOOH), suggesting overoxidation of a sulfenic acid intermediate. However, even at stoichiometric quantities of H_2O_2 , no evidence for dimerization was observed by mass spectrometry. This result led us to speculate whether *Bc*BCP could utilize a small molecule thiol as a resolving partner in a manner previously described for 1-Cys peroxiredoxins (*12-16*).

A mechanistic study of the 1-Cys D-Prx has highlighted the interaction of glutathione (GSH) with the enzyme during turnover (*14*). Therefore, experiments with *Bc*BCP were repeated in the presence of 1mM GSH. Mass spectrometry shows that in the presence of GSH, but absence of H₂O₂, *Bc*BCP exists as a monomer with two reduced cysteine thiols (Figure 4A). After addition of 1 molar equivalent of H₂O₂, we observed the appearance of a species with Δ mass +305 Da, consistent with the formation of a *Bc*BCP-GSH heterodimer (Figure 4B). Presumably, GSH modification is via a mixed disulfide bond with one of the two cysteine residues within BCP. To map the site of glutathionylation, top-down fragmentation was employed. The *Bc*BCP-GSH heterodimer was treated with the alkylating reagent N-ethyl maleimide (NEM) to covalently modify any free cysteine residue. The NEM-labeled GSH-modified species was then reduced by treatment with 20 mM DTT, resulting in a Δ mass of - 305 Da, which we attribute to loss of GSH. This mass change confirmed that the glutathionylation modification was indeed via a disulfide bond. This final species was subject to CID fragmentation using FT-ICR MS. Analysis of the resulting fragments allowed the

NEM modification to be mapped to Cys-98, indicating that the glutathionylation occurred on the active cysteine, Cys-44 (Figure 4C and Supporting Information Figure S2).

Glutaredoxin can act as the reductive partner to BcBCP in a glutathione-dependent catalytic pathway. These mass spectrometric results suggest that GSH acts as a small molecule resolving cysteine, reducing the transient sulfenic acid intermediate on Cys-44 releasing H₂O and producing a mixed disulfide between Cys-44 and GSH. If this hypothesis is correct, the resulting BcBCP-GSH intermediate requires a reductive partner. This partner must reduce the intermolecular disulfide bond, regenerating a reduced peroxidatic cysteine on BcBCP, which would allow further catalytic cycles. Using high-resolution mass spectrometry, we monitored the ability of thioredoxin to reduce the BcBCP-GSH intermediate. BcBCP-GSH was prepared by treating reduced BcBCP with 2 equivalents of H₂O₂ in the presence of 1mM GSH (as shown in Figure 4B). Excess GSH and peroxide were removed using size exclusion chromatography before this species was incubated with an equimolar ratio of reduced thioredoxin. The mixture was equilibrated at room temperature for 2 min before analysis by FT-ICR MS (Supporting Information Figure S3). Analysis of the isotopic distributions within the resulting mass spectrum revealed no mass shift for either species, suggesting that thioredoxin is not an efficient reducing partner for BcBCP-GSH. This observation seemed to contradict results from the Trx/TrxR enzyme assay that demonstrated enzyme turnover using thioredoxin as the reducing partner. However, repeating the enzyme assay in the presence of 1mM GSH resulted in a dramatic loss of enzyme activity, recorded at $0.19 \pm 0.05 \,\mu\text{Mmin}^{-1}$ ${}^{1}\mu M^{-1}$ (Figure 1). This result suggests that glutathionylation of the peroxidatic cysteine inhibits the thioredoxin-dependent reductive step of catalysis. Together, these observations imply that thioredoxin can act as a reductant in BcBCP catalysis by directly reducing the transient Cys44-S-OH intermediate to Cys44-SH. However, in the presence of GSH, the

transient Cys44-S-OH is quickly glutathionylated. The resulting *Bc*BCP-GSH mixed disulfide is not a substrate for reduction by thioredoxin and *Bc*BCP turnover is inhibited.

As stated above, glutathione-dependent catalysis has been reported previously for peroxiredoxin enzymes - the poplar phloem 1-Cys D-Prx has been extensively studied and displays glutathione dependence (*15, 16*). Furthermore, NMR and MS studies of this enzyme have revealed that the poplar Prx can be glutathionylated on its peroxidatic cysteine residue and the enzyme accepts glutaredoxin (Grx) as a proton donor (*14*). Using high resolution MS, we monitored the ability of glutaredoxin to reduce the *Bc*BCP-GSH intermediate. Equimolar ratios of reduced Grx and *Bc*BCP-GSH were mixed and equilibrated at room temperature for 2 min. After allowing the reaction to proceed, FT-ICR MS analysis revealed that Grx displayed a Δ mass of -2Da, consistent with oxidation of Grx and formation of an intramolecular disulfide bond; and the *Bc*BCP species underwent a mass shift of -305 Da, consistent with reduction of the glutathione moiety, and displayed an isotope distribution consistent with *Bc*BCP containing two reduced cysteines (Figure 5). This result indicates that Grx can efficiently reduce the *Bc*BCP-GSH mixed disulfide.

In light of these mass spectrometric results, *Bc*BCP was assayed for peroxiredoxin activity using an NADPH-linked GSH/Grx/glutathione reductase system, and was monitored spectrophotometrically by measuring the decrease in A_{340} . As expected, NADPH oxidation was observed in the presence of BCP and activity increased in a BCP concentrationdependent manner (Figure 6). *Bc*BCP displayed an activity of $6.10 \pm 0.04 \,\mu\text{Mmin}^{-1}\mu\text{M}^{-1}$ and was dependent on the presence of Grx and GSH in the reaction. Interestingly, this activity is greater than 5-fold higher than the activity using Trx/TrxR as a reducing system under similar conditions.

Introduction of Cys49 converts Burkholderia BCP to an atypical 2-Cys Prx. To investigate the differences between the *E. coli* and *Burkholderia* BCP homologues, a resolving cysteine was introduced into *Bc*BCP (L49C) at an equivalent position to the resolving cysteine of the *E. coli* enzyme (Cys-50).

The activity and catalytic mechanism of L49C *Bc*BCP was examined by mass spectrometry. Oxidation of L49C *Bc*BCP with H_2O_2 resulted in a Δ mass of -2 Da, which we attribute to the loss of two hydrogen atoms, consistent with the formation of an intramolecular disulfide bond between two cysteines (Figure 7A and 7B). This disulfide bond was formed upon treatment with peroxide both with and without the presence of 1 mM GSH. Disulfide bond formation was confirmed by treatment of the oxidized protein with NEM, which resulted in the modification of only a single thiol group (data not shown). The site of NEM-modification was examined by top-down fragmentation. CID fragmentation revealed that Cys-98 was modified by NEM, and that Cys-44 and Cys-49 formed the intramolecular disulfide bond (Figure 7C and Supporting Information Figure S4). Together, these results demonstrate that the L49C *Bc*BCP mutant acts via the atypical 2-Cys Prx mechanism, in a manner similar to *Ec*BCP.

Having demonstrated by MS that L49C *Bc*BCP functions as an atypical 2-Cys Prx, we investigated the activity of the enzyme using the previously described enzyme assays. Although less active than the wildtype enzyme, L49C *Bc*BCP displays an activity of $0.59 \pm 0.06 \,\mu\text{Mmin}^{-1}\mu\text{M}^{-1}$ using Trx/TrxR as the reductive partners (Figure 1), confirming that the Cys44-Cys49 intramolecular disulfide bond can be resolved by the Trx/TrxR redox system. This activity was not inhibited by the presence of GSH. L49C BCP was also compared to the native *E. coli* enzyme, as the catalytic mechanism of these enzymes appears identical by MS. The activity of the native *E. coli* enzyme was 4-fold greater than that of L49C *Bc*BCP (Figure 1). Indeed, *Ec*BCP displayed greater activity than all isoforms of *Bc*BCP (WT and sitedirected mutants) using this assay with Trx/TrxR as the reductive partners (Figure 1). Analyzing the activity of the BcBCP L49C using the GSH/Grx system, we found that introduction of the cysteine resulted in an activity of $0.47 \pm 0.09 \,\mu\text{Mmin}^{-1}\mu\text{M}^{-1}$, a 15-fold decrease in comparison to wild-type BcBCP (Figure 6). We also assayed the activity of WT *Ec*BCP using this assay. Again, this enzyme showed significantly lower activity than WT BcBCP (Figure 6). These results suggest that the GSH/Grx system is not an efficient reducing partner for peroxiredoxins that catalyze peroxide reduction via the 2-Cys mechanism. Presumably this system cannot reduce the intramolecular disulfide formed in the reaction pathway of these enzymes. To test this hypothesis, the redox exchange reaction between oxidized BcBCP L49C and reduced glutaredoxin/GSH was monitored by MS. An equimolar mixture of oxidized BcBCP L49C and reduced glutaredoxin was allowed to equilibrate for 2 min in the presence of 1 mM GSH, before analysis by FT ICR-MS. As expected, analysis of the isotopic abundances within the resulting mass spectrum revealed that both species were unchanged (Supporting Information Figure S5, top) - highlighting the inability of Grx/GSH to reduce the Cys44-Cys49 disulfide bond within BcBCP L49C. In a similar experiment, no reduction of oxidized wild-type EcBCP by Grx/GSH was observed by FT-ICR MS (data not shown). In contrast, MS analysis of the redox exchange reaction between BcBCP L49C and reduced thioredoxin, demonstrates that thioredoxin can reduce oxidized *Bc*BCP L49C. However, it is interesting to note that to fully reduce BcBCP L49C, thioredoxin was added in 5-fold molar excess (Supporting Information Figure S5, bottom). These MS results correlate with the enzyme assay results (presented here and in ref. 3) which suggest that Trx is the preferred redox partner for the 2-Cys Prxs *Ec*BCP and *Bc*BCP L49C.

The Burkholderia bcp gene shows unique linkage and coexpression with the arn locus. We previously reported that the gene encoding *Bc*BCP (BCAL1936; *B. cenocepacia* J2315 gene nomenclature) is coexpressed with the *arn* locus (BCAL1929-1935) (7). Here, we further investigated this intriguing linkage of two different, but complementary, resistance determinants through comparative genomic analyses and quantitative gene expression studies.

Analysis of publicly available genomes that contain a *bcp* homologue revealed that the linkage of *bcp* with the *arn* locus is unique to β -proteobacteria that exhibit innate resistance to the polymyxin class of antimicrobials (*Burkholderia* species and the closely-related *Ralstonia pickettii* lineage). The genetic organization of the locus is shown in Figure 8. In the majority of polymyxin-sensitive β -proteobacteria studied, including *Bordetella*, *Comamonas* and *Delftia* species, the *bcp* gene is flanked by homologues of BCAL1928 and BCAL1937 without the intervening *arn* locus. In contrast, in γ -proteobacteria (including *Salmonella*, *Shigella* and *Pseudomonas* species), there is conserved linkage of the *bcp* and *gcvR* genes, the latter coding for the glycine cleavage repressor protein (*17*). This is consistent with the previous report that the *E. coli bcp* gene is located immediately downstream of the *gcvR* gene, but expressed independently (*18*). It should be noted that β -proteobacteria generally do not possess orthologs of GcvR (*19*).

Northern blot analysis of *B. cenocepacia* K56-2 at logarithmic phase revealed the presence of an abundant *bcp* transcript of approximately 830 bp, suggesting that *bcp* is strongly expressed independently, in addition to its previously described coexpression with the *arn* locus (7). We previously reported that the *Burkholderia arn* locus is expressed as two transcriptional units. Consequently, the independent expression of *bcp* was investigated further by QRT-PCR, comparing the level of *bcp* expression with that of representative genes from each transcriptional unit of the *arn* locus (*arnT* and *arnB*). The abundance of *bcp* transcript was approximately 11-fold higher than both *arnT* and *arnB* (Supporting Information Figure S6),

confirming the strong independent expression of *bcp*. A comparable ratio of transcripts was observed irrespective of the growth phase studied (data not shown).

Burkholderia BCP confers growth-phase-dependent resistance to oxidative killing. Insertional inactivation of the *bcp* gene in *B. cenocepacia* K56-2 resulted in strain XOA14. To explore the role of *Bc*BCP, the sensitivity of K56-2 and XOA14 to oxidative killing was assessed at regular timepoints throughout the growth phase. As shown in Figure 9, XOA14 is hypersensitive to H_2O_2 -mediated killing, however this hypersensitivity is restricted to the lag phase / early log phase of growth. At stationary phase, there is no difference in the survival of XOA14 and K56-2 following treatment with H_2O_2 . The hypersensitivity of XOA14 to oxidative killing at lag phase could be rescued through complementation with WT *Bc*BCP encoded on plasmid pARB10 (Supporting Information Figure S7).

Burkholderia BCP is not required for intracellular survival. Having demonstrated hypersensitivity to oxidative killing, the ability of XOA14 to survive within macrophages was assessed, and compared with K56-2. Cumulative evidence suggests that Bcc do not actively replicate intracellularly (20, 21). Consequently, we consider the use of stationary phase bacteria to be most physiologically relevant for Bcc intracellular survival assays. However, given the growth phase-dependent hypersensitivity of XOA14 to peroxide-mediated killing, we investigated intracellular survival and/or intracellular localization using both early log phase and stationary phase bacteria.

As shown in Figure 10A, intracellular survival of early log phase XOA14 is no different from that of K56-2, suggesting that the role of BcBCP is to protect from oxidative stresses in the extracellular rather than intracellular environment. Stationary phase bacteria were then used to compare the localization of K56-2 and XOA14 within macrophages (Figure 10B). The persistence of Bcc within macrophages in the apparent absence of replication has been

associated with their ability to delay the maturation of the phagosome, persisting instead within *B. cepacia*-containing vacuoles (BcCVs) (for review, see 22). Key events in the maturation of phagosomes include their progressive acidification and ultimately fusion with lysosomes. These processes can be monitored using LysoTracker Red and TMR-dextran, which (when visualized by fluorescence microscopy) label acidic organelles and lysosomes respectively. Microscopy of infected macrophages allows BcCVs to be visualized, and the percentage of BcCVs colocalizing with either LysoTracker Red or TMR-dextran can be calculated. A strain that is impaired in its ability to persist intracellularly (by delaying phagosomal maturation) would be expected to yield a higher proportion of LysoTracker Red and/or TMR-dextran-colocalizing BcCVs. The BCP-deficient XOA14 displays a marginally higher proportion of LysoTracker Red- and TMR-dextran-colocalizing BcCVs 4 h post infection compared to K56-2, but these differences are not statistically significant (Figure 10B). Therefore, we conclude that *Bc*BCP is not critical for the intracellular survival of *B. cenocepacia* in murine macrophages.

DISCUSSION

BCP is a thiol-dependent Prx of the AhpC/TSA family that is present in diverse bacterial species. Recently, we re-classified EcBCP, the founder member of this group of enzymes, as an atypical 2-Cys Prx on the basis of high resolution MS analysis (*3*). In the present study, we describe the BCP of *B. cenocepacia*, a novel member of the BCP family and an addition to the repertoire of 'superoxide detoxification' gene products recently identified within the *B. cenocepacia* J2315 genome (*23*). We provide the first detailed characterization of the 1-Cys BCP-peroxidatic mechanism, and demonstrate that BCP homologues fall into one of two functional groups, based on the presence or absence of a resolving cysteine (Figure 11). In the presence of the resolving cysteine, as typified by EcBCP, the sulfenic acid intermediate that forms following oxidation of the peroxidatic cysteine is rapidly reduced through the formation

of an intramolecular disulfide bond with the resolving cysteine (atypical 2-Cys Prx mechanism). Crucially, this occurs even in the presence of excess GSH (*3*), indicating that in the presence of the resolving cysteine, the formation of the intramolecular disulfide bond is strongly favored over the formation of a BCP-GSH heterodimer. This intramolecular disulfide bond is subsequently reduced by thioredoxin, completing the catalytic cycle. In contrast, *Bc*BCP described in the present study, lacks the resolving cysteine. This enzyme displays moderate enzymatic activity using thioredoxin, presumably with Trx directly reducing the sulfenic acid intermediate. However, here we demonstrate that in the presence of GSH this Trx-dependent catalysis is inhibited. Furthermore, the enzyme forms a rapid BCP-GSH heterodimer following oxidation of the peroxidatic cysteine, in a manner previously described by Noguera-Mazon *et al. (14)*. We show that this resolving complex is not a substrate for reduction by thioredoxin, but by the redox partner glutaredoxin. Introduction of a resolving cysteine (L49C) converts the *Burkholderia* enzyme to an atypical 2-Cys Prx that can be reduced by thioredoxin, thus retaining enzyme activity. A schematic of the WT and L49C *Bc*BCP reaction mechanisms is shown in Figure 11.

BCP homologues found in the majority of bacterial species lack the equivalent of the *Ec*BCP resolving cysteine (Cys-50) (Supporting Information Figure S8 and ref. *3*). Of 59 BCP proteins represented in Figure S8, only 18 have a cysteine residue at the equivalent position of the *E. coli* Cys-50, and these 18 BCP homologues are almost exclusively found in a restricted set of gamma proteobacteria. The majority of bacterial BCP homologues, from taxonomically diverse bacterial species, lack an apparent resolving cysteine and are presumed to exhibit the 1-Cys mechanism, typified by the native *Burkholderia* enzyme.

We previously reported that the engineered C50S *Ec*BCP retains activity through a novel catalytic mechanism that is distinct from the 1-Cys pathway (*3*). In contrast, we have demonstrated in the current study that the native *Bc*BCP that lacks the equivalent resolving

cysteine functions via a GSH/Grx dependent 1-Cys pathway. Interesting, we also assessed the activity of the EcBCP C50S mutant in the Grx assay (Figure 6). This single-cysteine BCP mutant displays activity approaching the level of BcBCP in this assay, suggesting it can also act via a BCP-GSH mixed disulfide resolving complex. In our previous study, MS analysis of the *Ec*BCP C50S was performed without the presence of GSH. MS analysis of this enzyme after oxidation with peroxide was repeated in the presence of 1 mM GSH and reveal a mixture of both BCP-GSH and BCP-BCP mixed disulfide dimers (data not shown). Thus, it seems that two resolving-pathways can be used by this BCP-variant. In contrast, no evidence for intermolecular disulfide bond formation and BCP dimers was found for WT BcBCP. Comparison of the amino acid sequences of representative BCP homologues shows that in atypical 2-Cys enzymes that possess an equivalent resolving cysteine to *EcBCP* (Cys-50), the conserved region that encompasses the peroxidatic cysteine extends up to (and beyond) the resolving cysteine (Supporting Information Figure S9). In contrast, BCP homologues that lack the resolving cysteine are divergent after the active TPGCT motif. It may be that these differences in amino acid sequence and thus architecture around the peroxidatic cysteine account for the subtle differences observed between the engineered C50S EcBCP enzyme and the native Burkholderia enzyme. There may be merit in high-resolution 3D structural studies of these BCPs to further explore the link between these isoform-specific sequence differences and catalytic mechanism. Such studies of the BCP homologue of Xanthomonas campestris (XcBCP) have recently been reported (24). Whilst XcBCP possesses a resolving cysteine and functions as an atypical 2-Cys Prx, it differs from *Ec*BCP in the positioning of the resolving cysteine relative to the peroxidatic cysteine. The peroxidatic and resolving cysteines of *EcBCP* (Cys-45 and Cys-50, respectively) are present within the same α -helix, but the equivalent cysteines of XcBCP (Cys-48 and Cys-84, respectively) are in different α -helices (24). Consequently, the proposed pathway of alternate opening and closing of the substrate

entry channel and disulfide bond pocket in XcBCP (24) is unlikely to be directly applicable to the *E. coli* enzyme or, indeed, the 1-Cys *Bc*BCP. Equivalent studies of the *Ec*BCP and *Bc*BCP homologues are required to establish the structural changes that occur in the catalytic cycles of these 1-Cys and 2-Cys BCP variants.

The BCP-associated phenotype shows variation between different bacterial species. Whilst BCP-deficient B. cenocepacia and E. coli (2) are hypersensitive to hydrogen peroxide, BCPdeficient Helicobacter pylori are not (25). This variable phenotype is most likely a consequence of differences in the repertoire and relative abundance of other anti-oxidant proteins within the cell. In the case of *H. pylori*, the 2-Cys AhpC peroxidase is considerably more abundant than BCP. Consequently, an *ahpC*-deficient *H. pylori* mutant is significantly more sensitive to oxidative killing than the bcp-deficient mutant (25). Indeed, given that we observed consistent expression (at the transcript level) of bcp throughout the growth phase of B. cenocepacia, we believe that the growth phase-dependent phenotype of the BCP-deficient strain XOA14 is a consequence of other anti-oxidant factors being strongly induced at late exponential / stationary phase, thus rendering BcBCP essentially redundant. Several antioxidants identified within the Bcc as playing a decisive role in intracellular survival show dramatically increased activities at stationary phase, including superoxide dismutase (SodC), catalase enzymes (including the bifunctional catalase-peroxidase, KatB) and a melanin-like pigment (26-29). Similarly, an apparent AhpC-deficient variant of B. cenocepacia C1394 is hypersensitive to oxidative killing at stationary phase relative to the parent strain (30). In B. cenocepacia, we propose that the Prx activity of BCP fulfils a protective role under physiological conditions that precede the induction of these stationary phase anti-oxidants, and that this protective role is restricted to the extracellular environment under conditions that favour active bacterial replication. It has been established that intracellular B. cenocepacia only minimally replicates in macrophages (21, 31). Therefore, it is possible that BcBCP

antioxidant activities are physiologically relevant in actively metabolic bacteria, as it has been observed for the *B. cenocepacia* KatA protein that is required for anti-oxidant protection of enzymes of TCA cycle (29). How *Bc*BCP activity, and indeed that of other anti-oxidants, contributes to the survival of *B. cenocepacia* within the specific environment of the CF lung is unclear. Intriguingly, *Pseudomonas aeruginosa* isolates from CF infants show reduced expression of *bcp* relative to that observed in the wound isolate, and common laboratory strain, PAO1 (RK Ernst; Personal Communication). Furthermore, loss of the related antioxidant AhpC in *B. cenocepacia* has been associated with the ability to establish persistent pulmonary infection in a murine host (*30*). This latter observation suggests that *Burkholderia* AhpC may be immunogenic, as has been observed for *Salmonella typhimurium* AhpC (*32*). The immunogenicity of BCP has yet to be addressed, and further studies are required to establish the role that these bacterial anti-oxidants play in the adaptation to CF airways.

In conclusion, we have described a novel Prx enzyme of *Burkholderia*, and characterized its catalytic activity. Although *Bc*BCP does not appear to play a pivotal role in the intracellular survival of *B. cenocepacia*, it confers resistance to ROS at a time when other known anti-oxidant mediators are not expressed, or expressed only at a low level. Whilst homologous to the previously described atypical 2-Cys BCP peroxidase of *E. coli*, the *Burkholderia* enzyme lacks the resolving cysteine and functions via a 1-Cys peroxidatic mechanism utilising an external small molecule thiol redox partner, glutathione, to resolve the oxidized peroxidatic cysteine. The resulting glutathionylated-*Bc*BCP intermediate is subsequently reduced by glutaredoxin. These studies provide the first detailed characterization of the 1-Cys Prx pathway in a BCP, and highlight the mechanistic differences that exist within the BCP family of bacterial peroxiredoxins and support their subdivision into two classes based on their catalytic activity.

SUPPORTING INFORMATION AVAILABLE

Top-down fragmentation of H₂O₂-treated WT *Bc*BCP, NEM-modified WT *Bc*BCP and NEMmodified L49C *Bc*BCP. MS analysis of the reduction of BcBCP-GSH heterodimer by thioredoxin. MS analysis of the reduction of oxidized BcBCP L49C by glutaredoxin and thioredoxin. Quantitative RT-PCR analysis of *bcp* expression. Complementation of BCPdeficient *B. cenocepacia* XOA14 with WT *Bc*BCP. Phylogenetic tree of bacterial BCP homologues. Amino acid alignments of representative BCP homologues with and without the resolving cysteine. This material is available free of charge via the Internet at http://pubs.acs.org.

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Strain or plasmid	Relevant characteristics	Source and/or reference
B. cenocepacia		
K56-2	ET12 clone related to genome sequence strain,	BCCM; (33)
XOA14	K56-2, <i>bcp</i> ::pGPΩTp, Tp ^r	(7)
ARB201	XOA14, pARB10 (encoding a functional <i>Bc</i> BCP protein), Tp ^r , Tet ^r	This study
E. coli strains		
DH5a	F Φ 80dlacZ Δ M15 Δ (lacZYA-argF)U169 recA1 endA1 hsdR17(r_k , m_k) phoA supE44 λ thi-1 gyrA96 relA1	Invitrogen
BL21(DE3)	$F mpT hsdS_B(r_B m_B) gal dcm$ (DE3)	Novagen
Plasmids		
pET-28a	ori_{pBR322} , Kan ^r , P_{T7lac}	Novagen
pDA17	ori_{pBBR1} , Tet ^r , mob^+ , P_{dhfr}	D. Aubert
pARB1	pET28a encoding WT <i>Bc</i> BCP	This study
pARB2	pET28a encoding C44S BcBCP	This study
pARB3	pET28a encoding C98S BcBCP	This study
pARB4	pET28a encoding L49C BcBCP	This study
pARB5	pET28a encoding WT <i>Ec</i> BCP	(3)
pARB10	pDA17 encoding WT BcBCP	This study

TABLE 1. Strains and plasmids

Tp^r, trimethoprim resistance; Tet^r, tetracycline resistance; Kan^r, kanamycin resistance;

BCCM, Belgian Co-ordinated Collection of Micro-organisms.

FIGURE LEGENDS

Figure 1. Thiol-dependent peroxidase activity of *Bc*BCP using thioredoxin as the reducing partner. The thioredoxin linked peroxidase activity of recombinant BCP proteins was assessed in an assay coupled to the oxidation of NADPH (monitored by a decrease in A_{340nm}) and measured in μ M NADPH oxidized per min per μ M of enzyme (μ Mmin⁻¹ μ M⁻¹). The assay contained 0.36 μ M TrxR, 1.7 μ M Trx, 0.1 mM H₂O₂ and 0.2 mM NADPH in 50 mM Hepes-NaOH (pH 7). '+GSH' indicated the presence of 1 mM reduced GSH in the reaction mixture. In the absence of BCP, a background (B/ground) level of NADPH oxidation occurs, equating to 0.1 μ Mmin⁻¹. Enzyme activity is stated as the mean of three independent results and the error bars indicate one standard deviation.

Figure 2. Mass spectrometry analysis of *Bc*BCP and H₂O₂-treated *Bc*BCP. *Bc*BCP was analysed by ESI-FT-ICR MS in H₂O:MeOH:HCOOH (v/v/v 50:48:2) at a final concentration of 10 μ M. (*A*, *insert*) A typical spectrum of *Bc*BCP displaying a charge state distribution from [M+11H]¹¹⁺ to [M+23H]²³⁺ and an average deconvoluted mass of 18992 Da. (*A*) The most abundant ion, [M+17H]¹⁷⁺, has an isotope distribution consistent with a theoretical isotope distribution of *Bc*BCP containing 2 reduced cysteine residues (red circles; empirical formula [C₈₄₉H₁₃₅₈N₂₃₉O₂₄₆S₅]¹⁷⁺). (*B*) *Bc*BCP after addition of 500 μ M H₂O₂. The isotope distribution of the [M+17H]¹⁷⁺ ion reveals a mass shift of +32 Da, to an average mass of 19024 Da, which is consistent with the addition of 2 oxygen atoms, attributed to oxidation of cysteine to sulfinic acid (blue circles, empirical formula [C₈₄₉H₁₃₅₈N₂₃₉O₂₄₈S₅]¹⁷⁺). (*C*) Prosight-PTM output file (cleavage map) showing the top-down fragmentation of sulfinic acid-modified *Bc*BCP. ECD of H₂O₂-treated *Bc*BCP allowed the modification to be mapped to a 3-amino acid section containing Cys-44 (see also Supporting Information Figure S1).

Figure 3. Amino acid alignment of *B. cenocepacia* **and** *E. coli* **BCP.** *Bc*BCP and *Ec*BCP enzymes have two cysteine residues in common (black arrowheads), at positions Cys-44 and Cys-98 (*B. cenocepacia* co-ordinates). *Ec*BCP has an additional cysteine at Cys-50 (grey arrowhead) that is not conserved in the *Burkholderia* enzyme. Both amino acid sequences are shown without the N-terminal methionine, which was cleaved from all purified BCP proteins studied.

Figure 4. Glutathione can act as a resolving-thiol in *Bc*BCP catalysis. *Bc*BCP was analysed by ESI-FT-ICR MS in H₂O:MeOH:HCOOH (v/v/v 50:48:2) at a final concentration of 10 μ M. (*A*) Typical mass spectrum of *Bc*BCP in the presence of 1mM GSH. The isotope distribution of the [M+19H]¹⁹⁺ charge state is consistent with *Bc*BCP containing two reduced cysteine residues (red circles; empirical formula [C₈₄₉H₁₃₆₀N₂₃₉O₂₄₆S₅]¹⁹⁺), average deconvoluted mass 18992 Da. For comparison, the theoretical isotope distribution of *Bc*BCP containing one disulfide bond ([C₈₄₉H₁₃₅₈N₂₃₉O₂₄₆S₅]¹⁹⁺), is shown in green circles. (*B*) Addition of 1 molar equivalent of H₂O₂ resulted in a second species of average mass 19297 Da (Δ mass +305 Da). This species displays an isotope distribution consistent with *Bc*BCP containing one reduced cysteine and one glutathionylated cysteine (claret circles, empirical formula [C₈₅₉H₁₃₇₅N₂₄₂O₂₅₂S₆]¹⁹⁺). (*C*) This species was subsequently derivatized with the alkylating agent NEM and reduced with DTT before top-down fragmentation using CID. The Prosight-PTM output file (cleavage map) shows the assignment of the NEM modification to Cys-98 allowing the assignment of the glutathionylated cysteine to Cys-44 (see also Supporting Information Figure S2).

Figure 5. Analysis of potential reductive partners for *Bc*BCP catalysis using mass spectrometry. Glutaredoxin can efficiently reduce glutathionylated *Bc*BCP. An equimolar mixture of *Bc*BCP-GSH and reduced Grx was incubated at room temperature for 2 min before being quenched with H₂O:MeOH:HCOOH (v/v/v 50:48:2). The resulting mixture was analyzed by FT-ICR MS at a final protein concentration of 10 μ M. (*A*) Analysis of the isotope distributions of *Bc*BCP-GSH and Grx before mixing. *Bc*BCP-GSH displays an isotope distribution consistent with the glutathionylated protein (claret circles;

 $[C_{859}H_{1375}N_{242}O_{252}S_6]^{19+})$. The $[M+9H]^{9+}$ charge state of the glutaredoxin species displayed an isotope distribution consistent with reduced Grx (red circles; $[C_{429}H_{671}N_{116}O_{134}S_3]^{9+})$. The theoretical isotope distribution of oxidized Grx is shown in green circles for comparison $([C_{429}H_{669}N_{116}O_{134}S_3]^{9+})$. (*B*) Analysis of the isotope distributions of *Bc*BCP and Grx after mixing. The *Bc*BCP species displays an isotope distribution consistent with the reduced protein, indicating loss of the GSH moiety (red circles; $[C_{849}H_{1360}N_{239}O_{246}S_5]^{19+}$). The $[M+9H]^{9+}$ charge state of the glutaredoxin species displayed an isotope distribution consistent with oxidised Grx (green circles; $[C_{429}H_{669}N_{116}O_{134}S_3]^{9+}$) consistent with the formation of a disulfide bond. (*C*) Broad-band mass spectrum showing the charge state distributions of reduced *Bc*BCP and oxidized Grx. BCP charge states are indicated by red squares; Grx charge states are indicated by green triangles.

Figure 6. Peroxidase activity of BCP enzymes using glutaredoxin as the reducing

partner. The GSH/glutaredoxin linked peroxidase activities of recombinant BCP proteins were assessed in an assay coupled to the oxidation of NADPH (monitored by a decrease in A_{340nm}) and measured in μ M NADPH oxidized per min per μ M of enzyme (μ Mmin⁻¹ μ M⁻¹). The assay contained 1 unit glutathione reductase, 12 μ M Grx, 1mM GSH, 0.1 mM hydrogen peroxide and 0.15 mM NADPH in 50 mM Hepes-NaOH (pH 7). Enzyme activity is stated as the mean of three independent results and the error bars indicate one standard deviation.

Figure 7. Introduction of a resolving cysteine (L49C) converts BcBCP from a 1-Cys Prx to an atypical 2-Cys Prx. BcBCP L49C was analysed by ESI-FT-ICR MS in H₂O:MeOH:HCOOH (v/v/v 50:48:2) at a final concentration of 10 µM. (A) Typical mass spectrum of L49C *Bc*BCP displaying a charge state distribution from $[M+8H]^{8+}$ to [M+21H]²¹⁺ and an average mass of 18982 Da. Analysis of the isotope distribution of the [M+19H]¹⁹⁺ charge state reveals a distribution consistent with L49C BcBCP containing three reduced cysteines (red circles; empirical formula $[C_{846} H_{1354} N_{239} O_{246} S_6]^{19+}$). (B) Mass spectrum of L49C BcBCP after addition of 1 molar equivalent of H₂O₂, in the presence of 1mM GSH, displays a charge state distribution from $[M+8H]^{8+}$ to $[M+17H]^{23+}$. The isotope distribution of the $[M+19H]^{19+}$ reveals a Δ mass of -2 Da and is consistent with L49C *Bc*BCP containing one disulfide bond and one reduced cysteine (green circles; empirical formula [C₈₄₆ H₁₃₅₂ N₂₃₉ O₂₄₆ S₆]¹⁹⁺). (C) H₂O₂-treated L49C BcBCP was subsequently derivatized with the alkylating agent NEM and reduced with DTT before top-down fragmentation using ECD. The Prosight-PTM output file (cleavage map) shows the assignment of the NEM modification to Cys-98 allowing the assignment of the disulfide bond to Cys-44 and Cys-49 (see also Supporting Information Figure S4).

Figure 8. Genetic linkage of *bcp* **in the genus** *Burkholderia***.** Within β-proteobacteria that exhibit innate resistance to cationic antimicrobial peptides (*Burkholderia* species and organisms of the *Ralstonia pickettii* group) *bcp* (black arrow) is located immediately downstream of the *arn* locus (grey arrows). The flanking genes BCAL1928 and BCAL1937 (white arrows) encode a hypothetical protein and a PhoH-like protein respectively. 'BCAL' gene nomenclature refers to the genome sequence of *B. cenocepacia* J2315, chromosome 1.

Figure 9: Growth-phase dependent role of *Bc***BCP in protecting against oxidative stress.** At regular intervals throughout the growth phase, percentage survival of *B. cenocepacia* K56-2 and the BCP-deficient strain XOA14 was assessed following challenge of each strain with 1 mM H_2O_2 for 30 min. Both K56-2 and XOA14 showed comparable growth in aerobic conditions (closed squares and closed circles respectively). XOA14 is hypersensitive to H_2O_2 mediated killing at lag phase / early log phase of growth (open circles), compared to K56-2 (open squares). At stationary phase, both strains are equally resistant to H_2O_2 -mediated killing.

Figure 10: Comparison of intracellular survival and localization within murine macrophages of *B. cenocepacia* K56-2 and the BCP-deficient XOA14. Macrophage infection assays demonstrate that *Burkholderia* BCP is not required for intracellular survival. (*A*) Intracellular survival of early log phase *B. cenocepacia* K56-2 (grey bars) and XOA14 (white bars) was quantified 2 h and 4 h postinfection (MOI 50). The BCP-deficient XOA14 is not impaired in intracellular survival at early log phase relative to K56-2. (*B*) The number of *B. cenocepacia*-containing vacuoles (BcCVs) colocalizing with either TMR-Dextran (dex) or LysoTracker Red (Lys) was compared following either 2 h or 4 h infection of murine macrophages (ANA1 or Raw264.7) with stationary phase K56-2 (grey bars, 'K') or XOA14 (white bars, 'X'). The number of BcCVs colocalizing with TMR-dextran or LysoTracker Red is not significantly different when comparing the two strains, indicating that the stationary phase BCP-deficient XOA14 is not impaired in its ability to persist within macrophages.

Figure 11: Schematic of the peroxidatic mechanisms of 1-Cys and atypical 2-Cys BCPs.

(*A*) Catalytic mechanism of a 1-Cys BCP, as exemplified by wildtype *Bc*BCP. The enzyme functions via a GSH/Grx dependent 1-Cys Prx mechanism. In the presence of peroxide, the peroxidatic cysteine (Cys-44) is oxidized to a sulfenic acid intermediate. In the absence of a small molecule thiol, this sulfenic acid intermediate is irreversibly oxidized to sulfinic acid. However, in the presence of glutathione (GSH), a *Bc*BCP-GSH heterodimer is formed, which is subsequently resolved by reduced glutaredoxin. (*B*) Catalytic mechanism of an atypical 2-Cys BCP, as exemplified by L49C *Bc*BCP or *Ec*BCP. The peroxidatic cysteine is oxidized to

the sulfenic acid intermediate prior to disulfide bond formation with the resolving cysteine (Cys-49). The formation of this intramolecular disulfide bond is highly favoured, occurring even in the presence of excess GSH. The disulfide bridge between Cys-44 and Cys-49 can be resolved by the Trx/TrxR redox system.

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