

1 **Characterisation of a Bi-functional Catalase-peroxidase of *Burkholderia***  
2 ***cenoepecia***

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8 Panagoula Charalabous\*<sup>1</sup>, Janet M. Risk<sup>2</sup>, Rosalind Jenkins<sup>3</sup>,  
9 Andrew J. Birss<sup>1</sup>, C. Anthony Hart<sup>4</sup> and John W. Smalley<sup>1</sup>

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15 <sup>1</sup>Microbiology, and <sup>2</sup>Molecular Genetics and Oncology Groups, School of Dental  
16 Sciences, <sup>3</sup>Biomedical Sciences Proteomics Facility, Department of Pharmacology &  
17 Therapeutics,  
18 and <sup>4</sup>Department of Medical Microbiology and Genito-Urinary Medicine, The  
19 University of Liverpool, Daulby Street, Liverpool L69 3GN, UK

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26 Author for Correspondence: Dr John W Smalley

27 Tel +44 151 706 5272;

28 FAX: +44 151 706 5809;

29 E-mail: [josmall@liv.ac.uk](mailto:josmall@liv.ac.uk)

30  
31 \* Dr P Charalabous, current address: Department of Physiology, School of  
32 Biomedical Sciences, Crown Street, Liverpool L69 3BX, UK.  
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**1 Abstract**

2 Isolates of *Burkholderia cenocepacia* express a putative haem-binding protein  
3 ( $M_r$  97kDa) which displays intrinsic peroxidase activity. Its role has been re-  
4 evaluated, and we now show that it is a bi-functional catalase-peroxidase, with  
5 activity against tetramethylbenzidine (TMB), *o*-dianisidine, pyrogallol, and 2,2'-azino-  
6 bis(3-ethylbenzthiazoline-6-sulphonic acid (ABTS). Both peroxidase and catalase  
7 activities are optimal at pH 5.5-6.0. The gene encoding this enzyme was cloned and  
8 expressed in *E. coli*. We have named it *katG* because of its similarity to other *katGs*  
9 including that from *Burkholderia pseudomallei*. It is substantially similar to a  
10 previously described catalase-peroxidase of *B. cenocepacia* (*katA*). Mass  
11 spectrometric analysis indicated that the initial *katG* translation product may be post-  
12 translationally modified in *B. cenocepacia* to give rise to the mature 97kDa catalase-  
13 peroxidase.

14

## 1 INTRODUCTION

2 *Burkholderia cenocepacia* (genomovar IIIa) isolates are associated with life-  
3 threatening lung infections that may progress to septicaemia (“cepacia syndrome”) in  
4 cystic fibrosis (CF) patients [1]. These bacteria were shown to express a 97kDa  
5 putative haem-binding protein [2], but more recent studies [3] showed that this  
6 protein does not bind haem dose-dependently, and is peroxidase positive even  
7 without prior exposure to exogenous iron(III) protoporphyrin IX. These newer  
8 findings lead to the possibility that it is not a true haem-binding protein. In the  
9 present study, we re-evaluated the properties of the 97kDa protein and demonstrated  
10 it to be a bi-functional catalase-peroxidase. The gene encoding this protein was  
11 cloned and expressed in *Escherichia coli*, and a survey of its distribution in bacterial  
12 strains of the “*Burkholderia cepacia* complex” was undertaken.

13

## 14 Materials and Methods

15 **Bacterial strains and growth conditions.** *B. cenocepacia* clonal isolates  
16 strains BC7, C5424, C6433 and J2315 (expressing the 97kDa putative haem-binding  
17 protein) were maintained by subculture on horse-blood agar. For peroxidase and  
18 catalase studies the cells were sub-cultured three times on Columbia or M9 Minimal  
19 Salts Medium agar (Sigma Chemical Company) before growth in bulk on these solid  
20 media as lawn growths for 3 days. Cells were harvested into and washed twice in  
21 0.14M NaCl, 0.1M Tris-HCl, pH 7.5 to remove any contaminating growth medium  
22 constituents. For molecular genetic studies, *E. coli* strains were grown in Luria-  
23 Bertani (LB) broth or agar, supplemented with 100µg ampicillin ml<sup>-1</sup> where  
24 appropriate.

25 **SDS-PAGE and staining for peroxidase and catalase activity.** Cell  
26 samples of strain J2315 were solubilised in non-reducing sample buffer (37°C for 1h),  
27 electrophoresed on 10% acrylamide gels and stained with 3,3',5,5'-  
28 tetramethylbenzidine (TMB)/H<sub>2</sub>O<sub>2</sub> [2]. Gels were counter-stained with Coomassie  
29 Blue [3] to precisely identify the positions of peroxidase bands. Chromogenic  
30 peroxidase substrates *o*-dianisidine (3,3'-dimethoxybenzidine), pyrogallol (1,2,3-  
31 trihydroxybenzene), 4-chloronaphthol, guaiacol (2-methoxyphenol), and ABTS [2,2'-  
32 azino-bis(3-ethylbenzthiazoline-6-sulphonic acid (all at 6.3mM) were also tested  
33 using the same procedure as for TMB above. Catalase activity was detected in

1 samples as above using the  $K_3Fe(CN)_6/FeCl_3-H_2O_2$  method of Katsuwon and  
2 Anderson [4], in which catalase-positive bands appear as clear zones against a dark  
3 brown background.

4 **Effect of pH on the peroxidase and catalase activities.** Replicate gel  
5 tracks of cell samples (strain J2315; 25 $\mu$ g protein) were incubated, after  
6 electrophoresis, in 0.14M NaCl buffered at pH 9.0, 8.5, 8.0, or 7.5 with 0.1M Tris-HCl,  
7 or at pH 7.0, 6.5, 6.0, or 5.5, buffered with 0.1M sodium acetate/acetic acid, for 1h.  
8 The gel strips were developed in  $K_3Fe(CN)_6/FeCl_3-H_2O_2$  or TMB- $H_2O_2$  to reveal  
9 catalase and peroxidase bands, respectively.

10 **pH-activity profiles for whole cell catalase and peroxidase.** Peroxidase  
11 activity was measured by monitoring  $\Delta A_{645nm}$  after incubation of cell suspensions  
12 (J2315; 250 $\mu$ g protein  $ml^{-1}$ ) in the above buffers at 20°C with 6.3 mM TMB plus  
13 10mM  $H_2O_2$  [5]. Catalase activity was measured at 20°C in the above suspensions  
14 by monitoring  $\Delta A_{240nm}$  [6], using 8mM  $H_2O_2$  as substrate. Catalase and peroxidase  
15 activities were expressed as either pmole  $H_2O_2$  degraded or pmole TMB oxidised per  
16 minute.

17 **Protein separation and mass spectrometry.** Protein bands from  
18 Coomassie Blue-stained 1D gels were excised and destained and dehydrated by  
19 incubation with 50% acetonitrile/50mM ammonium bicarbonate for 1h at room  
20 temperature, followed by vacuum drying in a SpeedVac (Eppendorf). The gel pieces  
21 were rehydrated in 50mM ammonium bicarbonate containing 40ng/ $\mu$ L modified  
22 trypsin (Promega) and incubated for 16h at 37°C. Peptides were extracted from the  
23 gel by incubation with 2 changes of 60% acetonitrile/1% trifluoroacetic acid, and the  
24 resulting supernatants were again dried in a SpeedVac. The extracts were desalted  
25 using C18 ZipTips according to the manufacturer's instructions (Millipore), and were  
26 reconstituted in a final volume of 30 $\mu$ L 5% acetonitrile/0.1% trifluoroacetic acid.  
27 Aliquots of 0.5 $\mu$ L sample were spotted onto a MALDI target plate together with an  
28 equal volume of 5mg/mL  $\alpha$ -cyano-4-hydroxycinnamic acid (LaserBiolabs, France) in  
29 50% acetonitrile/0.1% trifluoroacetic acid. Peptide mass fingerprints were acquired  
30 either on a Voyager DE Pro MALDI (Applied Biosystems, California, USA) or a  
31 M@LDI (Micromass, Manchester, UK) instrument in positive ion reflector mode.  
32 Data were submitted for screening via the Mascot search engine (Matrix Science,  
33 London). The mass tolerance was set to 100ppm, and 1 missed cleavage and no

1 modifications were allowed. For LC-MS/MS analysis, aliquots of 5µL sample were  
2 delivered into a QSTAR Pulsar i hybrid mass spectrometer (Applied Biosystems) by  
3 automated in-line liquid chromatography (integrated LCPackings System, 5mm C18  
4 nano-precolumn and 75µm x 15cm C18 PepMap column (Dionex, California, USA))  
5 via a nano-electrospray source head and 10µm inner diameter PicoTip (New  
6 Objective, Massachusetts, USA). A gradient from 5% acetonitrile/0.05% TFA (v/v) to  
7 48% acetonitrile/0.05% TFA (v/v) in 60mins was applied at a flow rate of 300nL/min,  
8 and MS and MS/MS spectra were acquired automatically in positive ion mode using  
9 information-dependent acquisition (IDA) (Analyst, Applied Biosystems). Database  
10 searching was carried out using Mascot with mass tolerances set to 1.2Da for MS  
11 and 0.6Da for MS/MS, and with deamidation as a variable modification.

12 **Cloning of the catalase-peroxidase (*katG*) gene from *B. cenocepacia***  
13 **strain J2315.** Genomic DNA was amplified by PCR using the primers  
14 CACCATGTCTGAACGAAGGGCAGT and CGATGTACCACCGCTTTT. PCR was  
15 performed with an initial denaturation step at 94°C for 5 min, followed by 35 cycles  
16 each of 1 min at 94°C, 1 min at 58°C, 3.5 min at 72°C, and a final extension step at  
17 72°C for 5 min. PCR products were electrophoresed on a 1% (w/v) agarose gel and  
18 visualised using ethidium bromide to confirm a single product of the correct size  
19 before cloning into pET-100/D-TOPO (Invitrogen) to produce pKatG and  
20 transformation of One Shot<sup>®</sup> TOP 10 chemically competent *E. coli* cells (Invitrogen).  
21 Successful insertion of the gene in the correct orientation was confirmed by  
22 restriction analysis.

23 **Sequencing of *katG*.** The *katG* was sequenced from pKatG plasmid DNA  
24 using multiple overlapping IRD-700-labelled forward and reverse primers (MWG-  
25 Biotech Ltd., UK; details on request) and employing the SequiThermEXCEL<sup>™</sup> II DNA  
26 Sequencing Kit-LC (Epicentre Technologies) in a LI-COR 4200S auto-sequencer.  
27 The sequencing data were viewed using the LI-COR Base ImagIR<sup>™</sup> software in  
28 conjunction with the Sequencher programme (<http://www.genecodes.com>).

29 **Expression analysis.** BL21 Star<sup>™</sup> One Shot *E. coli* cells were transformed  
30 with the pKatG plasmid and induction of expression was undertaken by growth at  
31 37°C in the presence of 0.1M isopropyl β-D-thioglucoopyranoside (IPTG; Sigma).  
32 Pelleted cells were electrophoresed as above and assessed for catalase and  
33 peroxidase activity by in-gel staining.

34

## 1 Results

2 As previously reported [2], a major peroxidase-positive protein of 97kDa was  
3 seen for *B. cenocepacia* strain J2315 (Fig. 1, tracks b and h). The 97kDa enzyme  
4 also showed peroxidase specificity towards *o*-dianisidine, ABTS and pyrogallol (Fig.  
5 1, tracks c, e, and g), but guaicol (methoxyphenol) and 2-chloronaphthol were not  
6 peroxidised (Fig. 1, tracks d and f). Using in-gel  $K_3Fe(CN)_6/FeCl_3/H_2O_2$  staining, the  
7 97kDa peroxidase protein was also shown to display catalase activity (Fig 1, track a),  
8 a phenomenon also demonstrated by the clonal isolates BC7, C5424 and C6433  
9 (data not shown). The catalase activity of the 97kDa enzyme of strain J2315 was not  
10 inhibited by pre-exposure of the cells for 1h to the specific mono-functional catalase  
11 inhibitor 3-amino-1, 2, 4-triazole (20mM). Peroxidase activity against substrates  
12 TMB, ABTS and *o*-dianisidine was also unaffected by the inhibitors isonicotinic acid  
13 and niacinamide (all 20mM) [8] (data not shown). In general, the catalase activity  
14 visualised by the in-gel assay was stronger than the peroxidase staining.

15 Both catalase and peroxidase activities were observed in SDS-PAGE gels  
16 over the pH range 5.5 to 8.5, whilst little or no activity was seen at pH 9.0 (Fig. 2).  
17 Both enzyme activities were generally low at alkaline pHs, and highest over the acid  
18 pH range, and scanning densitometry confirmed maximal catalase and peroxidase  
19 activities at pH 6.0 and 5.5, respectively. Peroxidase activity of suspensions of whole  
20 cells of strain J2315 against TMB was not detected at neutral or alkaline pHs, but  
21 maximal activity was seen at pH 6.0 (Fig. 3a). In contrast, low levels of catalase  
22 activity of whole cells were observed at alkaline pH, rising in the acid pH range to a  
23 maximum at pH 6.0 (Fig 3b).

24 Masses of tryptic peptides of the 97kDa catalase-peroxidase were obtained by  
25 MALDI-TOF mass spectrometry and matches were obtained to the KatG catalase-  
26 peroxidases of *Burkholderia pseudomallei* [9] and *Mycobacterium tuberculosis* [10],  
27 and to an archeal catalase-peroxidase of *Haloarcula marismortui* [11]. This genomic  
28 data was aligned with the sequence of the J2315 strain  
29 ([http://www.sanger.ac.uk/Projects/B\\_cenocepacia/](http://www.sanger.ac.uk/Projects/B_cenocepacia/)) and used to design primers to a  
30 putative open reading frame (ORF) of 2211 bases (768 amino acids) on chromosome  
31 2 which was cloned and re-sequenced (accession number DQ112341). This ORF  
32 was subsequently identified in the published *B. cenocepacia* genome (Sanger  
33 Institute) and was calculated to have a size of 80.5kDa. A second ORF was also

1 identified at 3612912-3615098 on chromosome 1, which possessed 73% identity and  
2 81% similarity to the first catalase-peroxidase gene at the amino acid level and may  
3 represent another catalase-peroxidase protein.

4 Paired amino acid alignments (BLASTP) revealed a high degree of homology  
5 between the *B. cenocepacia* catalase-peroxidase and the minor catalase-peroxidase  
6 described by Lefebvre *et al.* [12], those of other selected bacterial species including  
7 the cell-surface catalase-peroxidase of *B. pseudomallei* (KatG), and the 77kDa iron  
8 (III) protoporphyrin IX monomer binding protein (accession number DQ114424;  
9 Smalley *et al.*, 2005) (Table 1). Multiple sequence alignment analysis using  
10 CLUSTALW revealed striking similarities between the *B. cenocepacia* catalase-  
11 peroxidase and the other selected enzymes in both the C- and N-terminal regions.  
12 These included the conserved amino acid triad Arg<sup>104</sup>-Trp<sup>107</sup>-His<sup>108</sup>, and the second  
13 haem ligand (His<sup>270</sup>) of *M. tuberculosis* KatG [13]. Because of the similarities of the  
14 catalase-peroxidase to these well-characterised KatG proteins, the gene encoding  
15 the *B. cenocepacia* enzyme was named *katG*.

16 IPTG induction of the *E. coli* BL21Star cells carrying the 80.5kDa catalase-  
17 peroxidase gene resulted in a protein product which electrophoresed on 10% gels as  
18 a single band with an apparent molecular mass of ~ 80kDa (Fig 4). The  
19 recombinant protein stained positively for both peroxidase and catalase showing the  
20 gene product to be functionally active (Fig 4). To confirm the identity of the  
21 recombinant enzyme expressed in *E. coli* as the product of *katG*, MS/MS analysis  
22 was performed after SDS-PAGE and trypsin digestion. On 7% acrylamide gels it was  
23 found that the recombinant protein was separated into two bands with calculated  
24 molecular masses of 79 and 83kDa, denoted R1 and R2, respectively (Fig 5), both  
25 of which were positive for peroxidase (data not shown). The observation of the band  
26 R2 is in keeping with the expected size of an initial translation product based on the  
27 vector system employed which results in the addition of 36 amino acids to the N-  
28 terminus of the expressed protein. We speculate that the lower molecular weight  
29 band R1 arises as a result of proteolytic cleavage of the initial translation product. In  
30 addition to proteins R1 and R2, a very faint Coomassie Blue stained band of  
31 approximately 97kDa (denoted R3) was observed which was not expressed by *E. coli*  
32 cells carrying the empty plasmid. This band was peroxidase positive as revealed by  
33 TMB staining for a longer time period (data not shown). MS/MS analysis of these  
34 three proteins showed them to match *B. cenocepacia* KatG (Table 2). Taken

1 together these data confirmed that the product of *katG* was the bifunctional catalase-  
2 peroxidase. The presence of a higher molecular weight form of the enzyme suggests  
3 that the initial translation product may be post-translationally modified to give the  
4 mature 97kDa catalase-peroxidase.

5

## 6 **Discussion**

7 We have re-evaluated the role of the 97kDa putative haem-binding protein of  
8 *B. cenocepacia* [2]. This protein shows peroxidase specificity towards  
9 tetramethylbenzidine, *o*-dianisidine, pyrogallol, and 2,2'-azino-bis(3-  
10 ethylbenzthiazoline-6-sulphonic acid (ABTS), but not against 4-chloronaphthol and 2-  
11 methoxyphenol (guaicol). It also has catalase activity, but is not inhibited by the  
12 specific mono-functional catalase inhibitor 3-amino-1,2,4,-triazole. This protein does  
13 not show dose-dependent binding of iron(III) protoporphyrin IX in either the  
14 monomeric or  $\mu$ -oxo oligomeric form [3], and does not bind to haem-agarose  
15 (Smalley et al., unpublished findings). Collectively, these data show this component  
16 is not a true haem-binding protein, but a bi-functional catalase-peroxidase, in contrast  
17 to the 77- and 149kDa iron(III) protoporphyrin IX-binding, outer-membrane  
18 components which do not possess intrinsic catalase activity, and which are only  
19 peroxidase-positive after exposure to, and binding of, iron(III) protoporphyrin IX  
20 monomers [3].

21 Multiple amino acid alignment analysis of the translated *B. cenocepacia*  
22 catalase-peroxidase gene revealed a strong homology with other bacterial catalase-  
23 peroxidases, and supported the above biochemical observations. It possessed the  
24 Arg<sup>88</sup>-Trp<sup>91</sup>-His<sup>92</sup> triad which is conserved among all known catalase-peroxidases  
25 [14], and displayed the greatest cross-species amino acid homology (70.7% identity  
26 and 79% similarity) to that of *B. pseudomallei* KatG, a homo-dimer of subunit size  
27 81.6kDa [15, 16], which plays a role in protecting against hydrogen peroxide [17].  
28 For this reason, the *B. cenocepacia* catalase-peroxidase gene was named *katG*.  
29 A catalase-peroxidase gene *katA* (accession number AF317697) has recently been  
30 described in *B. cenocepacia* strain C5424 by Lefebvre *et al.* [12], which is similar to  
31 the gene identified herein.

32 MS/MS analysis clearly demonstrated that the recombinant catalase-  
33 peroxidase was the product of *katG*. The detection of a higher molecular weight  
34 enzyme matching KatG shows that expression of *katG* in *E. coli* is also accompanied



1 some post-translational processing and suggests that this step may be more efficient  
2 in *B. cenocepacia*, giving rise to the mature 97kDa catalase-peroxidase. At present  
3 however, the nature of any post-translational modifications is unclear. The  
4 electrophoretic mobility of the native 97kDa enzyme from *B. cenocepacia* does not  
5 change upon reduction with dithiothreitol, nor does it react with phosphoprotein stains  
6 or periodic acid-Schiff reagent (data not shown).

7         Recent *B. cenocepacia* J2315 sequence database releases indicate that the  
8 *katG* gene (BCAM2107) may actually be extended by 20 amino acids at the N  
9 terminus to give a 756 amino acid, 82.6kDa, protein. This, together with other  
10 sequence differences that we have noted between our data, the *B. cenocepacia*  
11 J2315 database sequence and the *katA* gene of Lefebvre *et al.* [12], may indicate that  
12 the ORF is not yet correctly identified. Lefebvre *et al.* [12] also demonstrated a  
13 second catalase-peroxidase gene in *B. cenocepacia* J2315. We confirm the  
14 presence of this second gene (BCAL3299) and observe that it is 73% identical and  
15 81% similar to our *B. cenocepacia* *katG* at the amino acid level.

16         *B. cenocepacia* *katA* mutants are sensitive to H<sub>2</sub>O<sub>2</sub> and *katA* also appears to  
17 contribute to the normal functioning of the TCA cycle [12], but the extent to which the  
18 catalase-peroxidase described herein contributes to growth and survival *in vivo* is not  
19 clear. Although the pH of the liquid surface layer of the lung in health is  $\approx$  6.9 [18],  
20 endo-bronchial pHs of around 6.5 have been recorded. In addition, respiratory  
21 mucins, to which *B. cenocepacia* binds specifically [19], are highly sulphated,  
22 especially those produced by CF patients [20, 21, 22], and this also contributes to the  
23 acidity of the secretions. The pH is also reduced as a result of defective  
24 transmembrane conductance regulator function [23] and the mucopurulent secretions  
25 formed in the CF lung during infection also have an acid pH [24, 25]. In view of the  
26 above and the acid pH optima of the catalase-peroxidase, it is likely that bacterial  
27 cells expressing this enzyme would be advantaged in enduring attack by  
28 macrophage-derived H<sub>2</sub>O<sub>2</sub> in the slightly acidic conditions prevailing in the lung  
29 during chronic infection and inflammation. Although members of the "*B. cepacia*  
30 complex" display catalase and peroxidase activities [8], we have generally found that  
31 these activities in other species of the complex are very low compared to *B.*  
32 *cenocepacia* strains [26]. Bacterial catalase-peroxidases display wide substrate  
33 specificities [14, 27], but it is not known which compounds represent natural  
34 peroxidase substrates for the *B. cenocepacia* enzyme or whether it plays any role in

1 attacking and degrading other host (macro)molecules for defensive or nutritive  
2 purposes.

3

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## 1 **References**

- 2 [1] Hart CA & Winstanley C (2002) Persistent and aggressive bacteria in the lungs of  
3 cystic fibrosis children. *Brit Medical Bull* **61**: 81-96.  
4
- 5 [2] Smalley JW, Charalabous P, Birss AJ & Hart CA (2001) Detection of heme-  
6 binding proteins in epidemic strains of *Burkholderia cepacia*. *Clin Diag Lab Immunol* **8**:  
7 509-514.
- 8 [3] Smalley JW, Charalabous P, Hart CA & Silver J (2003). Transmissible  
9 *Burkholderia cepacia* genomovar IIIa strains bind and convert monomeric iron(III)  
10 protoporphyrin IX into the  $\mu$ -oxo oligomeric form. *Microbiology* **149**: 843-853.  
11
- 12 [4] Katsuwon J & Anderson AJ (1992) Characterisation of catalase activities in a  
13 root-colonizing isolate of *Pseudomonas putida*. *Can J Microbiol* **38**: 1026-1032.  
14
- 15 [5] Josephy PD, Eling T & Mason RP (1982) The horseradish peroxidase-catalyzed  
16 oxidation of 3,5,3', 5'-tetramethylbenzidine: Free radical and charge-transfer  
17 intermediates. *J Biol Chem* **257**: 3669-3675.  
18
- 19 [6] Beers RF & Sizer IW (1952) A spectrophotometric assay for measuring the  
20 breakdown of hydrogen peroxide by catalase. *J Biol Chem* **196**: 133-140.  
21
- 22 [7] Southern EM (1975) Detection of specific sequences among DNA fragments  
23 separated by gel electrophoresis. *J Mol Biol* **98**: 503-517.  
24
- 25 [8] Lefebvre MD & Valvano MA (2001) *In vitro* resistance of *Burkholderia cepacia*  
26 complex isolates to reactive oxygen species in relation to catalase and superoxide  
27 dismutase production. *Microbiology* **147**: 97-109.

28

- 1 [9] Donald LJ, Krokhin OV, Duckworth HW, Wiseman B, Deemagarn T, Singh R,  
2 Switala J, Carpena X, Fita I & Loewen P (2003) Characterization of the catalase-  
3 peroxidase KatG from *Burkholderia pseudomallei* by mass spectrometry. *J Biol*  
4 *Chem* **278**: 35687-35692.
- 5
- 6 [10] Sonnenberg MG & Belisle JT (1997) Definition of *Mycobacterium tuberculosis*  
7 culture filtrate proteins by two-dimensional polyacrylamide gel electrophoresis, N-  
8 terminal amino acid sequencing and electrospray mass spectrometry. *Infect Immun*  
9 **65**: 4515-4524.
- 10
- 11 [11] Cannac-Caffrey V, Hudry-Clergeon G, Petillot Y, Gagnon J, Zaccari G & Franzetti  
12 B (1998) The protein sequence of an archaeal catalase-peroxidase. *Biochimie* **80**:  
13 1003-1011.
- 14
- 15 [12] Lefebvre MD, Flannagan R S & Valvano MA (2005) A minor catalase/peroxidase  
16 from *Burkholderia cenocepacia* is required for normal aconitase activity. *Microbiology*  
17 **151**: 1975–1985.
- 18
- 19 [13] Zamocky M, Janecek S & Koller F (2000) Common phylogeny of catalase-  
20 peroxidases and ascorbate peroxidases. *Gene* **256**: 169-182.
- 21
- 22 [14] Zamocky M, Regelsberg G, Jakopitsch C & Obinger C (2001) The molecular  
23 peculiarities of catalase-peroxidases. *FEBS Lett* **492**: 177-182.
- 24
- 25 [15] Carpena X, Switala J, Loprasert S, Mongkolsuk S, Fita I & Loewen PC (2002)  
26 Crystallisation and preliminary X-ray analysis of the catalase-peroxidase KatG from  
27 *Burkholderia pseudomallei*. *Acta Crystallogr D Biol Crystallogr* **58**: 2184-2186.
- 28
- 29 [16] Carpena X, Loprasert S, Mongkolsuk S, Switala J, Loewen PC & Fita I (2003)  
30 Catalase-peroxidase KatG of *Burkholderia pseudomallei* at 1.7Å resolution. *J Mol*  
31 *Biol* **327**: 475-489.
- 32

- 1 [17] Loprasert S, Whangsuk W, Sallabhan R & Mongkolsuk S (2003)  
2 Regulation of the katG-dpsA operon and the importance of KatG in survival of  
3 *Burkholderia pseudomallei* exposed to oxidative stress. *FEBS Lett* **542**: 17-21.
- 4 [18] Jayaraman S, Song Y & Verkman AS (2001) Airway surface liquid pH in well-  
5 differentiated airway epithelial cell cultures and mouse trachea. *Am J Cell Physiol*  
6 **281**: C1504-C1511.
- 7 [19] Sajjan US, Corey M, Karmali MA & Forstner JF (1992) Binding of *Pseudomonas*  
8 *cepacia* to normal intestinal and respiratory mucin from patients with cystic fibrosis. *J*  
9 *Clin Invest* **89**: 648-656.
- 10 [20] Chace KV, Leahy DS, Martin R, Carubelli R, Flux M & Sachdev P (1983)  
11 Respiratory mucous secretions in patients with cystic fibrosis: relationship between  
12 levels of highly sulfated mucin component and severity of the disease. *Clin Chim*  
13 *Acta* **132**: 143-155.
- 14  
15 [21] Chace KV, Flux M & Sachdev P (1985). Comparisons of physicochemical  
16 properties of purified mucus glycoproteins isolated from respiratory secretions of  
17 cystic fibrosis and asthmatic patients. *Biochem* **24**: 7334-7341.
- 18  
19 [22] Cheng PW, Boat TF, Cranfill K, Yankaskas JR & Boucher RC (1989) Increased  
20 sulfation of glycoconjugates by cultures of nasal epithelial cells from patients with  
21 cystic fibrosis. *J Clin Invest* **84**: 68-72.
- 22 [23] Coakley RD & Boucher RC (2001) Regulation and functional significance of  
23 airway surface liquid pH. *J Pancreas (Online)* **2**: 294-300.
- 24 [24] Yoon SS, Coakley R, Lau GW, Lyman SV, Gaston B, Karabulut AC, Hennigan  
25 RF, Hwang SH, Buettner G, Schurr MJ, Mortensen JE, Burns JL, Speert D, Boucher  
26 RC & Hassett DJ. (2006) Anaerobic killing of mucoid *Pseudomonas aeruginosa* by  
27 acidified nitrite derivatives under cystic fibrosis airway conditions. *J Clin Invest* **116**:  
28 436-446.
- 29

- 1 [25] Bodem CR, Lampton L.M, Miller DP, Tarka EF & Everett ED (1983)  
2 Endobrochial pH relevance to aminoglycoside activity in gram-negative bacillary  
3 pneumonia. *Am Rev Respir Dis* **127**: 39-41.  
4
- 5 [26] Charalabous, P. (2004). Catalase-peroxidases of *Burkholderia cenocepacia*.  
6 PhD Thesis, The University of Liverpool, Liverpool, UK.  
7
- 8 [27] Marcinkeviciene JA, Magliozzo RS & Blanchard JS (1995) Purification and  
9 characterisation of the *Mycobacterium smegmatis* catalase-peroxidase involved in  
10 isoniazid activation. *J Bacteriol* **270**: 22290-22295.
- 11 [28] Long S & Salin M L (2001) Molecular cloning, sequencing analysis and  
12 expression of the catalase-peroxidase gene from *Halobacterium salinarum*. *DNA*  
13 *Sequencing* **12**: 39-51.  
14
- 15 [29] Heym B, Alzari P M, Honore N & Cole ST (1995) Missense mutations in the  
16 catalase-peroxidase gene, *katG*, are associated with isoniazid resistance in  
17 *Mycobacterium tuberculosis*. *Mol Microbiol* **15**: 235-245.  
18
- 19 [30] Menendez MC, Ainsa JA, Martin C & Garcia MJ (1997) *katGI* and *katGII*  
20 encode two different catalases-peroxidases in *Mycobacterium fortuitum*. *J Bacteriol*  
21 **179**: 6880-6886.  
22
- 23 [31] Morris SL, Nair J & Rouse DA (1992) The catalase-peroxidase of  
24 *Mycobacterium intracellulare*: nucleotide sequence analysis and expression in  
25 *Escherichia coli*. *J Gen Microbiol* **138**: 2363-2370.  
26
- 27 [32] Heym B, Zhang Y, Poulet S, Young D & Cole ST (1993) Characterization of the  
28 *katG* gene encoding a catalase-peroxidase required for the isoniazid susceptibility of  
29 *Mycobacterium tuberculosis*. *J Bacteriol* **175**: 4255-4259.  
30
- 31 [33] Triggs-Raine BL, Doble BW, Mulvey MR, Sorby PA & Loewen PC (1988)

1 Nucleotide sequence of *katG*, encoding catalase HPI of *Escherichia coli*. *J Bacteriol*  
2 **170**: 4415-4419.

3

4 [34] Brunder W, Schmidt H & Karch H (1996) KatP, a novel catalase-peroxidase  
5 encoded by the large plasmid of enterohaemorrhagic *Escherichia coli* O157:H7.  
6 *Microbiology* **142**: 3305-3315.

7

8 [35] Bandyopadhyay P & Steinman HM (1998) *Legionella pneumophila* catalase-  
9 peroxidases: cloning of the *katB* gene and studies of KatB function. *J Bacteriol* **180**:  
10 5369-5374.

11

12 [36] Loewen PC & Stauffer GV (1990) Nucleotide sequence of *katG* of *Salmonella*  
13 *typhimurium* LT2 and characterization of its product, hydroperoxidase I. *Mol Gen*  
14 *Genetics* **224**: 147-151.

15

16 [37] Zou P, Borovok I, Ortiz de Orue Lucana D, Muller D & Schrempf H (1999) The  
17 mycelium-associated *Streptomyces reticuli* catalase-peroxidase, its gene and  
18 regulation by FurS. *Microbiology* **145**: 549-559.

19

20 [38] Garcia E, Nedialkov Y A, Elliott J, Motin V L & Brubaker R R (1999) Molecular  
21 characterization of *KatY* (antigen 5), a thermoregulated chromosomally encoded  
22 catalase-peroxidase of *Yersinia pestis*. *J Bacteriol* **181**: 3114-3122.

23

24 [39] Loprasert S, Sallabhan R, Whangsuk W & Mongkolsuk S (2002) The  
25 *Burkholderia pseudomallei oxyR* gene: expression analysis and mutant  
26 characterization. *Gene* **296**: 161-169.

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28

1 Table 1

2

3 Amino acid homology analysis of the *B. cenocepacia* catalase-peroxidase KatG  
 4 (GenBank accession DQ112341) with bacterial and archaeal catalase-peroxidases  
 5 using paired alignment comparisons performed in BLASTP. Accession numbers are  
 6 from GenBank (gb) or Swiss-Prot (sp).

Organism	Accession no.	Reference	% Identity	% Similarity
<i>Haloarcula marismortui</i>	gbY16851	Cannac-Caffrey <i>et al.</i> [11]	56.8	69.4
<i>Halobacterium salinarum</i>	gbAF069761	Long & Salin [28]	56.4	68.9
<i>Mycobacterium bovis</i>	spP46817	Heym <i>et al.</i> [29]	60.0	71.8
<i>Mycobacterium fortuitum</i>	gbY07865	Menendez <i>et al.</i> [30]	58.1	65.9
<i>Mycobacterium intracellulare</i>	spQ04657	Morris <i>et al.</i> [31]	58.6	69.3
<i>Mycobacterium tuberculosis</i>	spQ08129	Heym <i>et al.</i> [32]	59.9	71.6
<i>Escherichia coli</i>	spP13029	Triggs-Raine <i>et al.</i> [33]	59.9	71.8
<i>E. coli</i> (0157: H7)	gbX89017	Brunder <i>et al.</i> [34]	55.2	66.2
<i>Legionella pneumophila</i>	gbAF078110	Bandyopadhyay & Steinman [35]	57.1	69.3
<i>Salmonella typhimurium</i>	spP17750	Loewen & Stauffer [36]	60.6	71.9
<i>Streptomyces reticuli</i>	gbY14317	Zou <i>et al.</i> [37]	64.0	74.0
<i>Yersinia pestis</i>	gbAF135170	Garcia <i>et al.</i> [38]	55.6	67.0
<i>Burkholderia pseudomallei</i>	gbAAK72466	Loprasert <i>et al.</i> [37]	70.7	79.0
<i>Burkholderia cenocepacia</i>	gbDQ114424	Smalley <i>et al.</i> (2005)	71.0	79.0
<i>Burkholderia cenocepacia</i>	gbAF317697	Lefebvre <i>et al.</i> [12]	94.0	94.0

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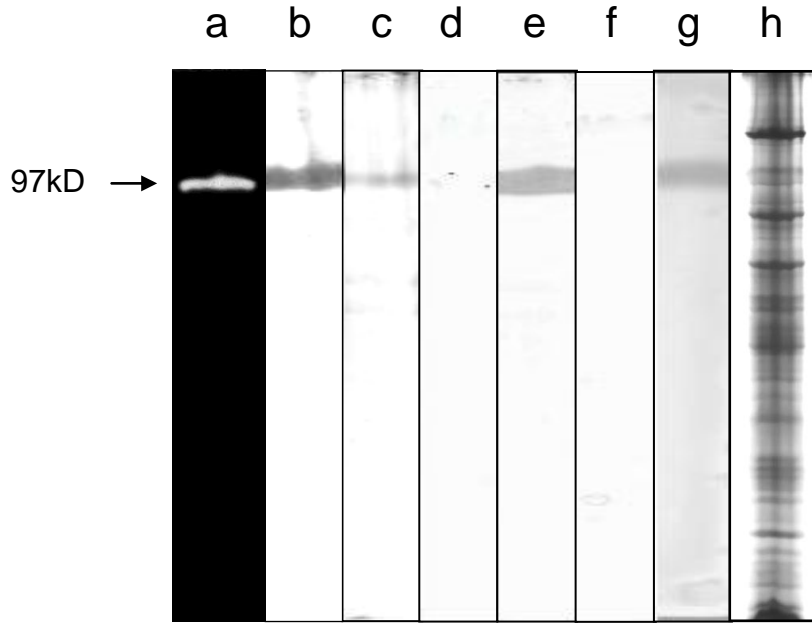
Table 2

Masses of tryptic peptides derived from the recombinant protein. Analysis was performed with one missed cleavage allowed and deamidation as the only variable modification. A mass tolerance of 1.2Da was allowed for MS and of 0.6Da for MS/MS analysis. R1, R2, and R3 refer to the arrowed bands on Figure 5; m/z, mass/charge; Mr(calc.), calculated relative molecular masses of the sequenced tryptic peptides; (D), deamidation.

Protein	Position	m/z	Mr(calc.)	Sequence	
<b>R1</b>	29 - 38	564.3347	1126.5982	R.LDLLSQHSSK.T	
	39 - 61	1277.6446	2553.2023	K.TDPLDPGFNYAEAFNSLDLDALR.K	
	273 - 299	901.4813	2701.3095	K.THGAGPADNVGLEPEAAGLEQQGLGWK.N	
	333 - 343	700.3453	1398.6819	K.NLFGYEWELTK.S	
	Overall	383 - 389	449.2351	896.4279	R.FDPVYEK.I
	Mascot	520 - 532	681.8758	1361.6575	R.IQGEFNSTQPGGK.K
	Score =	534 - 553	955.5686	1909.0883	K.ISLADLIVLAGGAGIEQAAK.R
	875	555 - 567	661.3781	1320.6938	R.AGHDVVVPFAPGR.M
		598 - 609	643.9088	1285.7281	K.FAVPAEALLIDK.A
		639 - 655	1005.0381	2007.9690	K.HGVFTDQPETLTVDFFR.N
<b>R2</b>	690 - 701	645.8812	1288.7139	R.VDLVFGSNAVLR.A	
	702 - 715	705.8608	1409.6674	R.ALSEVYASADGEAK.F	
	29 - 38	564.3241	1126.5982	R.LDLLSQHSSK.T	
	39 - 61	1277.6765	2553.2023	K.TDPLDPGFNYAEAFNSLDLDALR.K	
	113 - 127	851.9246	1701.8362	R.FAPLNSWPDNVSLDK.A	
	Overall	273 - 299	901.4947	2702.2935	K.THGAGPADNVGLEPEAAGLEQQGLGWK.N (D)
	Mascot	333 - 343	700.3926	1398.6819	K.NLFGYEWELTK.S
	Score =	383 - 389	449.2540	896.4279	R.FDPVYEK.I
	871	534 - 553	955.5824	1909.0883	K.ISLADLIVLAGGAGIEQAAK.R
		555 - 567	661.3666	1320.6938	R.AGHDVVVPFAPGR.M
<b>R3</b>	598 - 609	643.8862	1285.7281	K.FAVPAEALLIDK.A	
	639 - 655	1005.0239	2007.9690	K.HGVFTDQPETLTVDFFR.N	
	690 - 701	645.8698	1288.7139	R.VDLVFGSNAVLR.A	
	702 - 715	705.8490	1409.6674	R.ALSEVYASADGEAK.F	
	29 - 38	564.3127	1126.5982	R.LDLLSQHSSK.T	
	39 - 61	852.4210	2553.2023	K.TDPLDPGFNYAEAFNSLDLDALR.K	
	253 - 272	1032.0497	2060.9659	R.MAMNDEETVALIAGGHAFGK.T	
	Overall	273 - 299	901.4964	2701.3095	K.THGAGPADNVGLEPEAAGLEQQGLGWK.N
	Mascot	355 - 368	757.3829	1512.7321	K.NAEPTIPHAHDPSK.K
	Score =	370 - 382	737.4313	1472.8272	K.LLPTMLTTDLSLR.F
964	393 - 407	876.8862	1751.7725	R.HFMDNPDVFADAFAR.A	
	520 - 532	681.8408	1361.6575	R.IQGEFNSTQPGGK.K	
	534 - 553	955.5983	1909.0883	K.ISLADLIVLAGGAGIEQAAK.R	
	555 - 567	661.3550	1320.6938	R.AGHDVVVPFAPGR.M	
	598 - 609	643.8859	1285.7281	K.FAVPAEALLIDK.A	
	610 - 628	977.5799	1953.1081	K.AQLLTLTAPQMTALVGGLR.V	
	639 - 655	1004.9840	2008.9530	K.HGVFTDQPETLTVDFFR.N (D)	
	690 - 701	645.3592	1288.7139	R.VDLVFGSNAVLR.A	
	702 - 715	705.8491	1409.6674	R.ALSEVYASADGEAK.F	

Fig 1

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Fig 2

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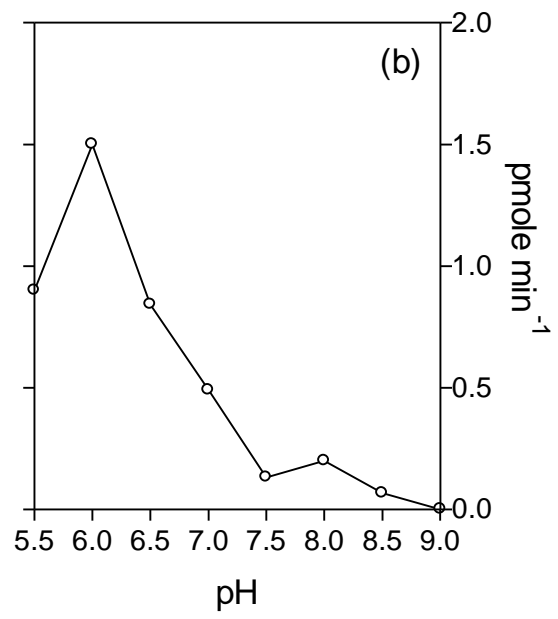
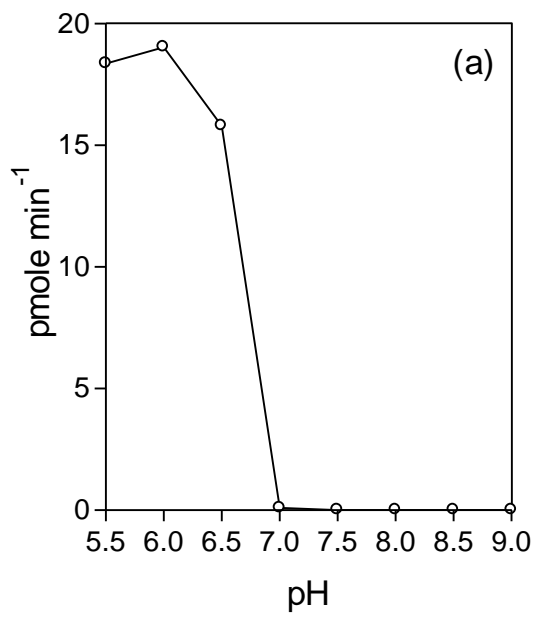
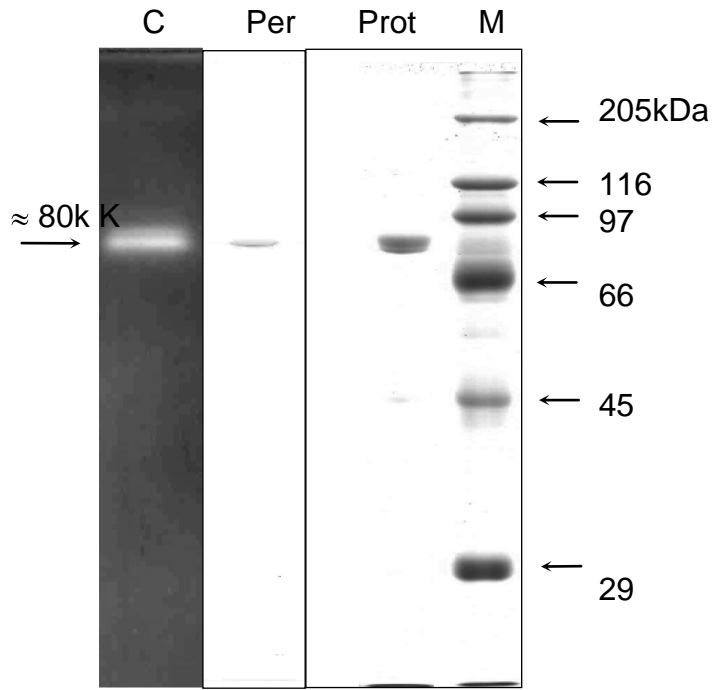
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Fig 4

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1 Fig 5

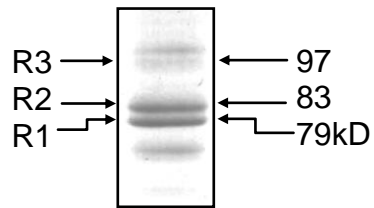
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1 Figure legends

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3 Figure 1: Catalase and peroxidase specificity of the 97kDa protein of *B. cenocepacia*  
4 strain J2315 as shown after SDS-PAGE under non-reducing conditions. Catalase  
5 activity was demonstrated using the potassium ferricyanide/ ferric chloride-H<sub>2</sub>O<sub>2</sub>  
6 method (track a), whilst peroxidase activity was assessed against (b) TMB, (c) *o*-  
7 dianisidine, (d) 2-methoxyphenol, (e) ABTS, (f) 4-chloronaphthol, and (g) pyrogallol.  
8 Track h, Coomassie Blue counterstaining for protein following the peroxidase  
9 reaction. See Methods for details.

10

11 Figure 2: Effect of pH on the activity of the bi-functional catalase-peroxidase as  
12 shown after SDS-PAGE under non-reducing conditions. Peroxidase activity (a) was  
13 revealed using tetramethylbenzidine-H<sub>2</sub>O<sub>2</sub> whilst catalase was assayed using the  
14 K<sub>3</sub>Fe(CN)<sub>6</sub>/FeCl<sub>3</sub>-H<sub>2</sub>O<sub>2</sub> staining method (b). Gel loadings were 25µg protein per  
15 track.

16

17 Figure 3: pH activity profile of catalase and peroxidase activities of suspensions of  
18 whole cells of *B. cenocepacia* strain J2315 grown on M9 Minimal Salts Medium agar.  
19 Peroxidase was measured using tetramethylbenzidine as substrate (shown as  
20  $\Delta A_{645\text{nm}}$ ). Catalase activity was assayed by UV absorbance (shown as  $\Delta A_{240\text{nm}}$ ). The  
21 reactions were carried out at 20°C.

22

23 Fig 4: SDS-PAGE on 10% polyacrylamide gels of BL21Star *E. coli* cells expressing  
24 the catalase-peroxidase, after growth in the presence of 0.1M IPTG. C, catalase;  
25 Per, peroxidase; Prot, protein staining; M, molecular weight markers. The in-gel  
26 catalase staining was performed at pH 6.0 in 0.5M sodium acetate.

27

28 Fig 5: SDS-PAGE of the recombinant *B. cenocepacia* KatG catalase-peroxidase  
29 enzyme on a 7% polyacrylamide gel. Recombinant protein bands R1, R2 and R3  
30 gave mass matches to KatG after trypsinisation and MS/MS analysis (see Table 2).  
31 The gel was stained with Coomassie Blue.

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