1	The moonlighting peroxiredoxin-glutaredoxin in Neisseria meningitidis
2	binds plasminogen via a C-terminal lysine residue and contributes to
3	survival in a whole blood model
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17 Abstract

18

cross the blood-brain barrier resulting in life-threatening sepsis and meningitis. Meningococci 19 20 express a cytoplasmic peroxiredoxin-glutaredoxin (Prx5-Grx) hybrid protein that has also been identified on the bacterial surface. Here, recombinant Prx5-Grx was confirmed as a 21 plasminogen (Plg)-binding protein, in an interaction which could be inhibited by the lysine 22 analogue ϵ -aminocapronic acid. rPrx5-Grx derivatives bearing a substituted C-terminal lysine 23 residue (rPrx5-Grx^{K244A}), but not the active site cysteine residue (rPrx5-Grx^{C185A}) or the sub-24 terminal rPrx5-Grx^{K230A} lysine residue, exhibited significantly reduced Plg-binding. The 25 absence of Prx5-Grx did not significantly reduce the ability of whole meningococcal cells to 26 bind Plg, but under hydrogen peroxide-mediated oxidative stress, the N. meningitidis $\Delta pxn5$ -27 28 grx mutant survived significantly better than the wild-type or complemented strains. 29 Significantly, using human whole blood as a model of meningococcal bacteremia, it was found that the *N. meningitidis* $\Delta pxn5$ -grx mutant had a survival defect compared with the 30 31 parental or complemented strain, confirming an important role for Prx5-Grx in meningococcal pathogenesis. 32 33 Keywords: Neisseria meningitidis; protein moonlighting; peroxiredoxin; pathogenesis; whole 34 blood model; plasminogen 35 36 Highlights: 37 38 Meningococcal Prx5-Grx interacts with plasminogen via a C-terminal lysine residue • Loss of Prx5-Grx does not reduce the ability of meningococci to bind Plg 39 • 40 Loss of Prx5-Grx enhances meningococcal survival under oxidative stress ٠ Loss of Prx5-Grx results in a survival defect in human whole blood 41

Neisseria meningitidis is a human-restricted bacterium that can invade the bloodstream and

42 **1. Introduction**

Neisseria meningitidis is an important cause of sepsis and meningitis in susceptible 43 individuals, with the majority of cases occurring in children and young adults. The organism 44 frequently colonizes the nasopharynx asymptomatically and is transmitted from person-to-45 person by close contact [1]. Rarely, however, hypervirulent strains may invade the mucosal 46 epithelial barrier to reach subcutaneous tissues and subsequently enter the bloodstream [2]. 47 48 The ability of the meningococcus to survive in subcutaneous tissues and blood is therefore a prerequisite for invasive meningococcal disease. Various meningococcal components are 49 50 known to be essential for, or enhance, survival in non-immune blood, including capsular polysaccharide [3, 4], lipooligosaccharide (LOS) [5], the factor H-binding proteins (fHbp and 51 NspA) [5-8] and NalP [8, 9]. Various detoxifying enzymes are also used by N. meningitidis to 52 53 protect against reactive oxygen and nitrogen species produced by polymorphonuclear (PMN) 54 cells and monocytes/macrophages [10, 11]. Up-regulation of catalase (Kat), superoxide dismutase (Sod) and nitrite reductase (AniA) have been demonstrated following incubation in 55 56 human blood [8]. SodC has previously been shown to protect *N. meningitidis* from phagocytosis [12] and AniA protects N. gonorrhoeae in human serum [13], however Kat was 57 shown not to be required for meningococcal survival in human blood [8]. 58 Peroxiredoxins (Prxs) are a large family of cysteine-based peroxidases, present in all 59 kingdoms of life, that catalyse the reaction ROOH+2e⁻ \rightarrow ROH+H₂O and reduce hydrogen 60 61 peroxide (H₂O₂), peroxynitrite and a wide range of organic hydroperoxide compounds [14-16]. Prxs are ubiquitous and abundant proteins that are important for antioxidant defence and 62 the regulation of cell signalling pathways [17, 18]. Prxs can be divided into six subfamilies 63 64 (AhpC-Prx1, BCP-PrxQ, Tpx, Prx5, Prx6 and AhpE) with distinct amino acid sequences, but all contain an absolutely conserved active site cysteine in the motif P-XXX-T(S)-XX-C, 65 which reacts with H₂O₂ to form a cysteine sulfenic acid [19]. Prxs are described as either 1-66

67	Cys Prx or 2-Cys Prx, depending on whether a second cysteine residue participates in the
68	enzymatic reaction [19]. The 2-Cys Prxs are further subdivided into 'typical' and 'atypical'
69	forms, depending on the location of the second cysteine residue; in typical forms the second
70	cysteine is found in the C-terminus of the partner subunit, whereas in atypical enzymes it is
71	located at other positions [19]. Glutaredoxins (Grxs) are small redox enzymes that use
72	glutathione (GSH) as a cofactor [20]. Grxs possess an active site disulfide bond, which exists
73	in either a reduced or an oxidized form. Grxs act as antioxidants by reducing
74	dehydroascorbate, peroxiredoxins and methionine sulfoxide reductase. Some bacterial species
75	contain tetrameric Prx5-Grx hybrid enzymes in which the C-terminal Grx domain acts as an
76	electron donor for the N-terminal Prx domain [21].
77	Prx enzymes are increasingly being recognized as multifunctional or moonlighting
78	proteins, which are involved in various aspects of bacterial survival and pathogenesis. In
79	Helicobacter pylori, a BCP-PrxQ subfamily Prx was found to contribute to resistance to
80	oxidative stress and host colonization [22] and an AhpC-Prx1 subfamily Prx was shown to
81	oligomerise and acquire protein chaperone activity under oxidative stress conditions [23]. In
82	Haemophilus influenzae, a Prx-Grx hybrid enzyme was found to be expressed preferentially
83	during growth in biofilms [24]. A mutant strain was more susceptible to organic
84	hydroperoxides [25] and showed persistence defects in a chinchilla model of otitis media and
85	a murine model of chronic obstructive pulmonary disease [26]. In Streptococcus agalactiae, a
86	similar AhpC-Prx1 enzyme was shown to bind haem and participate in its intracellular
87	availability [27]. In Coxiella burnetii, a BCP-PrxQ Prx was shown to bind DNA and protect
88	the organism from oxidative stress during the exponential phase of growth [28]. A Prx6
89	subfamily protein, LsfA, in Pseudomonas aeruginosa, was found to reduce the oxidative
90	burst in macrophages and to be important for virulence in a murine model of pneumonia [29].
91	In Vibrio vulnificus, Prx3, a Grx3/GSH-dependent 1-Cys Prx, was shown to be required for

92 optimal growth in medium containing peroxides and the Prx3 mutant strain was less virulent in mice [30]. A 1-Cys Prx6 enzyme from Thermococcus kodakarensis, a thermophilic 93 anaerobic archaeon, was found to act as a molecular chaperone that blocked oxidative stress-94 95 mediated aggregation of proteins and DNA [31]. More recently, an AhpC-Prx1 subfamily Prx of F. tularensis LVS was shown to confer resistance against a wide range of reactive oxygen 96 and nitrogen species, and to serve as a virulence factor. In the highly virulent strain, F. 97 98 *tularensis* SchuS4, the enzyme also enhanced intra-macrophage survival [32]. N. meningitidis contains a Prx-Grx hybrid atypical 2-Cys enzyme in the Prx5 99 100 subfamily and a BCP-PrxQ subfamily Prx (designated NMB0946 and NMB0750, respectively, in strain MC58) [19, 33]. The Prx5-Grx hybrid enzyme was previously found to 101 be a putative moonlighting protein present on the meningococcal cell surface where it 102 103 contributes to plasminogen (plg)-binding [34] and in N. gonorrhoeae, the Prx5-Grx 104 orthologue was found to play a role in biofilm formation [35]. In this study, we examine the properties and role of Prx5-Grx in meningococcal pathogenesis. We confirm that Prx5-Grx 105 106 moonlights on the bacterial surface and show that a single lysine residue in the penultimate C-terminal position in Prx5-Grx is required for optimal interactions with human plasminogen. 107 Importantly, we also demonstrate that Prx5-Grx is required for meningococcal survival in 108 human non-immune whole blood. 109

110

111 **2. Experimental procedures**

112 2.1. Bacterial strains and culture conditions

Escherichia coli JM109 (Promega) was used as a host strain for the expression of 6 ×
histidine-tagged rPrx5-Grx and for mutagenic plasmid construction. *E. coli* XL10-Gold ultracompetent cells (Agilent Technologies) were used for site-directed mutagenesis of *rpsL*. NEB
5-alpha competent cells were used for cloning of Gibson assembly reactions. All *E. coli*

strains were grown at 37°C in Lysogeny Broth (LB) or on LB agar supplemented, where appropriate, with ampicillin (100 μ g ml⁻¹), kanamycin (30 μ g ml⁻¹) or erythromycin (200 μ g ml⁻¹). Strains of *Neisseria meningitidis* (Table S1) were grown at 37°C in air plus 5% CO₂ on Columbia agar with chocolated horse blood (Thermo Fisher Scientific), Brain Heart Infusion (BHI) agar or BHI broth supplemented with 1% Vitox (Thermo Fisher Scientific) and kanamycin (50 μ g ml⁻¹), streptomycin (100 μ g ml⁻¹) or erythromycin (5 μ g ml⁻¹), where appropriate.

124

125 2.2. Construction of plasmids encoding recombinant Prx5-Grx

The prx5-grx gene was amplified from N. meningitidis MC58 using oligonucleotides 126 NMB0946F1 and NMB0946R1 (Table S2) using Phusion High-Fidelity DNA polymerase 127 128 (New England Biolabs). After digestion with BamHI, the PCR product was ligated into BamHI-digested pQE30 to yield pMAJ2 (Table S3). The Prx5-Grx lysine residues (²³⁰K, ²⁴⁴K 129 and ${}^{230}\text{K}/{}^{244}\text{K}$) and the glutaredoxin active-site cysteine residue (${}^{185}\text{C}$) were replaced by 130 alanine residues using the Q5 Site-Directed Mutagenesis Kit (New England BioLabs) 131 following the manufacturer's instructions, yielding pMAJ2-230KA, pMAJ2-244KA, pMAJ2-132 230244KKAA and pMAJ2-185CA, respectively (Table S3). Oligonucleotides used are given 133 in Table S2. 134

135

136 2.3. Recombinant Prx5-Grx expression and purification

E. coli JM109 strains were grown to log phase, induced with 0.5 mM isopropyl β-D-1thiogalactopyranoside (IPTG) for 3 h, and harvested by centrifugation (4,200 × g for 10 min).
Recombinant 6 × histidine-tagged proteins were then affinity-purified under native
conditions. Briefly, *E. coli* cell pellets were resuspended in 30 ml lysis/wash buffer (50 mM
NaH₂PO₄, 300 mM NaCl, 40 mM imidazole, pH 7.4) followed by sonication using an MSE

142 SoniPrep 150 sonicator for 8 cycles (30s on, 30s off) on ice. The cell lysate was centrifuged $(4,500 \times g \text{ for } 15 \text{ min})$ and the cleared lysate was loaded onto a HisTrap FF column (GE 143 Healthcare Lifesciences) pre-packed with Ni Sepharose 6 Fast Flow (GE Healthcare 144 Lifesciences) connected to a ÄKTAprime plus liquid chromatography system (GE Healthcare 145 Lifesciences), equilibrated with 10 column volumes of lysis/wash buffer. Proteins were 146 eluted by step elution using elution buffer (50 mM NaH₂PO₄, 300 mM NaCl, 300 mM 147 imidazole, pH 7.4). A HiTrap column pre-packed with five millilitres of Sephadex G-25 148 Superfine (GE Healthcare Lifesciences) equilibrated with 5 column volumes of phosphate 149 150 buffered saline (PBS) was used for buffer exchange. Glycerol at a final concentration of 10% was added to inhibit aggregation. A pre-packed Superdex column (GE Healthcare 151 Lifesciences) mounted on an ÄKTAprime plus liquid chromatography system was then used 152 153 to carry out gel filtration, following the manufacturer's instructions. After equilibration with 154 150 ml PBS containing 10% glycerol, the IMAC-purified protein sample was injected into the column. Fractions contained the purified protein were collected and concentrated using 155 Vivaspin sample concentrators (Sartorius; 10,000 MWCO). 156

157

158 2.4. Production of a rabbit antiserum against rPrx5-Grx

Rabbit anti-Prx5-Grx antibodies were raised in a New Zealand White female rabbit immunized subcutaneously four times at 2-week intervals with 100 µg of rPrx5-Grx protein emulsified in Freud's complete (first immunization only) or incomplete adjuvant. After three injections, the animal was test bled, boosted once more and sacrificed 7 days later.

163

164 2.5. SDS-PAGE and immunoblotting

165 Proteins were electrophoretically separated using 10% polyacrylamide gels (Mini-

166 Protean III; Bio-Rad) and stained using SimplyBlueTM SafeStain (Thermo Fisher Scientific)

167 or transferred to nitrocellulose membranes using a Trans-Blot SD semidry transfer cell (Bio-Rad) according to the manufacturer's recommendations. Membranes were probed with mouse 168 anti-pentahistidine antibody (Qiagen) or rabbit anti-Prx5-Grx primary antibody (RaPrx5-Grx) 169 170 diluted 1:10,000 or 1:1,000, respectively, in blocking buffer (5% [w/v] non-fat dry milk, 0.1% [v/v] Tween 20 in $1 \times PBS$) and incubated for 2 h. After washing in PBS with 0.1% 171 Tween 20 (PBST), membranes were incubated for 2 h with 1:10,000-diluted goat anti-mouse 172 173 (or anti-rabbit) IgG-alkaline phosphatase conjugate (Sigma). After washing with PBST, blots were developed using BCIP/NBT-Blue liquid substrate (Sigma). 174

175

176 *2.6. ELISA*

Microplate wells (Nunc 96-well plates, PolySorp) were coated with 20 pmol glu-plg 177 178 (human plasma; Calbiochem), laminin (human placenta; Merck Millipore), fibronectin (human plasma; Sigma), collagen I (human placenta; Corning) or 1% BSA in sodium 179 carbonate buffer (142 mM NaHCO₃, 8 mM Na₂SO₃, pH 9.0) and incubated overnight at 4°C. 180 After washing with PBS-Tween 20 (0.05%; PBST), wells were blocked with 1% BSA in PBS 181 for 1 h. After removal of the blocking solution, 50 pmol rPrx5-Grx in 1% BSA/PBS was 182 added and incubated for 1 h. Following vigorous washing with PBST, RaPrx5-Grx (diluted 183 1:2,000 in 1% BSA/PBS) was added for incubation for 1 h. Plates were again vigorously 184 washed, before the addition of goat anti-rabbit IgG-alkaline phosphatase conjugate (Sigma; 185 186 diluted 1:20,000 in 1% BSA/PBS), was added and incubated for 1 h. Plates were again vigorously washed, and the color was developed by adding phosphatase substrate (Sigma), 187 dissolved in buffer containing 0.1 M glycine, 1 mM ZnCl₂ and 1 mM MgCl₂, pH 10.4, into 188 189 each well and absorbance at 405nm was measured using a Biotek EL800 spectrophotometer. Alternatively, binding assays were undertaken as above, but using rPrx5-Grx (5 pmol) or 1% 190 BSA as the immobilized ligand to capture glu-plg (5 pmol). Bound glu-plg was detected 191

using goat anti-human plg (1:10,000; Rockland Immunochemical) and donkey anti-goat IgGalkaline phosphatase conjugate (1:5,000; Promega). The lysine analogue, ε-aminocaproic
acid (EACA; 50 mM) (Sigma) was utilized for inhibition studies.

195

196 2.7. Generation of MC58 $rpsL^{-}$ and MC58 $Kan^{R} rpsL^{+} rpsL^{-}$

A 2.5-kb fragment consisting of *rpsL* and flanking DNA was amplified from 197 N. meningitidis MC58 using oligonucleotides rpsL-FlankF and rpsL-FlankR (Table S2). The 198 amplified gene was TA-cloned into pCR4-TOPO to generate plasmid pCF4-TOPOArpsL, 199 which was subjected to site-directed mutagenesis, designed to replace ⁴³K with T, using the 200 primers rpsLSD-F and rpsLSD-R (Table S2) and the QuikChange Lightning Site-Directed 201 Mutagenesis Kit (Agilent Technologies) according to the manufacturer's protocol. Following 202 203 mutagenesis, the reaction was incubated with DpnI to digest non-mutated parental DNA prior 204 to transformation of E. coli XL10-Gold ultra-competent cells. A plasmid extracted from a resulting clone was sequenced and confirmed to contain the desired mutation and designated 205 206 pCR4-TOPOArpsLT and used to transform N. meningitidis MC58. Genomic DNA from a resulting clone was sequenced, the desired mutation confirmed, and the resulting mutant 207 208 designated MC58 rpsL⁻ (Table S1).

An MC58 derivative containing a cassette consisting of a kanamycin resistance gene 209 and a wild-type rpsL gene was generated using a protocol based on that described by Gibson 210 211 [36]. Primer pairs Upstream343_F and Upstream343_R; Downstream343_F and Downstream343 R; rspL F and rspL R (Table S2) were designed using the web tool 212 NEBuilder to amplify fragments with appropriate overlaps required for assembly of 213 214 fragments corresponding to regions upstream and downstream of the gene NMB0343 and the rpsL gene from chromosomal DNA of N. meningitidis strain MC58. The aphA-3 gene of 215 plasmid pJMK30 [37] was amplified using the primers Kan cassette_F and Kan cassette_R 216

and all four fragments, together with pUC19 plasmid DNA linearized with EcoRI and BamHI, was performed using 0.2-0.25 pmol of each DNA fragment, 10 μ l of 2 × Gibson assembly master mix (NEB) and dH₂O to 20 μ l. After incubation at 50°C, the assembly reaction was used to transform NEB 5-alpha competent *E. coli* cells (NEB). The resulting plasmid was subsequently used to transform MC58 *rpsL*⁻ by selection for kanamycin resistance and screening for streptomycin sensitivity. The resulting strain was designated MC58 *Kan^R rpsL*⁺ *rpsL*⁻ (Table S1).

224

225 2.8. Construction of a markerless Δprx5-grx mutant

A 2.7-kb fragment consisting of *prx5-grx* and flanking DNA was amplified using 226 227 oligonucleotides UprxDF1 and UprxDR1 (Table S2) from N. meningitidis MC58. The amplified DNA was TA-cloned into pGEM-T Easy to generate pGUD1 (Table S3). This was 228 then subjected to inverse PCR using oligonucleotides DeprxF1 and DeprxR1 (Table S2) 229 resulting in the amplification of a ca. 5-kb amplicon in which the prx5-grx coding sequence 230 was deleted and a unique BamHI restriction site had been introduced. The BamHI site was 231 used to introduce a DNA fragment containing a kanamycin resistance cassette and $rpsL^+$ (a 232 dominant selectable marker) which was generated by PCR from N. meningitidis MC58 Kan^R 233 rpsL⁺ rpsL⁻ (Table S1) using oligonucleotides KnsLF1 and KnsLR1 (Table S2), in place of 234 prx5-grx, resulting in plasmid pGUD2. Plasmid pGUD3 was generated by self-ligation of the 235 same inverse PCR product. 236

pGUD2, containing the resistance cassette in the same orientation as the deleted gene, was subsequently used to mutate meningococcal strain MC58 $rpsL^{-}$ by natural transformation and allelic exchange as previously described [38] generating MC58 $Kan^{R} rpsL^{+} rpsL^{-} \Delta prx5$ -

240 grx. pGUD3 was then used mutate MC58 $Kan^{R} rpsL^{+} rpsL^{-} \Delta prx5$ -grx to generate the

241 markerless mutant MC58 $rpsL^{-}\Delta prx5$ -grx.

242

243 2.9. Complementation of prx5-grx

244	A fragment corresponding to the <i>prx5-grx</i> coding sequence and upstream sequence was
245	amplified from strain MC58 using oligonucleotides PerCompF1 and PerCompR1 (Table S2)
246	incorporating BglII sites into the amplified fragment. The BglII-digested fragment was then
247	introduced into a unique BglII-site in pYHS25 (Table S3). This vector contains an
248	erythromycin resistance gene flanked by the MC58 genes NMB0102 and NMB0103. The
249	resulting plasmid, pLD1, was used to transform MC58 $rpsL^{-} \Delta prx5$ - grx by natural
250	transformation, thus introducing a single chromosomal copy of <i>prx5-grx</i> and the downstream
251	erythromycin resistance cassette into the intergenic region between NMB0102 and
252	NMB0103, generating MC58 $rpsL^{-} \Delta prx5$ - $grx prx5$ - grx^{ECT} .
253	
254	2.10. Whole cell ELISA
255	Mid-log phase (OD ₆₀₀ \ge 0.5) liquid cultures of meningococcal strains were
256	centrifuged at 2087 × g at 4°C for 10 min. After discarding the supernatant, the bacterial
257	pellet was washed twice with sterile-filtered PBS and resuspended in 0.5% v/v formaldehyde
258	in PBS. The cells were incubated at 4°C for 30 min with gentle agitation. Cells were then
259	centrifuged at 2087 \times g for 10 min at 4°C, washed one more time with PBS, resuspended in
260	sodium carbonate buffer and the OD_{600} adjusted to 0.2. 100 µl aliquots of formaldehyde-fixed
261	meningococcal cells in carbonate buffer were then used to coat microplate wells overnight at
262	4°C with gentle agitation. Plates were then washed three times with PBST and further steps
263	were carried out as described in section 2.6.

264

265 2.11. Hydrogen peroxide survival assay

266 Sensitivity to hydrogen peroxide stress was determined by a method based on that described by Tala et al [39]. Briefly, meningococcal strains were grown to mid-log phase 267 $(OD_{600} \ge 0.5)$. Suspensions were then adjusted to *ca*. 10⁸ CFU ml⁻¹ (confirmed retrospectively) 268 269 by plating out serially diluted aliquots). 100 µl aliquots of the meningococcal cell suspension were added to a sterile 96-well polystyrene round bottom microwell plate (Thermo Fisher 270 Scientific) containing 150 µl BHI. Fifty microliters of H₂O₂ was then added to yield a final 271 concentration of 12.5 mM, and the plate was incubated at 37°C in 5% CO₂ atmosphere with 272 gentle shaking for 1 h. At 20 min intervals, 10 µl aliquots were taken, serially diluted, and 273 274 plated out. After overnight incubation at 37°C in 5% CO₂, colonies were counted and expressed as CFU ml⁻¹. 275

276

277 2.12. Ex vivo human whole blood model of bacteremia

278 Whole venous blood was collected from healthy individuals (not immunized against *N*. 279 *meningitidis*) and supplemented with heparin (17 IU ml⁻¹) to inhibit coagulation. 280 Meningococci were grown to mid-log phase and then diluted in BHI broth to approximately 281 10^8 CFU ml⁻¹. The assay was started by the addition of 190 µl whole human blood to 10 µl of 282 bacterial suspension. Suspensions were incubated at 37°C and 5% CO₂ with gentle agitation 283 for 2 h and at various time points an aliquot of the sample was removed and the number of

viable CFU determined by plating out of serial dilutions.

285

286 **3. Results**

287 *3.1. Prx5-Grx is a plasminogen-binding protein*

To examine putative host ligand-binding functions of meningococcal Prx5-Grx, and to raise specific antibodies, prx5-grx from the group B meningococcal strain MC58 was expressed in *E. coli* and purified under non-denaturing conditions to yield N-terminally 6 ×

histidine-tagged recombinant Prx5-Grx (Fig. S1). Since surface-localized bacterial
moonlighting proteins are recognized to bind a variety of host proteins [40] we screened the
ability of rPrx5-Grx to bind to human plasminogen (glu-plg), fibronectin, laminin and
collagen. Of these potential ligands, Prx5-Grx was only able to significantly bind to glu-plg
(Fig. 1A). Binding to glu-plg could be inhibited by the lysine analogue, ε-aminocaproic acid
(EACA) (Fig. 1B).

297

3.2. Lysine 244 is required for optimal binding of meningococcal Prx5-Grx to plasminogen 298 299 Given the likely involvement of lysine residues in the binding of rPrx5-Grx to glu-plg, the two C-terminal lysine residues (²³⁰K and ²⁴⁴K) of rPrx5-Grx were individually, or in 300 combination, replaced with alanine using site-directed mutagenesis and the effects on glu-plg 301 binding were examined. rPrx5-Grx^{K244A} exhibited a significantly reduced ability to bind 302 glu-plg, confirming an important role for this lysine residue at the penultimate position in 303 meningococcal Prx5-Grx (Fig. 2). In contrast, mutation of ²³⁰K had no significant effect on 304 glu-plg binding, while the double lysine mutant behaved similarly to the ²⁴⁴K mutant (Fig. 2). 305 No statistically significant difference in glu-plg binding was apparent in assays utilizing 306 rPrx5-Grx^{C185A}, in which the second cysteine residue of the Grx active site was substituted, 307 confirming that glutaredoxin activity is not required for the binding of glu-plg (Fig. 2). 308

309

310 *3.3. Generation of prx5-grx knock out and complemented strains*

To examine the roles of Prx5-Grx in the meningococcus, a knockout derivative of *N*. *meningitidis* MC58 was generated using a markerless mutation strategy. Briefly, prx5-grxplus flanking DNA was cloned and inverse PCR used to remove the entire ORF. The product was then ligated to a DNA fragment harbouring a kanamycin resistance cassette plus the dominant marker, $rpsL^+$, (conferring streptomycin sensitivity), and the resulting plasmid used

316 to transform a streptomycin-resistant N. meningitidis MC58 derivative (MC58 rpsL⁻) yielding MC58 $Kan^{R} rpsL^{+} rpsL^{-} \Delta prx5$ -grx. The latter strain was then further transformed using the 317 self-ligated inverse PCR product to yield the final streptomycin resistant, but markerless 318 319 MC58 $rpsL^{-}\Delta prx5$ -grx mutant. Immunoblotting using RaPrx5-Grx showed that a ca. 27-kDa 320 protein corresponding to Prx5-Grx could be detected in whole cell lysates of wild-type but not MC58 $rpsL^{-}\Delta prx5$ -grx (Fig. S2), confirming that expression had been abolished in the 321 322 mutant. To further confirm that the ca. 27-kDa immuno-reactive protein was Prx5-Grx, a wild-type copy of prx5-grx was introduced in trans into MC58 rpsL⁻ $\Delta prx5$ -grx restoring 323 324 Prx5-Grx expression, albeit at lower expression levels than the wild-type (Fig. S2).

325

326 3.4. Loss of Prx5-Grx does not significantly reduce the ability of meningococci to bind
327 plasminogen

A whole cell ELISA assay suggested localisation of meningococcal Prx5-Grx on the surface of wild-type, MC58 *rpsL*⁻ and the complemented mutant (Fig. 3), with the latter exhibiting reduced levels of anti-Prx5-Grx reactivity compared to the parental strain in line with reduced overall Prx5-Grx expression. Importantly, there was no significant difference in the Plg-binding ability of $\Delta prx5$ -grx compared to strains expressing surface-exposed Prx5-Grx (Fig. 4) consistent with previous observations that meningococci possess a range of surface-exposed glu-plg ligands which may be functionally redundant [34, 41].

335

336 3.5. *Prx-Grx-deficient meningococci are protected from killing in a hydrogen peroxide assay* 337 Given the likely involvement of Prx5-Grx in oxidative stress responses, hydrogen 338 peroxide (H₂O₂) killing assays were performed. After 20 min of exposure to H₂O₂, survival of 339 MC58 *rpsL⁻* $\Delta prx5$ -*grx* was significantly enhanced compared to the parental or 340 complemented strains (Fig. 5). With the exception of the *prx5*-*grx* mutant, each strain tested

was undetectable at the 40 and 60 min time points. In contrast, growth curve assays carried out in the absence of H_2O_2 showed no significant differences between strains (Fig. S3). Taken together, the absence of Prx5-Grx expression had no effect on *in vitro* growth under standard conditions, but Prx-Grx-deficient meningococci were protected from killing when exposed to H_2O_2 .

346

347 3.6. Prx5-Grx is important for survival in an ex vivo human whole blood model of 348 meningococcal septicemia

An ex vivo human whole blood model of meningococcal bacteremia was used to assess the contribution of Prx5-Grx to survival on exposure to cellular and humoral mechanisms of killing. In both donors tested, there was $a \ge 2$ -log₁₀ difference in CFU between MC58 *rpsL*⁻ $\Delta prx5$ -*grx* and the parental strain at 30 and 60 min time points. Survival of the complemented mutant was similar to that of the parental strain, suggesting that even the lower level of expression of Prx5-Grx observed in the complemented strain was sufficient to provide full protection against killing in human blood (Fig. 6).

356

357 **4. Discussion**

Prx5-Grx is a highly conserved protein in Neisseria species [42] and a previous study 358 demonstrated that both the Prx and Grx domains are biochemically functional in the 359 meningococcal enzyme [33]. Prx5-Grx is upregulated in response to heat shock in N. 360 meningitidis [43] and a deletion mutant in N. gonorrhoeae was found to have a defect in 361 biofilm formation [35]. In N. meningitidis, Prx5-Grx was also identified as a putative plg-362 binding protein on the bacterial surface [34]. In this study, we confirm that Prx5-Grx is a 363 multifunctional protein with plg-binding activity, but show that it is functionally redundant 364 on the bacterial cell surface in the presence of alternative plg-binding proteins. Importantly, 365

we demonstrate that Prx5-Grx is also required, however, for survival of *N. meningitidis* innon-immune blood.

368 Having confirmed that meningococcal Prx5-Grx is able to bind human plg, we 369 investigated which amino acid residues in Prx5-Grx were required for the plg-binding. We observed that binding of plg was sensitive to the lysine analogue EACA, suggesting that 370 certain lysine residues in Prx5-Grx were responsible for the interaction. Knaust et al. reported 371 372 that internal lysine residues (rather than the lysine residues in terminal or penultimate positions) might be important for plg-binding by enolase, DnaK and Prx5-Grx [34]. In 373 374 contrast, we demonstrated that the lysine residue in the penultimate position in Prx5-Grx rather than an internal lysine residue was primarily responsible for plg-binding. The 375 difference in findings may be explained by our use of an EIA-based assay to detect plg-376 377 binding to non-denatured rPrx5-Grx rather than a dot blot assay, or possibly the replacement 378 of both the penultimate lysine residue and the asparagine residue in the terminal position of Prx5-Grx in the study by Knaust et al [34]. 379

380 *N. meningitidis* normally inhabits the mucosal surface of the human nasopharynx or subcutaneous tissues. In these niches, meningococcal cells are continuously exposed to 381 reactive oxygen species (ROS), produced by internal metabolic functions as a natural by-382 product of aerobic respiration, by other commensal microorganisms, or by host metabolism 383 and immune responses, and must adapt rapidly to varying levels of oxidative stress [44]. In 384 385 the upper respiratory tract (and in the bloodstream), macrophages and polymorphonuclear cells, which utilize oxygen-dependent pathways to generate large quantities of ROS, facilitate 386 the clearance of invading organisms [45]. 387

The response to oxidative stress in pathogenic *Neisseria* is controlled by the OxyR regulon, which has been studied in *N. gonorrhoeae* [35], *N. meningitidis* [46] and

390 *H. influenzae* [47], and has been shown to operate in a distinctive manner in these organisms

391 compared to the OxyR regulon in Escherichia coli. The OxyR regulon controls expression of kat in response to H₂O₂ [48, 49], however, it was found that oxyR mutant strains of 392 393 N. gonorrhoeae [35, 50], N. meningitidis [46] and H. influenzae [47] have considerably 394 higher levels of kat expression and are significantly more resistant to H₂O₂ killing than their respective wild-type strains. In N. gonorrhoeae, the OxyR regulon was suggested to be 395 limited to kat and two other genes: prx (designated prx5-grx in this study) and the 396 397 downstream gene, gor [35]. In H. influenzae, the Prx5-Grx homologue was suggested to be involved in scavenging low levels of endogenous H_2O_2 , thus limiting the activation of kat, 398 399 and, thereby, enabling the response to oxidative stress to be tightly controlled at a level proportionate to the stress [25, 47]. In this study, we found that the meningococcal prx5-grx 400 mutant is also more resistant to exogenously supplied H₂O₂ compared to the wild-type and 401 402 complemented strains, which is in keeping with the findings described in N. gonorrhoeae and 403 H. influenzae [25, 35]. Increased catalase expression was determined in both the N. gonorrhoeae and H. influenzae mutants lacking Prx5-Grx expression and suggested as a 404 405 likely explanation for increased H₂O₂ resistance [25, 35]. Further experimentation will be required to establish if catalase expression is also elevated in the meningococcal prx5-grx 406 407 mutant. Interestingly, in H. influenzae, the majority of scavenging of metabolically-generated H₂O₂ was shown to be due to expression of kat or the prx5-grx homologue, however, deletion 408 of both genes did not impair virulence in the infant rat model of infection [51]. 409 410 Ex-vivo whole blood survival assays have been used widely as models of invasive meningococcal disease [52-55]. They are considered to be useful as they are relatively facile 411 and reproduce many of the important constituents of the host-pathogen interaction using cells 412 413 from the host relevant to this highly human-adapted pathogen. Killing of N. meningitidis in a whole blood survival assay is mediated by both cellular and humoral components of the 414

415 immune system, although capsulated N. meningitidis are protected from complement-

416	mediated killing in the absence of specific antibody. It has also been shown recently that the
417	presence of heparin, as used in our assay to prevent coagulation, may also reduce
418	complement-mediated killing [56]. The observation that the <i>prx5-grx</i> mutant was more
419	sensitive to killing than its wild-type parent is unlikely to be explained by differences in
420	sensitivity to H ₂ O ₂ as this mutant was less sensitive in an H ₂ O ₂ killing assay. It also cannot be
421	explained by changes in levels of plasminogen binding, which might act to inhibit
422	interactions with phagocytes, as the mutant cells bound as much plasminogen as their wild-
423	type parents in ELISA assays, presumably due to the presence of other identified
424	plasminogen-binding proteins on the meningococcal surface. The mechanism by which Prx5-
425	Grx promotes meningococcal survival remains to be determined, but our findings
426	demonstrate that it is likely to play an important role in invasive meningococcal disease.
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431	
432	Conflicts of interest

433 The authors declare that there are no conflicts of interest.







Fig. 1. rPrx5-Grx is a plasminogen-binding protein. (A) Binding of rPrx5-Grx to 436 immobilized human ligands examined by ELISA. rPrx5-Grx bound significantly to glu-plg 437 compared to the negative control immobilized ligand (BSA). ** P value < 0.01 (Student's t-438 test). Binding of rPrx5-Grx to laminin, fibronectin or collagen was not significantly different 439 440 to BSA. (B) Binding of rPrx5-Grx to glu-plg in the presence or absence of 50 mM ϵ aminocapronic acid (EACA). * *P* value < 0.05 (Student's *t*-test) compared to binding in the 441 absence of EACA. Means are from \geq 4 independent experiments carried out in triplicate 442 wells. Error bars indicate SE. 443



Fig. 2. Lysine 244 is required for optimal binding of rPrx5-Grx to plasminogen. No significant differences were detected in the ability of rPrx5-Grx^{K230A} and rPrx5-Grx^{C185A} to bind glu-plg compared to rPrx5-Grx as determined by ELISA. The significant reduction in binding of rPrx5-Grx^{K230A, K244A} and rPrx5-Grx^{K244A} shows that lysine 244 is required for optimal glu-plg binding. BSA was used as a negative control. *, **, *** denote *P* values < 0.05, < 0.01 and < 0.001, respectively (Student's *t*-test) compared to rPrx5-Grx. Means are from \geq 4 independent experiments carried out in triplicate wells. Error bars indicate SE.

455 Wells were coated with formaldehyde-fixed whole meningococcal cells before being probed

456 with rabbit anti-rPrx5-Grx antibodies. Binding of the antiserum was significantly reduced in

457 strains lacking wild-type levels of rPrx5-Grx expression. Complementation of *prx5-grx*

458 restored expression, albeit not to wild-type levels. Values shown are minus

459 the values obtained from control wells coated with 1% BSA. *** and **** denote P values <

460 0.001 and < 0.0001, respectively (Student's *t*-test) compared to MC58. Means are from ≥ 4

461 independent experiments carried out in triplicate wells. Error bars indicate SE.

464 Fig. 4. Loss of surface-localized Prx5-Grx does not significantly reduce the ability of

465 meningococci to bind plasminogen. No significant differences were detected in the ability

466 of the MC58 strains tested to bind glu-plg. In each case, the addition of ϵ -aminocapronic acid

- 467 (EACA) abolished the interaction confirming that all significant interactions at the
- 468 meningococcal surface with glu-plg are lysine-mediated. **** denote P value < 0.0001
- 469 (Student's *t*-test) compared to MC58 $rpsL^2$. Means are from ≥ 3 independent experiments
- 470 carried out in triplicate wells. Error bars indicate SE.

472

473 Fig. 5. Prx5-Grx deficient meningococci have increased tolerance to hydrogen peroxide.

474 Meningococcal cells were resuspended in BHI broth and exposed to a final concentration of

475 12.5 mM H₂O₂. After 20 min of exposure, the survival of MC58 $rpsL^{-}\Delta prx5$ -grx was

476 significantly higher (Student's *t*-test *p* value < 0.05) than MC58 *rpsL*⁻. With the exception of

- 477 MC58 $rpsL^{-}\Delta prx5$ -grx, all strains were undetectable at 40 and 60 min. Three experiments,
- 478 each in triplicate, were performed. Error bars indicate SE of the mean.

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