Reverse transcriptase-PCR differential display analysis of meningococcal transcripts during infection of human cells: Up-regulation of priA and its role in intracellular replication

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Abstract

Background: In vitro studies with cell line infection models are beginning to disclose the strategies that Neisseria meningitidis uses to survive and multiply inside the environment of the infected host cell. The goal of this study was to identify novel virulence determinants that are involved in this process using an in vitro infection system.

Results: By using reverse transcriptase-PCR differential display we have identified a set of meningococcal genes significantly up-regulated during residence of the bacteria in infected HeLa cells including genes involved in L-glutamate transport (gltT operon), citrate metabolism (gltA), disulfide bond formation (dsbC), two-partner secretion (hrpA-hrpB), capsulation (lipA), and DNA replication/repair (priA). The role of PriA, a protein that in Escherichia coli plays a central role in replication restart of collapsed or arrested DNA replication forks, has been investigated. priA inactivation resulted in a number of growth phenotypes that were fully complemented by supplying a functional copy of priA. The priA-defective mutant exhibited reduced viability during late logarithmic growth phase. This defect was more severe when it was incubated under oxygen-limiting conditions using nitrite as terminal electron acceptors in anaerobic respiration. When compared to wild type it was more sensitive to hydrogen peroxide and the nitric oxide generator sodium nitroprusside. The priA-defective strain was not affected in its ability to invade HeLa cells, but, noticeably, exhibited severely impaired intracellular replication and, at variance with wild type and complemented strains, it co-localized with lysosomal associated membrane protein 1.

Conclusion: In conclusion, our study i.) demonstrates the efficacy of the experimental strategy that we describe for discovering novel virulence determinants of N. meningitidis and ii.) provides evidence for a role of priA in preventing both oxidative and nitrosative injury, and in intracellular meningococcal replication.

Background

Neisseria meningitidis (meningoccus) is a transitory colonizer of the human nasopharynx that sporadically provokes life-threatening disease. This microorganism has to interact with cellular barriers for its life cycle [1]. After crossing the nasopharyngeal mucosa, meningococci occa-
sionally spread into the blood stream before moving across the blood-brain barrier causing fatal sepsis and meningitis in otherwise healthy individuals [2-5].

The early stages of an infection with \textit{N. meningitidis} are governed by specific interactions between the pathogen and the epithelial tissues and are quite well known. Indeed, this microorganism has evolved a diverse array of surface structures subjected to phase- and antigen-variation, which promote adherence and entry into human cells, although it is still unclear whether cell invasion is important in human infections. Initial adherence of encapsulated bacteria requires type IV pili, fine hair-like structures protruding from the bacteria surface [3]. Then, interaction with host cells is achieved by several meningococcal surface adhesins (Opa and Opc) that have been extensively studied [1,6]. In contrast, later stages of infection, including the intracellular location of the meningococci and their strategies for intracellular survival, are only poorly understood [1].

\textit{In vitro} studies with cell line infection models are beginning to disclose the essential role of the metabolic adaptation of meningococci to the intracellular environment of the infected cell. The adaptive response includes the stimulation of the capsular biosynthetic genes leading to increased resistance to cationic antimicrobial peptides (CAMPs), important components of the host innate defense system against microbial infections [7], and the activation of the GdhR regulon whose members are involved in metabolism of the available host carbon sources [8,9].

In this work by using reverse transcriptase-PCR differential display (RT-PCR-DD) we have identified a set of meningococcal genes up-regulated during residence of the bacteria in the intracellular host environment including genes involved in L-glutamate transport (\textit{gltT} operon), citrate metabolism (\textit{gltA}), disulfide bond formation (\textit{dsbC}), two-partner secretion (\textit{hrpA-hrpB}), capsulation (\textit{lipA}), and DNA replication/repair (\textit{priA}). PriA is a single-stranded DNA-dependent ATPase, and a 3’ to 5’ DNA translocase/helicase that was discovered originally because of its requirement \textit{in vitro} for the conversion of bacteriophage phiX174 viral DNA to the duplex replicative form [10,11]. In \textit{Escherichia coli} this protein, at the crossroads of DNA replication and recombination, plays a central role in origin-independent, replication restart of collapsed or arrested DNA replication forks and is also involved in DNA recombination [12-14]. These activities rely on the ability of PriA to load replication forks at a D loop, an intermediate that forms during homologous recombination, double-strand break-repair, and stable DNA replication. We investigated the role of \textit{priA} in the meningococcal infectious cycle using a human cell line infection model.

Results

\textbf{RNA differential display analysis of meningococcal gene expression in the intracellular host environment}

In an attempt to identify meningococcal genes selectively up- or down-regulated in the intracellular environment of infected host cells we used a previously published RT-PCR-DD strategy that relies on the presence of highly and moderately repetitive transcribed DNA sequences in the meningococcal genome [8]. Oligonucleotides designed on the basis of the DUS (DUS-IN and DUS-OUT) or the 26L, 27L \textit{nemis} sequences (26L-IN, 26L-OUT, 27L-IN, 27L-OUT) were used as primers in reverse-transcriptase (RT) assays to prepare cDNAs from intracellular meningococci (strain B1940) recovered from saponin-lysed HeLa cells after 7 h of infection or control bacteria grown in DMEM without HeLa cells as previously described [7]. After 7 h of infection bacterial recovery from infected cells reached the highest values [7]. We decided to use HeLa cells because studies with these cells have contributed to our understanding of the molecular mechanisms underlying the infectious cycle of \textit{N. meningitidis} [15-20]. The analysis of the meningococcal transcriptome during the early stages of an infection has been previously performed using HeLa cells [21].

The cDNAs from intracellular meningococci were then amplified by PCR using the corresponding oligonucleotides and a mixture of random hexamers as primers, and the PCRs were analyzed by polyacrylamide gel electrophoresis. By this approach, several bands corresponding to either up-regulated (\textit{a}, \textit{c}, \textit{d}, \textit{e}, \textit{f}, \textit{g}, \textit{l}, \textit{m}, \textit{n}, \textit{o}) or down-regulated genes (\textit{b}, \textit{h}, \textit{i}) were detected in the intracellular bacteria (Fig. 1A). The bands \textit{a}, \textit{c}, \textit{f}, \textit{g}, \textit{l} and \textit{n} were then excised from the gels, the corresponding cDNAs were cloned and subjected to nucleotide sequences analysis. Three clones were sequenced from each band. The bands \textit{d}-, \textit{e}-, \textit{m}-, and \textit{o}-associated cDNAs, when subjected to cloning, each gave rise to more than one clone, and therefore they were not further analyzed in this study. Bands corresponding to down-regulated genes were not examined.

The nucleotide sequence analysis demonstrated that the band \textit{a} and \textit{f}-associated cDNA corresponded to genes that have been shown to be induced in the intracellular environment and have been also implicated in meningococcal pathogenesis (Fig. 1C). The band \textit{a}-associated cDNA corresponded to NMB1964, a gene of the GdhR-regulated \textit{gltT} operon coding for the L-glutamate ABC transporter that is critical for meningococcal adaptation in the low-sodium intracellular environment [8]. The band \textit{f} corresponded to NMB0082 coding for LipA, a protein required for proper translocation and surface expression of the lipidated (\(\alpha_2\rightarrow\beta\))-linked polysialic acid capsule polymer [7,22]. The band \textit{g}-associated cDNA sequence fell within NMB1780-NMB1779 (\textit{hrpB-hrpA}) operon cod-
Figure 1
RT-PCR-DD and limited transcriptional analysis of meningococcal transcripts up- and down-regulated during the intracellular phase. (A) RT-PCR-DD analysis. Oligonucleotides DUS-IN, DUS-OUT, RS3-IN, RS3-OUT or oligonucleotide mixtures 26L-IN/27L-IN, 26L-OUT/27L-OUT were used as primers in first-strand cDNA synthesis using total RNAs from intracellular meningococci (strain B1940) recovered from saponin-lysed HeLa cells after 7 h of infection (intracellular) or control bacteria grown for 7 h in cell culture medium without HeLa cells (control) as templates. Then second-strand cDNAs synthesis was carried out using the corresponding oligonucleotides and a mixture of random hexamers as primers, and the PCR products were analyzed by polyacrylamide gel electrophoresis. In lanes 1 and 6 molecular weight ladders, whose sizes are indicated on the left of each panel, were run in parallel. The arrows indicate bands corresponding to either up-regulated (a, c, d, e, f, g, i, m, n, o) or down-regulated genes (b, h, i) in the intracellular environment. Molecular weight DNA ladders were run in parallel. (B) Limited transcriptional analysis. A partial Sau3AI-restricted genomic library from the strain B1940 was constructed. Individual plasmid clones were digested with Sau3AI and a Southern blot analysis was performed using 32P-labeled cDNA probes derived from intracellular bacteria after 8 h of infection of HeLa cells or control bacteria grown for 8 h in cell culture medium. In the figure only a limited number of clones (22 clones out of 3000) on duplicate filters are shown. The arrow indicates the cloned NMB0551 ORF corresponding to priA, up-regulated by intracellular meningococci. Asterisks mark the 983 and 1123 bp-long Sau3AI fragments contained in the plasmid clone. (C) Genetic map of meningococcal genes up-regulated in the intracellular environment. The positions of the cDNAs (closed rectangles) corresponding to up-regulated genes that were identified by RT-PCR-DD (a, c, g, f, i, m) are indicated with respect to the available genetic map of MC58. Open rectangles with asterisks locate the 983 and 1123 bp-long Sau3AI fragments contained in the priA plasmid clone shown in panel B.
ing for a two-partner secretion (TPS) system that has been recently shown to contribute to adhesion of un-encapsu-
lated bacteria to epithelial cells [23]. The bands c-, I- and n-associated cDNAs corresponded to genes whose role in meningococcal pathogenesis has never been reported so far, including NMB0551 (band c sequence) encoding PriA, the linked NMB0549 and NMB0550 (band I sequence) coding, respectively, for a thiolsulfide oxidi-
doreductase (DsbC) and the ATP-binding protein of an unknown ABC-type transporter (YbjZ), and NMB0954 (band n sequence) encoding GltA, the putative meningococ-
coccal citrate synthase (Fig. 1C). Fig. 1C also shows the genes surrounding priA (NMB0551). Upstream from
NMB0549 (ybjZ) a gene coding for an AcrA/AcrE family protein (NMB0548) is located; downstream to priA and in
the opposite direction a gene coding for a protein of unknown function (NMB0552) and that for a putative transposase (NMB0553) map.

**The role of priA in meningococcal pathogenesis**

The role of PriA in meningococcal pathogenesis was explored in more detail using the HeLa infection model. It is noteworthy that over-expression of priA was also observed in an independent screening by limited transcrip-
tional analysis (Fig. 1B) that was useful to demon-
strate up-regulation of lipA in a previous work [7]. This screening was performed using a partial Sau3AI-restricted genomic library from B1940. The library was screened by Southern blot using cDNA probes derived from intracellu-
lar bacteria after 8 h of infection of HeLa cells or control bacteria grown for 8 h in cell culture medium. The screening led to isolation of a plasmid harboring two Sau3AI fragments, 983 and 1123 bp-long (marked by asterisks in
Fig. 1B and 1C), spanning almost the entire priA gene.

Nevertheless, in order to confirm the results of the RT-
PCR-DD analysis, slot blot and RT real-time PCR experi-
ments were performed (Fig. 2). Semi-quantitative analysis by slot blot experiments demonstrated that the priA-spe-
cific RNA was about three-fold more abundant in menin-
gococcal meningococci grown from saponin-lysed HeLa cells than in control bacteria grown in DMEM without HeLa cells (Fig. 2A–B). This result was confirmed by RT real-time PCR demonstrating an about four-fold increase in the amount of priA transcripts in intracellular meningococci (Fig. 2C). It should be noted that RT-PCR-DD, limited transcrip-
tional analysis and RT real-time PCR were performed on RNAs from different sets of infection experiments thus enhancing the argument for a role of this gene in the intracellu-
lar environment.

To gain functional information about the role of priA in
*N. meningitidis*, we decided to inactivate it insertional by transformation with pDEXpriA (Fig. 3A and Materials and Methods). Southern blot analysis confirmed the insertion

of this suicide plasmid into the priA coding region by a
single cross-over event (Fig. 3B). As a result of the recom-
bination event, two *Hinfl* DNA fragments of the expected
sizes (4019 bp and 852 bp) were detected by priA-specific probe in the transformed strains B1940*priA* (Fig. 3B, lanes 1–3) in place of the 1568 bp *Hinfl* fragment of the parental strain B1940 (Fig. 3B, lane 4). In order to be sure that priA inactivation was causal to all of the phenotypes observed below, a functional copy of priA was introduced into *leuS- *Flam* region of B1940*priA* by using the integra-
tive plasmid pACpriA. Successful gene delivery was demonstrated by Southern blot analysis (Fig. 3C). In two transformants (B1940*priA/priA*) the priA-specific probe detected the presence of both the 1568 bp *Hinfl* fragment (typical of the wild type priA) and the 4019 bp and 852 bp *Hinfl* fragments (typical of the inactivated
*priA*) (Fig. 3C, lanes 3 and 4).

Phenotypic analysis demonstrated that priA inactivation resulted in a growth defect. Indeed, under aerobic condi-
tions, B1940*priA* strain failed to grow in diluted inocula
(= or <10^6 CFU ml\(^{-1}\)), but when it was inoculated at higher densities (= or >10^7 CFU ml\(^{-1}\)) it grew at rates similar to those of the parental strain (Fig. 4A). However, in high-
density inocula viability of B1940*priA* as determined by
CFU was apparently reduced when compared to that of
B1940. In the experiment shown in Fig. 4C, left panel,
B1940, B1940*priA* and B1940*priA/priA* were grown aero-
bically (not shown) or oxygen-limiting conditions (Fig. 4C,
right panel). Under both these conditions the colony
number of B1940*priA* was reduced to about 15% when
compared to that of B1940 under the same conditions
(Fig. 4B). Statistical analysis with the Student’s T-test con-
firmed that the growth defects in the presence of nitrite in
oxygen limited conditions in the presence of 5% CO\(_2\), and the number of CFU determined after
24 h. Results demonstrated small colony size (Fig. 4C)
and reduced colony number (to about 34%) of the priA-defective strain compared to the parental one (Fig. 4B).
This phenotype was much more severe when GC agar
plates containing nitrite were incubated under either aer-
obic (not shown) or oxygen-limiting conditions (Fig. 4C,
right panel). Under both these conditions the colony
number of B1940*priA* was reduced to about 15% when
c ompared to that of B1940 under the same conditions
(Fig. 4B). Statistical analysis with the Student’s T-test con-
firmed that the growth defects in the presence of nitrite or
in oxygen limited conditions in the presence of nitrite
were more severe than under aerobic conditions without

nitrite (p value < 0.05). The growth defect of B1940*priA*
was not observed during middle logarithmic phase (data
not shown), and could be completely restored by comple-
mentation with pACpriA in B1940*priA/priA* (Fig. 4B
and 4C).

We next investigated the mechanism underlying the
growth defect of B1940*priA*. As there is evidence that in
*E. coli* priA inactivation results in defective cell division
that generates long filaments [24], we used scanning elec-
tron microscopy (SEM) to analyze morphology B1940priA. Indeed, the growth defect of this strain as determined by CFU method might have been caused by a cell division defect that in meningococci, which divide in alternating planes generates clumps. However, SEM analysis did not reveal any difference between the wild type and the priA-defective strain grown to late logarithmic phase (1.0 \ OD_{600\ nm}) (Fig. 5).

To confirm the hypothesis that the growth impairment of B1940priA might be due to reduced viability, the Live/Dead staining assay was carried out. This assay employs two fluorescent nucleic acid stains, SYTO9 and propidium iodide, which differ in their ability to penetrate healthy bacterial cells. When used alone, the green-fluorescent SYTO9 stain labels both live and dead bacteria. In contrast, the red-fluorescent propidium iodide stain penetrates only bacteria with damaged membranes, reducing SYTO9 fluorescence when both dyes are present. Thus, live bacteria fluoresce green, while dead bacteria fluoresce red (Fig. 6A). Using this system we did not observe any significant changes in viability between B1940 and B1940priA when stain was carried out with bacteria grown in GC medium to middle logarithmic phase (0.5 \ OD_{600\ nm}). Indeed, the percentage of nonviable cells was approximately 10% for both strains. In contrast, B1940priA exhibited marked loss of viability during late logarithmic phase (1.0 \ OD_{600\ nm}) when the percentage of nonviable cells approximated 35% (Fig. 6B). The defect in viability was completely restored by complementation (Fig. 6A and 6B). The Live/Dead staining assay also confirmed the absence of any cell division defect in B1940priA (Fig. 6A).

Assessment of viability after exposure to hydrogen peroxide (Fig. 7A) or the nitric oxide generator sodium nitroprusside (Fig. 7B) also demonstrated that B1940priA was much more sensitive to oxidative and nitrosative injuries when compared to the parental strain. The phenotype could be complemented back to wild type levels in B1940priA/priA+.

B1940, B1940priA and B1940priA/priA+ were then used in cell invasion and intracellular persistence assays. In these experiments, high-density-inocula of meningococci were used and infections were initiated by using comparable numbers of viable bacteria for both strains as determined by both CFU method and Live/Dead staining. After allowing meningococci to invade HeLa cells for 1 h, intracellular viability was assessed immediately after gentamicin (time 0 h) treatment and 3, 5 and 7 h post-gentamicin treatment (Fig. 8). Results demonstrated the absence of statistically relevant differences in the ability to invade the HeLa cells by the three strains (Fig. 8, time 0 h). At 3 h post-gentamicin treatment the number of recoverable wild type, mutant and complemented bacteria dropped about ten-fold. Normally, at this time, a decrease occurs in intracellular viable bacteria, which is also characteristic of gonococcal invasion [7,8,25,26]. Between 3 and 7 h B1940 and B1940priA/priA+ underwent fast duplication and the number of viable bacteria increased more than 50-fold. In particular, intracellular duplication was extremely fast during the first 2 h after gentamicin treatment (time 3–5 h) with an apparent generation time of about 24 min and slower during the last 2 h (time 5–7) consistent with previous findings [7,8]. In contrast, between 3 and 7 h the number of viable B1940priA bacteria increased less than 10-fold with an apparent generation time of 96 min during the first 2 h after gentamicin treatment (time 3–5 h). It should be noted that intracellular generation times are only apparent, as they reflect the sum of intracellular growth, intracellular death and, eventually, bacterial egression from HeLa cells and re-invasion. Therefore different processes may be affected in this mutant.

To further investigate the origin of the growth/survival defect of B1940priA inside HeLa cells we used immunofluorescence microscopy. Intriguingly, we noted that number of intracellular priA-defective bacteria 7 h after infection was about the same as those of wild type and complemented bacteria (Fig. 9 and data not shown). This seemed to be in contrast with the data obtained with the CFU method showing a marked decrease in the number of intracellular priA-defective bacteria 7 h after infection (Fig. 8). However, in contrast to CFU method, immunofluorescence analysis detects even dead bacteria, and this means that most priA-defective meningococci inside HeLa cells were actually not viable. The evidence that, at variance with wild type and complemented bacteria, most priA mutant bacteria co-localized with the lysosomal associated membrane protein 1 (LAMP1) strongly supported this hypothesis. This finding is noteworthy because live meningococci have been shown to use IgA protease to cleave LAMP1 protein [27], accounting for the lack of co-localization of wild type and complemented bacteria with this marker. On the basis of this evidence we concluded that the growth phenotype of the priA-defective mutant inside HeLa cells was mostly due to reduced viability than to reduced growth rate.

**Discussion**

The goal of our study was to assess the reliability of RT-PCR-DD as a tool to identify genes with increased or decreased expression during meningococcal growth in infected cells. Although this analysis was clearly not exhaustive as the repeat sequences that we used to prime the RT-PCR-DD reactions were not randomly distributed in the chromosome, and not found in every gene [28], nevertheless it enabled us to identify three up-regulated...
genes/operons (gltT, lipA, hrpA-hrpB) with known roles in virulence (thus validating this approach) and three novel potential meningococcal virulence factors (Fig. 1). One of
these, priA, was selected for extensive further validation (Fig. 2) and characterization.

In infected cells meningococci have to adapt their metabolism to the available host carbon and nitrogen sources and circumvent or subvert a number of host defense mechanisms including lysosomal killing, antimicrobial peptides and oxidative injury. Up-regulation of the gltT operon, coding for an L-glutamate ABC type transporter that is critical for meningococcal adaptation in the low-sodium intracellular environment [8], and of gltA, coding for the citrate synthase, catalyzing the first committed step of the citric acid cycle is part of a metabolic strategy to survive and grow in the intracellular environment. Indeed, in the intracellular milieu there is little available glucose, as glucose entering cells is rapidly phosphorylated to glucose-6-phosphate [29], a substrate that, as well as the other glycolytic intermediates, cannot be assimilated by meningococci. The best intracellular carbon sources are pyruvate and lactate or/and several amino acids such as L-glutamate that stimulate the citric acid cycle [26,30].

Stimulation of L-glutamate uptake and respiratory metabolism is also useful to prevent oxidative injury, a major source of DNA damage, as this amino acid is the precursor of glutathione [29]. In this context the up-regulation of thiol-disulfide oxidoreductase (DsbC)- and PriA-encoding genes (Figs. 1 and 2), and the impaired ability of the priA-defective mutant to replicate in infected HeLa cells (Figs. 8 and 9) may be relevant. PriA plays a central role in origin-independent replication restart of collapsed or arrested DNA replication forks [12-14]. There is evidence that, even under normal aerobic conditions, most, if not all, replication forks do not proceed from the origin of replication to the terminus without hitting a roadblock, in the form of a damaged DNA template, a nick in the template, a “frozen” protein-DNA complex, or local positive superhelicity, which have the potential to arrest or collapse the fork. Inability to reassemble the replisome and resume replication in a timely manner after repair of DNA damage can lead to chromosomal rearrangements and consequently, a loss in viability [31].

Oxidative injury is a major cause of DNA replication fork arrest [32]. This accounts for the growth defect mostly due to reduced viability of the priA-defective meningococcal strain (Figs. 4 and 6), which was much more sensitive to oxidative and nitrosative injuries when compared to the parental strain (Fig. 7). Reduced viability was also observed in E. coli priA-defective mutants [24,33]. A growth defect was also shown in the Neisseria gonorrhoeae priA-defective mutant [34]. This strain was also defective in DNA repair and DNA transformation compared to the isogenic parental strain. However, based on staining with Live/Dead BacLight kit the authors concluded that the growth defect could not be attributable to reduced viability. The different biology of the meningococcal and gonococcal infection may account for the different phenotype of the meningococcal and gonococcal priA-defective mutants. Indeed, substantial differences in oxidative defenses between meningococci and gonococci have been reported. When compared to meningococci, gonococci that are usually associated with inflamed urogenital tissues and activated polymorphonuclear leukocytes, have much higher catalase and cytochrome c peroxidase activities in addition to a specific manganese (II) uptake system involved in resistance to superoxide radicals [35].
The growth defect of the priA-defective meningococcal strain was more severe in the presence of nitrite under either aerobic or oxygen-limiting growth conditions (Fig. 4). Under these conditions meningococci use nitrite as a terminal electron acceptor in anaerobic respiration. Nitrite reduction via nitric oxide to nitrous oxide is the partial denitrification pathway for this mode of growth. In pathogenic Neisseriae the pathway involves both a copper-containing nitrite reductase (AniA) and a single subunit nitric oxide reductase (NorB) [36,37]. Nitrite reduction leads to the production of nitric oxide, a toxic molecule that exhibits synergistic activity in combination with reactive oxygen species [38] and that in Salmonella induces DNA replication arrest [39]. The inability to restart nitric oxide-promoted arrested replication forks may account for reduced viability of the priA-defective mutant under oxygen-limiting conditions.

The relevance of nitrite reduction and defense mechanisms against nitric oxide injury in neisserial pathogenesis is the subject of investigation. Nitrite is present in the diet and is naturally formed in human blood and tissues due to the oxygenation of nitric oxide that is produced by various cell types as a signaling molecule and as a weapon against pathogens and tumor cells. It has been proposed that in microaerobic environments including the human nasopharynx and in the nitric oxide-enriched intracellular environment, meningococcal partial denitrification contributes with oxygen respiration as a route for electron transfer in respiration [37,40]. Interestingly, inside HeLa cells at variance with wild type and complemented bacteria, priA-defective meningococci co-localized with LAMP1 (Fig. 9) that marks highly oxidative and nitrosative cell compartments [41]. Based on this evidence, it is reasonable to assume that failure of the priA-defective mutant to replicate efficiently in HeLa cells may reflect the reduced ability to restart arrested replication forks.

Conclusion

In conclusion, in this study we, on one hand, demonstrate the efficacy of the experimental strategy that we describe, i.e. the analysis of the meningococcal gene expression, for disclosing the mechanisms that N. meningitidis uses to survive and multiply inside the environment of the human cell. On the other hand, we provide evidence that priA, whose expression increases in the intracellular environment, plays a key role in preventing both oxidative and nitrosative injuries, and in intracellular meningococcal replication.

Methods

Bacterial strains and growth conditions

N. meningitidis serogroup B strain B1940 was used in this study. The origin and genotype of this strain have been previously reported [42]. Meningococci were cultured either aerobically in GC broth or agar with 1% Polyvitox, at 37°C in a 5% CO2 incubator or under oxygen-limiting conditions in GC agar containing 5 mM sodium nitrite with 1% Polyvitox, at 37°C in an anaerobic jar with an anaerobic atmosphere generator (GENbox anaer, Biomerieux) and an indicator (Anaer indicator, Biomerieux).

E. coli strain DH5α [F-Φ80d lacZΔM15 endA1 recA1 hsdR17 supE44 thi-1 h- gyrA96 Δ(lacZYA-argF) U169] was used in cloning procedures. This strain was grown in Luria Bertani (LB) medium. To allow plasmid selection, the LB medium was supplemented with ampicillin (50 μg ml⁻¹).

Cell culture and infection

For standard invasion and intracellular viability assays, HeLa cells (ATCC Number CCL-2) were used. Several experiments were also repeated with HEp2 (ATCC Number CCL-23) and Chang conjunctiva cells (ATTC
Effects of priA inactivation on meningococcal viability. Meningococcal strains B1940, B1940ΩpriA and B1940ΩpriA/priA+ were grown to either middle (0.5 OD$_{600}$ nm) or late (1.0 OD$_{600}$ nm) logarithmic phase and then viability was determined by using the Live/Dead BacLight bacterial viability kit (Molecular Probe). (A) Representative images of Live/Dead-stained meningococci: live bacteria fluoresce green, while dead bacteria fluoresce red. (B) Percentages of non-viable meningococci were equal to ratios between bacteria staining positive with propidium iodide (red) and the total stained (red + green) bacteria. Data are shown as mean ± standard deviation of three experiments. Statistically significant difference between values from B1940 (or B1940ΩpriA/priA+) and B1940ΩpriA (asterisk) is declared at a p value < 0.05.
It should be noted, however, that, as a result of a well known contamination event, HEp2 (thought to be derived from an epidermoid carcinoma of the larinx) and Chang conjunctiva (thought to be derived from normal conjunctiva) cells present HeLa markers and have been established via HeLa cells contamination, as stated by ATCC [http://www.atcc.org]. These cells were grown at 37°C in a 5% CO₂ incubator in DMEM supplemented with inactivated 10% fetal bovine serum (FBS), 2 mM L-glutamine, penicillin (50 U ml⁻¹) and streptomycin (50 μg ml⁻¹).

*N. meningitidis* invasion assays were performed as previously described [8,7]. In brief, HeLa (or HEp2 or Chang conjunctiva) cells were infected at a multiplicity of infection (MOI) of 50 for 1 h. To start the infection bacteria were centrifuged (60 × g) onto cells. Cells were washed twice with phosphate-buffered saline (PBS) to eliminate the majority of extracellular bacteria and then viability was determined by recovery of viable bacteria. Data are shown as mean ± standard deviation from three independent experiments, each with triplicate samples. Statistically significant differences between values from B1940 and B1940ΔpriA (asterisks) are declared at a p value < 0.05.

**DNA procedures and genetic manipulation**

High-molecular-weight genomic DNA from *N. meningitidis* strains was prepared as described previously [43]. Oligonucleotides used in this study as primers in PCRs are listed in Table 1. Oligonucleotide synthesis was performed as a service by MWG-Biotech AG Oligo Production. The amplification reactions generally consisted of 40 cycles including 45 s of denaturation at 94°C, 45 s of annealing at 65°C, and 45 s of extension at 72°C. They were carried out in a Perkin-Elmer Cetus DNA Thermal Cycler 2400. DNA from strain B1940 was used as a template. Southern blot hybridizations were carried out according to standard protocols [44]. 32P labeling of the DNA fragments was performed by random priming using
Immunofluorescence analysis of HeLa cells during infection. HeLa cells were infected with B1940, B1940ΩpriA or B1940ΩpriA/priA+ as indicated. Gentamicin selection was used to eliminate extracellular bacteria. Images were taken 7 h after infection. Antibodies against *N. meningitidis* were used before permeabilization with saponin in combination with a secondary antibody conjugated with Cy5 and, after permeabilization, with a secondary TRITC-conjugated antibody. Intracellular bacteria were therefore red while extracellular bacteria were expected to be purple (the combination of TRITC and Cy5 signal). In the images all bacteria were intracellular. To detect cellular markers we used anti-LAMP1 and a secondary antibody conjugated with FITC.
the Klenow fragment of the *E. coli* DNA polymerase I and 
$^32P$-dATP and $^32P$-dGTP (3,000 Ci mmol$^{-1}$) [44].
DNA sequencing was performed as a service by the MWG
Biotech Custom Sequencing Service. Processing of the
DNA sequences was performed with the software Gene-
Jockey Sequence Processor (published and distributed by
Biosoft). Deduced amino acid sequence similarity search-
ing was performed with the BLAST program using the
Conserved Domain Database available at the NCBI.

The *Neisseria*- *E. coli* shuttle vectors pDEX and pACNL1
have been previously described [9,45]. pACNL1 is a
pACYC184 derivative harboring a 1480 bp Sau3AI frag-
ment, spanning the meningococcal *leuS-Ψdam* region in
the BamHI site. To construct pDEXpriA, the DNA corre-
sponding to the central segment of the ORF NMB0551
was amplified using the primers NMB0551-1 and
NMB0551-2, and the BamHI-restricted 851 bp PCR prod-
uct was cloned into the BamHI site of pDEX. NMB0551
(*priA*) was inactivated in B1940 by single cross-over event
using pDEXpriA originating the strain B1940ΔpriA.
Transformations were performed by using 0.1 to 1 μg of
plasmid DNA. Transformants were selected on GC agar
medium supplemented with erythromycin (7 μg ml$^{-1}$).
Successful gene inactivation was demonstrated by South-
ern blot hybridization. NMB0551-specific probe was
obtained by 32P labeling of BamHI-restricted PCR frag-
ment. The size of this fragment was 851 bp (primers
NMB0551-1 and NMB0551-2). B1940ΔpriA was comple-
mented by transformation with the integrative plasmid
pACpriA. To construct this plasmid, the *priA* gene (with
flanking regulatory elements) was amplified from strain
B1940 using the primers NMB0551-3 and NMB0551-4
(Table 1). The XbaI/HindIII-restricted 2565 bp PCR prod-
uct was cloned into the XbaI/HindIII sites of pACNL1.
Transformants were selected on GC agar medium supple-
mented with chloramphenicol (5 μg ml$^{-1}$).

**RNA procedures**

Total RNAs were extracted from intracellular bacteria during
the infection of HeLa cells or control bacteria grown in
culture medium as described [8].

The procedure for limited transcriptional analysis has been previously detailed [9,8]. Briefly, a partial Sau3AI-
restricted genomic library from the serogroup B strain
B1940 was constructed. Individual clones were digested and a Southern blot analysis was performed using cDNA
probes derived from intracellular bacteria after 8 h of
infection of HeLa cells or control bacteria grown for 8 h in
cell culture medium. This procedure detects mRNA spe-
cies present in the initial population at a frequency of at
least 1 in 200.

The protocol for RT-PCR-DD analysis has been also
reported [8]. Briefly, total RNAs were extracted from intra-
cellular bacteria during the infection of HeLa cells or con-
trol bacteria grown in cell culture medium, and were used
as templates for first-strand cDNA synthesis in the pres-
ence of the primers DUS-IN, DUS-OUT, 26L-IN plus 27L-
IN, 26L-OUT plus 27L-OUT, RS3-IN, RS3-OUT listed in
Table 1. Then second-strand cDNAs synthesis was carried
out using the corresponding oligonucleotides and a mix-
ture of random hexamers as primers, and the PCR prod-
ucts were analyzed by polyacrylamide gel electrophoresis.
The gel was stained by the silver staining method [44]. The
slices of gel containing the differentially expressed tran-
scripts were cut with a sharp scalpel, and the DNA frag-

### Table 1: Oligonucleotides used in this study

| Name            | Sequence*       |
|-----------------|-----------------|
| NMB0551-1       | 5'-GATTGGGCGGATCCGGGTTGGATTGAAACACGGAAAGCCG' |
| NMB0551-2       | 5'-CCTTGGTCAGAGATCTTTTGCCGAGCCAGTTGGCCAGCCG' |
| NMB0551-f       | 5'-GCAGGTGTTGTTTTCTGTGTTGCCC' |
| NMB0551-r       | 5'-CATAATTCTCCTGCTGGCGTTG' |
| NMB0551-3       | 5'-GAATTAGTGTAGCACAACCCGCTTTCCAAGAGCCGG' |
| NMB0551-4       | 5'-GGCCGAGAAGTTTACAGACGATTCGGGAAATTTTTCTGAAATTTG3' |
| DUS-IN          | 5'-GCCGTCTGAA' |
| DUS-OUT         | 5'-TTCAGACGGG' |
| 26L-IN          | 5'-GTTGATTAACAAATCAGACGAC3' |
| 26L-OUT         | 5'-GTCGTGTTTGGTTAATCCAC3' |
| 27L-IN          | 5'-GTTGATTAATTTTTAAAATCACGGGAC3' |
| 27L-OUT         | 5'-GTCGCTGTTTAAATTTTAATCCAC3' |
| RS3-IN          | 5'-CATTTCCGCCGCAGGCGGAAATC3' |
| RS3-OUT         | 5'-GATTCCGCCGCTGCGGGAATG3' |
| 16S-univ-1      | 5'-CAGCGAGGCGGCGTAAATAC3' |
| 16S-univ-2      | 5'-CGTCAATCTCTTGTGAGTTTT3' |
| 16S-r           | 5'-CTACGCATTTCTACTGCTACAG3' |

*Restriction sites XbaI and HindIII are underlined.
ments were allowed to diffuse into 100 μl of sterile water at 37°C for 4 h under stirring. The eluted DNA was ethanol precipitated using glycerol as a carrier. Then, it was used as a template in a PCR with the corresponding primers. The positive amplicons were cloned in a pGEM easy vector (Promega) and sequenced.

RT-PCR slot blot analysis of meningococcal priA-specific transcripts during infection of HeLa cells was performed as previously described [8]. To prepare gene-specific cDNA probes, the RNAs were subjected to reverse transcription using the oligonucleotides NMB0551-2 or 16Suniv-2 (loading control) as primers. The labeling procedure consisted of a PCR, where 2 μl of cDNA was used as template in a 50 μl mixtures containing 0.2 mM of each dCTP and dTTP, 2.5 μM of each dATP and dGTP, 0.12 μM [α-32P]ATP (3000 Ci mmol-1) and [α-32P]GTP (3000 Ci mmol-1), 0.2 μM forward NMB0551-1 or 16Suniv-1 primers, and 0.05 U μl-1 of Taq polymerase (Perkin Elmer S.p.A). The amplification reaction consisted of 20–25 cycles including 45 sec of denaturation at 94°C, 45 sec of annealing at 55°C and 45 sec of extension at 72°C. The number of cycles was critical to operate in the linear range of the PCR, and it was determined in preliminary experiments. The 32P-labeled cDNA probes were hybridized to different amounts (0.05 to 100 ng) of denatured NMB0551 (priA)- or 16S rRNA gene-specific DNA fragments generated by PCR with the corresponding primer pairs NMB0551-1/NMB0551-2, and 16Suniv-1/16Suniv-2, and fixed onto positively charged Hybond-N+ nylon membranes. The hybridization reactions were carried out in Church buffer [44] at 63°C. Semi-quantitative analysis was performed by directly counting the radioactivity bands by a Phospholmager SI (Molecular Dynamics, Inc., Sunnyvale, CA).

Bacteria viability assay

The viability of meningococcal strains was determined by using the Live/Dead BacLight bacterial viability kit (Molecular Probe). To this purpose, meningococci were grown to either middle (0.5 OD600 nm) or late (1.0 OD600 nm) logarithmic phase in 50 ml of GC medium with 1% Polyvitox as described above. Then 1 ml of bacterial suspension was collected by low-speed centrifugation. Meningococci were washed, re-suspended in 1 ml 0.9% NaCl and mixed with an equal volume of 2X working solution of Live/Dead BacLight containing a 1:1 mixture of SYTO9 and propidium iodide. After 15 min dark incubation, 5 μl of mounted specimens were viewed with a Nikon Optiphot-2 microscope with an episcopic-fluorescence attachment (EFD-3, Nikon).

Hydrogen peroxide and sodium nitroprusside sensitivity assays

B1940, B1940opiA and B1940opiA/priA+ were grown to middle logarithmic phase (0.5 OD600 nm corresponding to about 1 × 109 CFU). Then, 5 ml aliquots (“input” CFU = 5 × 10⁹ CFU for each strains) were placed into 15-ml conical tubes. H2O2 was added to the tubes at final concentrations of 0, 0.5, or 1 mM, and tubes were incubated with shaking for 20 min at 37°C. Cultures were immediately centrifuged and pellets were re-suspended into 5 ml GC broth. These re-suspensions were serially diluted into GC broth and spotted onto GC agar. The plates were incubated at 37°C in a 5% CO2 incubator and the colonies were counted after 24 h of growth. For the sodium nitroprusside sensitivity assay, dilutions of freshly grown liquid cultures of N. meningitidis to middle logarithmic phase (0.5 OD600 nm corresponding to about 1 × 10⁹ CFU) were immediately spread onto GC agar plates containing 1% Polyvitox and the nitric oxide generator. Plates were incubated in an atmosphere of 5% CO2 at 37°C.

Immunofluorescence microscopy

Immunofluorescence analysis was performed as previously described to distinguish between extracellular and intracellular bacteria [7]. In brief, after infection, cells were washed once with PBS, and fixed with 3% paraformaldehyde. For staining of extracellular bacteria, incubation with primary antibodies was carried out for 20 min at room temperature. After washes in PBS, cells were incubated with secondary antibodies for 20 min in the dark, at room temperature. Subsequently, cells were permeabilized with saponin, incubated with primary antibodies to stain bacteria or intracellular markers and then incubated with secondary antibodies. Rabbit polyclonal anti-N. meningitidis antibody was from ViroStat while monoclonal H4A3 anti-LAMP1 was obtained from the Developmental Studies Hybridoma Bank at the University of Iowa. Primary and secondary antibodies were used at a 1:500 dilution. Mounted coverslips were examined using...
a Nikon Optiphot-2 microscope with an episcopic-fluorescence attachment (EFD-3, Nikon). Image processing was carried out with Adobe Photoshop version 7.0.

Scanning electron microscopy

Meningococcal strains B1940 and B1940QpriA were grown to late (1.0 OD\textsubscript{600 nm}) logarithmic phase in 50 ml of GC medium with 1% Polyvitox as described above. Then 1 ml of bacterial suspension was collected by centrifugation. For SEM observations, meningococci were fixed of GC medium with 1% Polyvitox as described above.

Authors' contributions

AI and MDS carried out the experiments, CB participated in the design of the study and its coordination, PA conceived the study and wrote the manuscript. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

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