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Genome Wide Expression Profiling Reveals Suppression of Host Defence Responses during Colonisation by Neisseria meningitides but not N. lactamica

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Abstract
Both Neisseria meningitidis and the closely related bacterium Neisseria lactamica colonise human nasopharyngeal mucosal surface, but only N. meningitidis invades the bloodstream to cause potentially life-threatening meningitis and septicemia. We have hypothesised that the two neisserial species differentially modulate host respiratory epithelial cell gene expression reflecting their disease potential. Confluent monolayers of 16HBE14 human bronchial epithelial cells were exposed to live and/or dead N. meningitidis (including capsule and pili mutants) and N. lactamica, and their transcriptomes were compared using whole genome microarrays. Changes in expression of selected genes were subsequently validated using Q-RT-PCR and ELISAs. Live N. meningitidis and N. lactamica induced genes involved in host energy production processes suggesting that both bacterial species utilise host resources. N. meningitidis infection was associated with down-regulation of host defence genes. N. lactamica, relative to N. meningitidis, initiates up-regulation of proinflammatory genes. Bacterial secreted proteins alone induced some of the changes observed. The results suggest N. meningitidis and N. lactamica differentially regulate host respiratory epithelial cell gene expression through colonisation and/or protein secretion, and that this may contribute to subsequent clinical outcomes associated with these bacteria.

Introduction
Neisseria meningitidis and Neisseria lactamica are commensal bacteria that colonise the mucosal surface of the human nasopharynx. On rare occasions, N. meningitidis can enter the bloodstream and cause invasive disease with a reported incidence from 1–3 per 100,000 cases in industrialised countries [1], whilst N. lactamica does not cause invasive disease. N. lactamica is associated with colonisation of the nasopharynx in the first few years of life which wanes with age, the converse being found with N. meningitidis [2,3,4]. Hence, it has been suggested that colonisation with N. lactamica can protect against meningococcal disease, supported by recent studies which show that carriers of N. lactamica develop cross-reacting opsonophagocytic antibodies to N. meningitidis [5]. Colonisation by N. meningitidis involves the adherence of host epithelial cells, which is mediated by components such as the pili [6].

Although N. meningitidis and N. lactamica are closely related bacteria, with 60% similarity in their genomes [7], there are important gene differences between the two neisserial species which affect their interactions with the host. For example, the genes required for capsule expression are present in N. meningitidis but not N. lactamica [7]. The capsule can protect N. meningitidis against phagocytosis [8], complement mediated lysis [9] and prevent desiccation during transmission [10], and is considered an important virulence factor. The presence of a capsule in meningococci reduces both adherence and invasion of nasopharyngeal epithelial cells by masking adhesins and invasins [11]. Mutation of the meningococcal pilE gene, encoding the major pilin subunit, results in greatly reduced adherence of bacteria to endothelial and epithelial cells [12]. It is not known whether an equivalent mutation in N. lactamica would have similar effects. In contrast to the amount of data available on the role of microbial factors in adherence and invasion of eukaryotic cells, there is a lack of information on the comparative host response, in particular with respiratory tract epithelial cells, to N. meningitidis and N. lactamica. A recent review by Schubert-Unkmeir et al. [13] outlines the various human gene expression studies that have been done so far using cell lines other than those representing the respiratory tract in response to N. meningitidis. Microarrays (comprising a limited number of inflammation, adhesion and iron-homeostasis genes) were used to investigate the host response of ME-180 epithelial-like human cervical carcinoma [14] and A431 human epidermoid carcinoma cells [15] to N. meningitidis. Bonnah et al. [14] showed that the mRNA expression of several host genes involved in iron homeostasis were altered upon infection with meningococci, while Plant et al. [15] showed that there was an induction of chemokine receptors and cytokines such as CXCR-4, CXCR-5, IL1A, IL1B, IL18 and IFN-γ , with most of the host genes induced early in infection.

Other microarray studies have investigated the responses of human endothelial cells to N. meningitidis. Using primary human umbilical vein endothelial cells, Linhartova et al. [16] have shown
that pilus-mediated adhesion and growth of meningococci in microcolonies on the host cell surface results in alteration of expression levels of human genes known to regulate apoptosis, cell proliferation, inflammatory response and adhesion, and of genes for signalling pathway proteins such as TGF-β/Smad, Wnt/β-catenin and Notch/Jagged. It was suggested that the response found increased the ability of host cells to withstand apoptotic signals induced by infection, thus allowing the maintenance of normal cell function, and subsequently bacterial colonisation. A human cDNA microarray of 11,835 genes was used to study the response of human brain microvascular endothelial cells to N. meningitidis [17]. Host genes involved in apoptosis, cell adhesion, downstream signalling of integrins (and their negative regulators) and cytoskeleton reorganisation were significantly differentially regulated at 4 and 8 hours post infection. The influence of capsule on host gene expression was investigated by comparing the host response to wild type (WT) MC58 strain with that in response to an isogenic siaD knockout mutant (also known as a cap- mutant), which does not possess a capsule. At 4 and 8 hours post infection, the expression of 49.4% and 45% of host genes, respectively, were considered to be capsule-dependent. Response to N. meningitidis in whole blood has also been investigated using custom-printed cDNA microarrays consisting of about 18,000 genes, with the aim of identifying a serum factor causing cardiac dysfunction in meningococcal septic shock [18]. Two studies have described the response of human meningothalial cells to N. meningitidis and its secreted proteins. Using human broad range cDNA expression arrays for 3528 genes, Wells et al. [19] observed an up-regulation of proinflammatory cytokines such as IL-6 and IL-8, as well as of anti-apoptosis genes in meningothalial cells, suggesting that genes involved in immunity and defence are activated in response to N. meningitidis, but at the same time the host cells are able to resist the damaging effects of the bacteria. A follow up study by Robinson et al. [20] using a microarray comprising of cytokine and apoptosis genes (573 in total) showed that secreted protein preparations from N. meningitidis induced host pro-inflammatory responses and resistance to apoptosis, suggesting that secreted proteins are important in meningooccal-host interactive biology.

More recently, there have been studies reported that have investigated the host response of respiratory tract epithelial cells to N. meningitidis, N. lactamica or components derived from them. Liu et al. [21] investigated the response of BEAS-2B human bronchial cells to purified PorB, a major outer membrane protein present in both N. lactamica and N. meningitidis. The N. lactamica PorB had a different Toll like receptor 2 (TLR2) binding specificity to that from the meningococcus. Compared to the PorB of N. meningitidis, the one from N. lactamica was a poorer inducer of proinflammatory mediators and of TLR2 expression in human airway epithelial cells, an effect also seen with live N. lactamica. With the nasopharyngeal cell line Detroit 562, Tezera et al. [22] found that N. lactamica induced a weak inflammatory response via attenuation of secretory cytokines such as TNF-α and IL-6, and to a lesser extent chemokines such as IL-8 and RANTES, compared to N. meningitidis. These authors have concluded that through TLR1/2 stimulation, by activating PPARγ and inhibiting NFκB activity, N. lactamica plays an important role in suppressing pathogen-induced inflammation in the nasopharyngeal mucosa. Both of these studies were confined to a limited set of genes and focused on late time points, typically at 24 hours post infection.

In this study we have compared the transcriptomes of 16HBE14 human bronchial epithelial cells at time points 0 to 7 hours in response to N. meningitidis and N. lactamica as a surrogate model for the initial stages of respiratory tract colonisation. Both bacteria have been shown to associate with 16HBE14 cells [23]. Unlike in the previous microarray studies which have analysed a comparatively limited set of human genes, the Illumina HumanRef-8 BeadChip covering most of the whole human genome, representing 20,509 genes, was used. We hypothesise that early interactions of N. meningitidis and N. lactamica with respiratory epithelial cells are characterised by differential gene expression and this affects subsequent outcomes. Host gene expression profiles in response to both live and dead WT N. lactamica and N. meningitidis, N. meningitidis capsule (cap-) and pil (pilE) mutants and secreted protein preparations from both bacteria were compared. In particular, using dead bacteria as a comparator, we have focused on host responses resulting from active bacterial processes since meningococcal gene expression, including those involved in processes such as host cell adhesion, amino acid and DNA metabolism is known to change after interaction with epithelial cells [24]. It has also been shown that when in contact with epithelial cells, N. meningitidis adds phosphoglycerol to its type IV pili and this posttranslational modification mediates a regulated detachment of the bacterium from the host, which is thought to facilitate its dissemination [25].

Our results suggest that both N. meningitidis and N. lactamica actively interact with respiratory tract epithelial cells and utilise host resources for energy, perhaps as a means of adaptation and colonisation. In addition, the data indicate that N. meningitidis down-regulates host defence genes whilst N. lactamica induces a proinflammatory response, suggesting specific colonisation processes that may lead to different clinical outcomes. Neisserial secreted proteins appear to be mediators of some of these differential host gene expression changes, suggesting novel mechanisms for modulation of the host response.

Results

Neisserial association and invasion of 16HBE14 bronchial epithelial cells

To assist interpretation of transcriptome data, association and invasion assays were carried out as shown in Figure 1. These allowed the extent of interaction of WT N. lactamica and N. meningitidis, as well as N. meningitidis cap- and pilE- mutants with confluent monolayers of 16HBE14 human respiratory bronchial epithelial cells to be determined. The N. meningitidis serogroup B strain MC58, the genome sequence of which is known [26] and has been widely utilised in pathogenicity research [16,17,19,20], was used. N. lactamica strain Y92-1009 was chosen as it has been evaluated as a vaccine strain for meningococcal disease [27] and its genome sequence has been determined [28]. There were no significant differences in association with epithelial cells between WT N. lactamica and N. meningitidis at 3 and 5 hours. However, significantly more N. lactamica associated at 7 hours (Figure 2A). While there was no significant difference in the numbers that invaded at 3 hours, significantly more N. lactamica, compared to N. meningitidis, invaded at 5 and 7 hours (Figure 2B). Association and invasion by the MC58 cap- mutant was significantly greater compared to the WT from 3 to 7 hours post infection (Figure 2 C and D). In addition, the association of the MC58 pilE- with epithelial cells was significantly less compared to the WT from 3 to 7 hours (Figure 2C), while the invasion by the MC58 pilE- was significantly less compared to the WT from 5 to 7 hours (Figure 2D).

Host responses specific to live N. meningitidis occur at an earlier time point compared to those specific to live N. lactamica

Confluent monolayers of 16HBE14 bronchial epithelial cells were studied under the following conditions from 0 to 7 hours:
mock-infected, infected with killed WT N. lactamica, with live WT N. lactamica, with killed WT N. meningitidis and with live WT N. meningitidis. Figure 3 shows how the lists of differentially expressed genes were compared to obtain genes regulated in response to live WT N. meningitidis, to live WT N. lactamica and common to both. Table 1 shows the number of genes in each group from 3 to 7 hours post infection. At an initial time point of 3 hours, there were 9 times more genes regulated specifically in response to live N. meningitidis (125 genes) compared to N. lactamica (14 genes). At 5 and 7 hours, however, the number of genes regulated specifically in response to live N. lactamica increased until it was almost comparable to those responding to N. meningitidis. At this same time point of 5 and 7 hours, there were also increased numbers of N. lactamica associated with and invading 16HBE14 cells (as shown in Figure 2 A and B). We next determined if there were any biological processes significantly over-represented by the genes regulated specifically in response to live WT N. meningitidis, to live WT N. lactamica and common to both, at each time point from 3 to 7 hours. The following sections describe these host cell responses determined by microarray, the expression of selected genes validated by quantitative real-time polymerase chain reaction (Q-RT-PCR), and in specific instances, by protein expression.

Host genes involved in metabolic and energy production processes were up-regulated in response to live N. meningitidis and N. lactamica at late time points

Microarray analysis identified host genes that were regulated in response to both live WT N. meningitidis and N. lactamica and which clustered using Panther software analysis [29] into the biological process categories of phosphate metabolism (5 to 7 hours) and glycolysis (6 to 7 hours) (Table 2). The expression of the six genes representative of these biological processes is shown in a heat map in Figure 4, with their fold changes listed in Table S1. The time course expression of these six genes from 0 to 7 hours (using Q-RT-PCR) is shown in Figure 5. The genes are STC1 and STC2 (stanniocalcin 1 and 2), which are involved in the regulation of phosphate metabolism, ENO2 (enolase 2), HK2 (hexokinase 2), PFKFB3 and PFKFB4 (phosphofructokinases 3 and 4), which encode enzymes involved in glycolysis. Significant up-regulation of these genes compared to the mock-infected controls in response to both live WT N. meningitidis and N. lactamica occurred from 5 to 7 hours for STC1 and PFKFB4 and from 4 to 7 hours for STC2, ENO2, HK2 and PFKFB3, with expression increasing with time. None of these genes was activated upon addition of killed N. meningitidis or N. lactamica, indicating that the up-regulation was specific to live bacteria.

Immunity and defence genes were down-regulated in response to live N. meningitidis but not N. lactamica at 3 hours

At 3 hours, there was a significant over-representation of genes in the immunity and defence category that were regulated specifically in response to live WT N. meningitidis (p = 3.8E-5). The expression of the genes validated by Q-RT-PCR at 3 hours is shown in a heat map in Figure 6, with their fold changes listed in Table 3. Fourteen out of 15 (93%) of genes in this category were down-regulated in response to live WT N. meningitidis (but not to N. lactamica).

The 3 hour time point was the only one where genes that were regulated specifically in response to live WT N. meningitidis clustered into the immune and defence category. The loss of capsule and pili respectively enhances and reduces the association and invasion of N. meningitidis (Figure 2 C and D). Genes whose expression is altered in response to WT N. meningitidis, N. meningitidis cap- and N. meningitidis pilE- mutants (and so are independent of the capsule and pili) could be mediated by a live, contact-independent process such as secreted proteins. Therefore, we infected 16HBE14 cells with N. meningitidis cap- or pilE- mutants and identified host genes that were similarly differentially expressed in response to live WT and mutant N. meningitidis, compared to mock-infected controls, at 3 hours. Table 4 shows the fold changes of these genes found from the microarray and validated by Q-RT-PCR at 3 hours post infection. Five out of 6 (83%) of these genes were down-regulated with respect to mock-infected controls. The expression of 3 of these validated immune-related genes, C1S (complement component 1, s subcomponent), LCN2 (lipocalin 2) and PI3 (peptidase inhibitor 3) were followed up from 0 to 7 hours (Figure 7). C1S is a component of the complement pathway while LCN2 and PI3 are antimicrobial peptides. Down-regulation of C1S, LCN2 and PI3 (with respect to mock-infected controls) was specifically associated with live WT and mutant N. meningitidis (cap- and pilE-). This occurred at a time point of 3 hours for C1S (Figure 7A), from 3 to 5 hours for LCN2 (Figure 7B) and at 3, 4 and 7 hours for PI3 at the transcript level (Figure 7C1). Measurement of PI3 at the protein level also indicated a down-regulation at 3, 6 and 7 hour time points (Figure 7C2).

N. lactamica activates proinflammatory cytokines and genes encoding transcription factors and a cytoplasmic protein at higher levels compared to N. meningitidis

Biological processes that were significantly over-represented in host genes regulated specifically in response to live WT N. lactamica include cytokine and chemokine mediated signalling pathway at 5 hours, cell proliferation and differentiation at 6 and 7 hours and
Figure 2. Comparing the association and invasion of *N. lactamica* and *N. meningitidis* with 16HBE14 cells. 16HBE14 cells were incubated with *N. meningitidis* and *N. lactamica* and association and invasion assays were carried out at various time points at 3, 5 and 7 hours. (A) and (B) compare the association and invasion of wild type (WT) *N. meningitidis* and *N. lactamica* respectively while (C) and (D) compares the association and invasion of WT *N. meningitidis* with the cap- and pilE- mutants, respectively. Values of colony forming units (cfu) per well represent means from at
mRNA transcription and its regulation at 7 hours (Table 5). The expression of genes validated by Q-RT-PCR and involved in these biological processes is shown in a heat map in Figure 8, with their fold changes to live WT *N. lactamica* listed in Tables 6, 7 and 8. Most of the genes (58%) were activated in response to live but not killed *N. lactamica* compared to the mock-infected controls. They were also activated more and for a longer period of time from 5 to 7 hours compared to the response to live WT *N. meningitidis*. Examples of these genes include IL1A and TNF-α, which were involved in the cytokine and chemokine mediated signalling pathway at 5 hours (Table 6), KLF6, ERRFI1 and IL8, which were involved in the cell proliferation and differentiation process at 7 hours (Table 8).

Figure 3. Identification of genes regulated specifically in response to live *Neisseria* determined by microarray analysis. Genes that were regulated specifically in response to live but not killed bacteria from 0 to 7 hours, for both *N. meningitidis* and *N. lactamica* were determined as follows: genes regulated in response to killed bacteria (compared to mock-infected controls) were subtracted from those regulated in response to live *N. meningitidis* (A) or *N. lactamica* (B) derived at each time point. These genes were then separated into three groups: those that were specific to *N. meningitidis*, those that were specific to *N. lactamica* and those that were common to both (C). These analyses were done for every time point up to 7 hours post infection.

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Secreted proteins of *N. meningitidis* and *N. lactamica* regulate C1S and TNF-α expression respectively in 16HBE14 cells

Microarray and Q-RT-PCR data indicated a differential expression of host genes involved in immunity and defence in response to *N. meningitidis* and *N. lactamica*. Some of these responses e.g. the down-regulation of C1S in response to *N. meningitidis* and the activation of TNF-α in response to *N. lactamica* were mediated by live bacteria, and in the case of *N. meningitidis* were independent of the presence of capsule or pili, suggesting a mechanism involving active production of mediators. Secreted protein preparations from WT *N. meningitidis* and *N. lactamica* were obtained from supernatants cultured in the exponential phase of growth, treated to remove outer membrane vesicles (by ultracentrifugation), depleted of lipooligosaccharides (by passing through polymyxin B columns) and added to the 16HBE14 cells.

*N. meningitidis* and *N. lactamica* secreted protein preparations contained less than 0.1 endotoxin units per ml as determined by the Limulus amoebocyte lysate (LAL) assay. Contamination of secreted protein preparations by outer membrane proteins and/or outer membrane vesicles was assessed by the presence or absence of NspA by Western blotting. NspA is an outer membrane protein present in both *N. meningitidis* and *N. lactamica* and is a major component of their outer membrane vesicles [30]. No NspA was detected in either the *N. meningitidis* and *N. lactamica* secreted protein preparations (data not shown). The addition of *N. meningitidis* and *N. lactamica* secreted protein preparations resulted in the down-regulation of C1S at 5 hours at the transcript level (Figure 10A) and the activation of TNF-α at 3 hours at both the transcript and protein levels (Figure 10B1 and B2), respectively.

**Discussion**

The influence of association and invasion of *N. lactamica* and *N. meningitidis* on gene expression in 16HBE14 epithelial cells

This is the first study to investigate the comparative host response of human respiratory tract cells to *N. lactamica* and *N. meningitidis* using a whole genome microarray platform. Emphasis has been placed on early events (0–7 hours). Firstly, the extent of association and invasion of the two bacteria with 16HBE14 bronchial epithelial cells were compared. There were no significant differences in association and invasion of *N. lactamica* or *N. meningitidis* at 3 hours. We have shown that the early suppression of host defence genes is specifically associated with live *N. meningitidis* 3 hours post infection, which suggests that this specific host response is due to bacterial differences rather than their numbers per se. At later time points of 5 and 7 hours however, more *N. lactamica* associated and invaded the 16HBE14 cells, compared to *N. meningitidis*. There was a concomitant increase in activation of proinflammatory processes associated with *N. lactamica* at these later time points and this may reflect the greater numbers of bacteria present. Another strain of *N. lactamica* (NL4.1) has also been recently shown to invade epithelial cells derived from the respiratory tract [22].

In this study, compared to WT *N. meningitidis*, the *cap* and *pilE* mutants associated and invaded 16HBE14 cells to greater and lesser extents, respectively. These findings are consistent with previous studies. For example, it is known that the expression of the capsular polysaccharide inhibits the invasion of the nasopharyngeal barrier by masking the meningococcal adhesins and invasins [11]. In addition, a role for *N. meningitidis* pili in adherence to epithelial cells has been well documented [6,12].

**Utilisation of host energy production processes by colonising Neisseria**

An up-regulation of host genes involved in the biological processes of phosphate metabolism and glycolysis was associated with both live *N. lactamica* and *N. meningitidis*. This suggests that the two bacteria share a common mechanism for successful colonisa-

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**Table 1.** Number of host genes regulated in response to live WT Neisseria.

<table>
<thead>
<tr>
<th>Time/hour</th>
<th>Specific to <em>N. meningitidis</em></th>
<th>Specific to <em>N. lactamica</em></th>
<th>Common to both</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>125</td>
<td>14</td>
<td>8</td>
</tr>
<tr>
<td>4</td>
<td>204</td>
<td>425</td>
<td>66</td>
</tr>
<tr>
<td>5</td>
<td>186</td>
<td>198</td>
<td>106</td>
</tr>
<tr>
<td>6</td>
<td>247</td>
<td>103</td>
<td>90</td>
</tr>
<tr>
<td>7</td>
<td>220</td>
<td>185</td>
<td>132</td>
</tr>
</tbody>
</table>

**Table 2.** Host biological processes associated with both live WT *N. meningitidis* and *N. lactamica*.

<table>
<thead>
<tr>
<th>Biological Process</th>
<th>Time points (hour)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3</td>
</tr>
<tr>
<td>Regulation of phosphate metabolism</td>
<td>-</td>
</tr>
<tr>
<td>Glycolysis</td>
<td>-</td>
</tr>
</tbody>
</table>

Significance of biological processes is expressed as p-values and dashes indicate no significance was identified at the time point.

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tion whereby they both actively utilise host resources as a way to survive and adapt in the host. The genes STC1 and STC2 (from the regulation of phosphate metabolism pathway) encode members of a family of secreted homodimeric glycoproteins which are involved in phosphate transport, cell metabolism, and cellular calcium/phosphate homeostasis. ENO2 (enolase 2), HK2 (hexokinase 2), PFKFB3 and PFKFB4 (phosphofructokinases 3 and 4) are genes involved in carbohydrate metabolism including glycolysis.

The clustering of host genes regulated in response to both bacteria into the categories of regulation of phosphate metabolism and carbohydrate metabolism suggests that the human host cell represents a milieu rich in nutrients for bacterial growth and a source of energy. Grifantini et al. [23] observed that cell contact of *N. lactamica* and *N. meningitidis* with 16HBE14 cells resulted in the down-regulation of several bacterial genes involved in metabolism. To explain this, the authors suggested that the bacteria were able to utilise part of the ATP synthesised by the host. Similarly, when Dietrich et al. [31] analysed the transcriptome of *N. meningitidis* after contact with epithelial cells (Hela cells) and human brain microvascular endothelial cells, a high proportion of the differentially regulated genes were involved in central metabolism. In addition, genes involved in cell metabolism, particularly in energy production, were found to have increased transcription in human brain endothelial cells in response to *N. meningitidis* [17].

The *pfk* gene (encoding phosphofructokinase) is not found in the genomes of *N. meningitidis* and *N. lactamica*, which explains the lack of a functional glycolytic pathway in these bacteria [26,32,33]. Under anaerobic conditions, where oxidative phosphorylation cannot occur, *pfk* is essential as the glycolytic process is important for the production of energy. For *Neisseria* however, which colonises the aerobic nasopharyngeal mucosa, sufficient energy can be liberated from the substrate by oxidative phosphorylation, and the glycolytic process (in which phosphofructokinase is an enzyme) is not essential. Although *Neisseria* does not carry out glycolysis, host glycolytic enzymes (ENO2, HK2, PFKFB3 and PFKFB4) were up-regulated in response to the bacteria in this study. It is tempting to speculate that the bacteria may be responsible for this up-regulation so as to increase the amount of ATP produced by the host, which in turn may be available for use by the bacteria. Our results are broadly in agreement with another study describing a commensal bacterium using a similar strategy to survive in the human host. *Bacteroides thetaiotaomicron*, a component of the intestinal microflora of humans uses epithelial fucosylated glycans as a source of energy in the highly competitive intestinal ecosystem. In this way, the host appears to be a participant in providing for the nutritional needs of the bacteria [34].

*N. meningitidis* down-regulates immune response genes while *N. lactamica* initiates a proinflammatory response

Results from the microarray validated by Q-RT-PCR, and in selected cases at the protein level, indicate a down-regulation of genes such as C1S, LCN2 and PI3 after infection with *N. meningitidis* relative to *N. lactamica*. In contrast, there was a greater activation of genes such as TNF-α, IL1A, IL8, JUN, ERFF11 and KLF6 after infection with *N. lactamica* relative to *N. meningitidis*. The results suggest that, in broad terms, *N. meningitidis* is associated with a suppression of the host defence response while *N. lactamica* is associated with an activation of the proinflammatory response.

Besides our study, there have been very few investigations comparing host responses to *N. meningitidis* and *N. lactamica*. In one such study looking at a range of limited cytokine and chemokine responses in human meningioma cells, *N. meningitidis* induced higher amounts of proinflammatory markers such as IL8 compared to *N. lactamica* at much later time points of up to 48 hours [35]. It is well documented that during invasive meningococcal disease, potentially life-threatening meningitis and septicaemia arises from the host due to the overwhelming amount of proinflammatory cytokines and chemokines produced [36], and thus it is not surprising host cells derived from the blood and the central nervous system respond analogously in *vitro*. In contrast to this study, we used an epithelial cell line from the respiratory tract and followed responses at early time points of up to 7 hours. Many factors will determine the relative host gene expression in response to *N. meningitidis* and *N. lactamica* and these include cell lineage and the time after infection sampled.

In this study, a specific down-regulation of host defence genes (C1S, LCN2 and PI3) was associated with live *N. meningitidis*. C1S encodes a serine protease, which is a major constituent of the human complement subcomponent C1. C1s associates with two other complement components C1r and C1q in order to yield the first component of the serum complement system. It is widely known that an effective complement system is pivotal for host resistance against *N. meningitidis*. It is, therefore, not surprising that *N. meningitidis* has been found to exploit two negative complement regulators from its human host, factor H [37] and C4 binding protein [9] to reduce the effectiveness of the host complement defence system. For example, *N. meningitidis* mimics the mechanism

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**Figure 4.** Microarray heat map of validated genes up-regulated in response to *N. lactamica* and *N. meningitidis*. The expression of the 6 validated genes from 5 to 7 hours is shown. Each column is the mean signal of 4 (Neisseria infected) to 8 replicates (mock-infected). Red indicates that the signal is higher relative to the rest of the samples while green indicates that the signal is lower.

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Figure 5. Q-RT-PCR of host genes up-regulated in response to both live *N. meningitidis* and *N. lactamica*. Expression at the transcript level is shown for STC1 (A) and STC2 (B), which are involved in the regulation of phosphate metabolism as well as ENO2 (C), HK2 (D), PFKFB3 (E) and PFKFB4 (F), which encode enzymes involved in glycolysis. There was significant up-regulation (with a p-value of less than 0.05) with respect to the mock-infected controls in response to both live but not killed *N. meningitidis* and *N. lactamica*. This occurred from 5 to 7 hours for STC1 (A) and PFKFB4 (F) and from 4 to 7 hours for STC2 (B), ENO2 (C), HK2 (D) and PFKFB3 (E) as indicated by the asterisks (*). M: mock-infected, YK: killed WT *N. lactamica*, YL: live WT *N. lactamica*, MK: killed WT *N. meningitidis*, ML: live WT *N. meningitidis*, RQ: relative quantification. An outlier (0) is defined as
between 1.5 to 3 \times \) the interquartile range from the 25th or 75th percentile while an extreme data point (E) is defined as more than 3 \times \) the interquartile range from the 25th or 75th percentile.

by which host cells regulate complement activation on their surface by facilitating the high affinity interaction between factor H and factor H binding protein on the bacterium. This suggests that \textit{N. meningitidis} could rapidly sequester factor H, an alternative pathway inhibitor, and avoid clearance by the complement system.

[38]. Suppression of C1S, another component of the classical complement pathway, (see Figure 7A and Figure 10A), could be another mechanism to subvert host defences during colonisation.

LCN2 and PI3 encode antimicrobial peptides. LCN2 inhibits microbial growth by limiting iron availability [39] while PI3 is a low molecular weight cationic peptide and acts as an antimicrobial defense-like molecule with the ability to eliminate pulmonary pathogens [40]. The expression and secretion of PI3 is induced in human keratinocytes [41] and in bronchial epithelial cells [42] after exposure to \textit{Pseudomonas aeruginosa}. Other pathogens e.g. adenoviruses have been found to be responsible for the suppression of PI3 in primary human bronchial epithelial cells [43]. In addition, another antimicrobial peptide LL-37 has also been found to be consistently suppressed by \textit{Neisseria gonorrhoeae} (another pathogenic species of \textit{Neisseria}) in a cervical epithelial cell line [44]. Down-regulation of antimicrobial peptides like LCN2 and PI3 by \textit{N. meningitidis} may be a mechanism to reduce the ability of the host to clear the bacterium and to promote colonisation.

The expression of C1S, LCN2 and PI3 is not only specifically down-regulated in response to live \textit{N. meningitidis}, but also in response to the \textit{N. meningitidis cap-} and \textit{pilE-} mutants, despite their different extent of association and invasion of host epithelial cells. This suggested an active bacterial process involved in the down-regulation of these genes. Co-incubation of \textit{16HBE14} cells with preparations of WT \textit{N. meningitidis} secreted proteins indicated their involvement in the down-regulation of C1S. However, they had no effect on the expression of LCN2 and PI3, indicating that other extracellular bacterial components, such as outer membrane vesicles (absent in our secreted protein preparations) may have a role in the differential expression of LCN2 and PI3. Other alternative explanations are that the meningococcal secreted protein(s) mediating differential gene expression of LCN2 and PI3 is produced by bacteria growing in the presence of 16HBE14 cells/serum (rather than defined medium) or when \textit{N. meningitidis} is attached to or has invaded host cells. \textit{N. meningitidis} and \textit{N. lactamica} were grown to log phase in defined medium to allow pure bacterial protein secreted preparations. We did consider the use of supernatants obtained from co-cultures of bacteria with 16HBE14 cells. However, it would have been less clear as to whether any changes in host gene expression were due to bacterial or host derived factors, and thus, this approach was not used.

It has been shown by Liu et al. [21] in another human bronchial epithelial cell line (BEAS-2B) that \textit{N. lactamica} PorB binds to TLR2 and is a poorer inducer of proinflammatory mediators compared to that from \textit{N. meningitidis}. Tezera et al. [22] reported that \textit{N. lactamica} NLA.1 protects mucosal barrier integrity by suppressing \textit{N. meningitidis}-induced inflammation by increasing the expression of PPARγ and inhibiting NFκB activity. Secretory TNF-α and IL-8 were found to be elevated after challenge with \textit{N. meningitidis} compared to \textit{N. lactamica} at 24 hours post infection. In our study, however, relative to \textit{N. meningitidis}, TNF-α and IL-8 expression was up-regulated in response to \textit{N. lactamica} at time points up to 7 hours post infection. Although our results appear to contrast with these two studies, it may not be reasonable to compare our findings with theirs, as our experimental set ups were different in terms of cell line, infection time points, bacterial strains and multiplicity of infection used. With 16HBE14 cells, proinflammatory molecule expression was inhibited by the bacterial pathogen \textit{T. pseudotuberculosis} [45]. In addition, our array data indicates that the expression of TLR2 at the transcript level could not be

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**Figure 6. Microarray heat map of immunity/defence genes regulated specifically in response to live WT \textit{N. meningitidis}.** The expression of the 15 validated genes at 3 hours is shown. Each column is the mean signal of 4 (\textit{Neisseria} infected) to 8 replicates (mock-infected). Red indicates that the signal is higher relative to the rest of the samples while green indicates that the signal is lower. M: mock, YL: Live WT \textit{N. lactamica}, MK: Killed WT \textit{N. meningitidis}, ML: Live WT \textit{N. meningitidis}. doi:10.1371/journal.pone.0026130.g006
detected in our cell line and both PPARγ and NFκB were not significantly differentially expressed in response to N. meningitidis or to N. lactamica, compared to mock-infected controls, at early time points from 3 to 7 hours post infection.

However, our results are not mutually incompatible with those from other studies. Our study found that whilst our N. lactamica strain can induce the expression of proinflammatory markers such as TNF-α, IL1A, IL-8 and JUN, it is also able to activate negative regulators of inflammation, preventing an excessive proinflammatory response within the respiratory tract epithelial cells [50] and ERRF1 has recently been suggested to act as a negative feedback inhibitor of epidermal growth factor receptor signalling through a direct, physical interaction with the epidermal growth factor receptor [51]. The other gene is KLF6, which belongs to the Kruppel-like protein family and inhibits the activity of JUN. Together with KLF2, they have an important regulatory role in controlling and inhibiting numerous host cellular processes, including phagocytosis, proinflammatory cytokine expression and cell proliferation [52].

Certain experimental models investigating host responses to respiratory pathogens have attempted to mimic the cooling effect of constant evaporation from respiratory mucosal surfaces. However, we chose, in line with other previous studies [53,54], to investigate airway epithelial colonisation by meningococci at 37°C. N. meningitidis strains grown on solid agar at 30°C and 37°C were described as having pilated and non-piliated phenotypes, respectively [55]. N. meningitidis MC58 was not used in the latter study. Whilst we did not quantify either pilin-related gene or protein expression, we observed a clear difference in association and invasion of WT N. meningitidis MC58 compared to the pilE-

<table>
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<th>N. meningitidis pilE-</th>
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<td>Array Q-RT-PCR</td>
<td>Array Q-RT-PCR</td>
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The microarray and Q-RT-PCR results show the fold changes of validated genes with respect to mock-infected controls.
doi:10.1371/journal.pone.0026130.t003

Table 3. Validated immunity and defence genes regulated specifically in response to live WT N. meningitidis at 3 hours.

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<th>Description</th>
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<td>Array Q-RT-PCR</td>
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<td>NM_001710</td>
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<td>complement factor B</td>
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<td>NM_002155</td>
<td>HSPA6</td>
<td>heat shock 70 kDa protein 6 (HSP70B)</td>
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<td>LCN2</td>
<td>lipocalin 2</td>
<td>−3.0 −3.5</td>
</tr>
<tr>
<td>NM_0016816</td>
<td>OA51</td>
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</tr>
<tr>
<td>NM_002572</td>
<td>PKD</td>
<td>paroxysmal nonkinesiogenic dyskinesia</td>
<td>−2.0 −1.7</td>
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<td>NM_005729</td>
<td>PPPI</td>
<td>peptidylprolyl isomerase F (cyclophilin F)</td>
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<td>NM_002908</td>
<td>REL</td>
<td>v-rel reticuloendotheliosis viral oncogene homolog (avian)</td>
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<td>SAA2</td>
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The microarray and Q-RT-PCR results show the fold changes of validated genes regulated in response to live WT N. meningitidis with respect to mock-infected controls.
doi:10.1371/journal.pone.0026130.t004

Table 4. Validated genes regulated similarly in response to WT and mutant N. meningitidis at 3 hours.
mutant. This suggests that in our study (at 37°C), WT N. meningitidis MC50 had a functional piliated phenotype.

Another aspect to consider is that differences in the numbers of adherent and invading N. meningitidis and N. lactamica per se might contribute to the host gene and/or protein expression changes observed. For example, the comparative higher activation of proinflammatory genes like TNF-α and IL8 by N. lactamica, compared to N. meningitidis, may reflect the greater numbers of N. lactamica associating and invading the host cells.

In our model both N. meningitidis and N. lactamica adhered to and invaded 16HBE14 cells. Whether bacteria that are adherent induce host gene responses that are different to those that have invaded is unknown but may contribute to the results obtained. One possibility, given that N. meningitidis has the potential to cause invasive disease while N. lactamica does not, is that host responses of respiratory epithelial cells induced by invasive bacteria may differ more between the two organisms than to extracellular colonisation.

**Model of early colonisation by N. meningitidis and N. lactamica**

Based on our results, we propose that N. meningitidis and N. lactamica use both shared and different mechanisms to colonise host respiratory epithelial cells (Figure 11). Both bacteria appear to use similar strategies to utilise host energy resources for growth. However, N. meningitidis appears to down-regulate host defence genes such as those encoding antimicrobial peptides such as PI3 and LCN2 and complement components such as C1S, as a strategy to maintain colonisation. In contrast, N. lactamica does not evade the host immune response, as indicated by activation of genes such as TNF-α, IL1A and IL-8, and that the expression of these cytokines may alert the host to its presence and potentially prevent or limit the bacterium from entering the blood and causing invasive disease. Despite this proinflammatory response, N. lactamica continues to colonise the nasopharynx. This
is likely to be due to an up-regulation of genes such as KLF6 that regulate and prevent an over activation of the proinflammatory response so that the commensal is not cleared completely by the host.

Neisserial secreted proteins appeared to be responsible for some of the host responses specific to N. meningitidis or N. lactamica. Bioinformatic and proteomic studies have identified proteins that are known or predicted to be secreted by N. meningitidis and N. lactamica. A review by van Ulsen and Tommassen [56] used the available genomes of N. meningitidis and of N. lactamica (ST640) to identify genes encoding predicted secreted proteins specific to each bacterium. Our results suggest that secreted proteins can regulate gene expression in respiratory epithelial cells.

In conclusion, our results show that whilst both N. lactamica and N. meningitidis colonise respiratory tract epithelial cells, they have both common and distinct effects on host gene expression and these may be associated with their respective ability to colonise or cause disease.

Materials and Methods

Bacteria and growth conditions
N. lactamica Y92-1009 was obtained from the Health Protection Agency (Porton Down, UK), WT N. meningitidis MC58 and its capsule (cap-) and pili (pilE-) mutants have been described previously [12,57,58]. Formaldehyde inactivation of WT N. meningitidis MC58 and N. lactamica Y92-1009 was carried out as described previously [59]. Neisseria were routinely propagated on gonococcal agar supplemented with 1% Vitox (sGC) or on brain heart infusion agar (BHI) (BD Diagnostics, USA) at 37°C in 5% CO2. Antibiotics were used at the following concentrations for selective growth of the N. meningitidis cap- and pilE- mutants: kanamycin 150 µg/ml and erythromycin 5 µg/ml.

Growth of 16HBE14 bronchial epithelial cells
The 16HBE14 bronchial epithelial cell line is derived from primary human bronchial epithelial cells and transformed by SV40 large T antigen [60]. It has been shown to retain differentiated epithelial morphology and functions and has also been used extensively in cystic fibrosis research [61]. 16HBE14 cells were cultured in Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 2 mM L-glutamine, 1% penicillin and streptomycin and 10% heat inactivated foetal bovine serum (HIFBS) in a humidified incubator at 37°C with 5% CO2. Growth media, antibiotics, supplements and foetal bovine serum were obtained from Invitrogen.

Epithelial association and invasion assays
16HBE14 cells were seeded in 24-well plates (Nunc, Thermo Fisher Scientific, USA) and grown to a confluent monolayer (approximately 2×10^5 cells per well) in DMEM supplemented with 2 mM L-glutamine and 10% HIFBS at 37°C in 5% CO2. Bacteria were added at a multiplicity of infection of 30 (6×10^6 cfu per well) for both association and invasion assays which were performed as previously described [62]. At time points of 3, 5 and 7 hours, wells were incubated with 1% saponin for 10 minutes at 37°C and appropriate dilutions (in DMEM) plated out to obtain viable counts.
Treatment of 16HBE14 human epithelial cells with N. meningitidis, N. lactamica or neisserial secreted protein preparations

16HBE14 cells were seeded into 6-well plates and incubated in DMEM supplemented with 2 mM L-glutamine and 10% HI-FBS at 37°C with 5% CO2. Confluent epithelial cells (approximately 9.5 x 10⁵ cells per well) were washed 3 times with phosphate buffered saline (PBS) and its medium changed before infection with 1.3 x 10⁷ cfu per well of N. meningitidis or N. lactamica (washed with DMEM prior to infection) or treatment with neisserial secreted protein preparations (2 μg per well). At various time points from 0 to 7 hours, the epithelial host cells were washed with PBS and harvested with Trizol (Sigma, UK). For each condition, there were 4 to 8 biological replicates. For neisserial infection

Table 7. Validated genes regulated specifically in response to live WT N. lactamica and associated with cell proliferation and differentiation process at 6 hours.

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doi:10.1371/journal.pone.0026130.t007

Table 8. Validated genes regulated specifically in response to live WT N. lactamica and associated with mRNA transcription regulation, cell proliferation and differentiation and mRNA transcription processes at 7 hours.

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doi:10.1371/journal.pone.0026130.t008
Figure 9. Expression of host genes up-regulated specifically in response to live *N. lactamica*. Activation of ERRFI1, IL8, IL1A, KLF6 and TNF-α was specifically associated with live but not killed *N. lactamica*. This occurred at a time point of 7 hours for ERRFI1 (A) and IL8 (B), at 5 hours for IL1A (C) and at 4 and 7 hours for KLF6 (D) and TNF-α (E1) at the transcript level using Q-RT-PCR. Measurement of TNF-α at the protein level also indicated an up-regulation at the 7 hour time point (E2). Asterisks (*) indicate statistical significance with a p-value of less than 0.05 with respect to mock-infected controls. M: mock-infected, YK: killed WT *N. lactamica*, YL: live WT *N. lactamica*, ML: live WT *N. meningitidis*. RQ: relative quantification. An outlier (O) is defined as between 1.5 to 3 × the interquartile range from the 25th or 75th percentile while an extreme data point (E) is defined as more than 3 × the interquartile range from the 25th or 75th percentile.

doi:10.1371/journal.pone.0026130.g009
experiments, initial bacterial inocula were determined by plating out on sGC or BHI agar for enumeration of viable organisms.

RNA extraction and quantification

Total cellular RNA from human epithelial cells was extracted using the Qiagen RNeasy Mini Kit using on-column DNase 1 treatment according to manufacturer’s instructions (Qiagen, USA). Quantification of RNA samples was performed by checking the absorbance at 260 nm using a NanoDrop 1000 instrument (NanoDrop Technologies, Wilmington, DE).

RNA amplification and microarray hybridisation

Whole genome microarray hybridisations were performed as described previously [63] on RNA samples from mock-infected host cells or those co-cultured with Neisseria. Briefly, total RNA (500 ng) was amplified in a single-round of in vitro transcription amplification that allowed incorporation of biotin-labelled nucleotides using the Illumina TotalPrep RNA Amplification Kit (Ambion, USA) according to the manufacturer’s instructions. Output cRNA was quantified using the NanoDrop ND-1000 UV-Vis Spectrophotometer. cRNA (750 ng) of each sample was hybridised to an Illumina HumanRef-8 V2 BeadChip (containing probes to 20,589 RefSeq gene sequences) at 58°C for 18 hours following the manufacturer’s instructions (Illumina, USA). This was followed by washing, blocking, and streptavidin-Cy3 staining steps, followed by scanning with a high resolution Illumina Bead Array Reader confocal scanner, all carried out following manufacturer’s instructions.

Microarray data analysis

The scanned microarray images were analysed, data extracted and background subtracted using the Illumina Bead Studio v3.1 software (Illumina, USA). The data exported from Bead Studio was then further analysed using Genespring GX7.3 software (Agilent Technologies, USA). Data transformation was corrected for low signals, setting measurement less than 0.01 to 0.01. The standard normalisation procedures were performed as recommended by the Genespring software for one colour array. Per-chip normalisation, where each measurement was divided by the 50th percentile of all measurements in that sample, and per-gene normalisation to median, where each gene was divided by the median of its measurements in all samples, were done. Genes that have a less than 99% confidence of detection above background levels in all the arrays were excluded from the final analysis, leaving 13,226 genes available for statistical differential expression analysis. The normalised signal of each condition was the geometric mean of the normalised signals of its individual biological replicates. Fully annotated microarray data have been deposited in Gene Expression Omnibus (GEO) under accession number GSE27557 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?token=jhullkkyqamygxek&acc=GSE27557).

Group comparisons were made between mock-infected cells and those challenged with bacteria or neisserial secreted protein preparations over a 7 hour time course. Differentially expressed genes were selected from the normalised data using the program Significance Analysis of Microarrays [64], which was incorporated into Genespring GX 7.3. For all comparisons, a false discovery rate of 5% was used. In addition, to remove low signal genes that could give false positive results due to a lack of sensitivity, significant genes, which did not have a raw intensity value of more than 50 in at least half of the samples in the smaller group of comparison, were discounted. For comparisons between mock-infected cells and cells challenged with (live or killed) WT bacteria, significant genes with a less than 2-fold change difference were removed to increase stringency.

Functional clustering of differentially expressed genes was assessed using the Applied Biosystems online program “Panther” gene expression analysis system [29]. These genes were compared with those in the Illumina array reference list to statistically determine over representation of functional categories. A Bonferroni corrected p-value of less than 0.05 was considered significant.

Heat maps were generated using log2 transformed normalised values. The genes were further clustered in Gene Cluster 3.0 (Stanford University, USA) with uncentered correlation as the similarity metric and average linkage as the clustering method. The output file was then visualised using TreeView (EisenSoftware, USA). Red indicates that the gene’s signal is higher relative to its average signal within its condition and blue is the opposite.

Figure 10. Secreted proteins of N. meningitidis and N. lactamica regulate C1S and TNF-α expression, respectively. Treatment of host epithelial cells with secreted protein preparations from WT N. meningitidis indicate that secreted proteins may be involved in the down-regulation of C1S at 5 hours at the transcript level (A). In contrast, secreted protein preparations from WT N. lactamica resulted in an activation of TNF-α at 3 hours at both the transcript (B1) and the protein level (B2). Asterisks (*) indicate statistical significance with a p-value of less than 0.05 with respect to mock-infected controls. M: mock-infected, MLS: WT N. meningitidis secreted protein preparations, YL: live WT N. lactamica, YLS: WT N. lactamica secreted protein preparations. RQ: relative quantification.

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Figure 11. Postulated model of early colonisation by N. meningitidis and N. lactamica. Host metabolic and energy production processes were associated with both neisserial species, suggesting that N. meningitidis and N. lactamica utilise host resources for energy. In addition, differential host responses to N. meningitidis and N. lactamica may indicate different colonisation processes. N. meningitidis down-regulates host defence genes such as complement (C1S) and antimicrobial peptides (e.g. PI3 and LCN2), reducing the ability of the host to clear the bacterium and thus promote colonisation. This may contribute to the ability of N. meningitidis to cause invasive disease in an environment where host defences are compromised. In contrast, N. lactamica does not evade the host immune response as seen from the activation of genes such as TNF-α, IL1A and IL-8, and that the expression of these cytokines may alert the host to its presence and the activation of genes such as TNF-α, IL1A and IL-8, and that the expression of these cytokines may alert the host to its presence and potentially prevent or limit the bacterium from entering the bloodstream and causing invasive disease. Despite this proinflammatory response, N. lactamica continues to colonise the nasopharynx. This is likely to be due to an up-regulation of genes such as KLF6 that regulate and prevent an over activation of the proinflammatory response so that the commensal is not cleared completely by the host.

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to the rest of the samples while green indicates that the gene’s signal is lower relative to the rest of the samples.

**Q-RT-PCR**

TaqMan Low Density Array cards were designed to validate a subset of genes that were differentially expressed in the array experiments (Applied Biosystems, USA). Total RNA (0.3 μg) was converted to cDNA using a High-Capacity cDNA archive kit (Applied Biosystems, USA). The Q-RT-PCR reactions were run on an ABI 7900 system (Applied Biosystems, USA). Data was analysed using the SDS2.2 software where baseline and threshold settings were automatically adjusted. Relative gene expression levels were obtained using the $2^{-\Delta\Delta CT}$ method [65] with the 18S housekeeping gene used for normalisation. In brief, this method uses a single sample, termed the “calibrator sample,” as a comparator for every unknown sample’s gene expression level. In this case, we chose a mock-infected sample at 0 hour time point to be the calibrator sample. The calibrator is analysed on every assay plate with the unknown samples of interest. The relative quantification (RQ) value is calculated using the following formula: $RQ = 2^{-\Delta CT}$, where $\Delta CT = (CT_{\text{gene of interest}} - CT_{18S})$ in unknown sample $- (CT_{\text{gene of interest}} - CT_{18S})$ in calibrator (CT of gene of interest in calibrator - CT of 18S gene in calibrator). Box plots indicate the median RQ value within a group of biological replicates.

The response of human epithelial cell genes to secreted neisserial proteins were determined by Q-RT-PCR using the Fluidigm chip technology (Fluidigm, USA). Briefly, 0.5 μg total RNA was converted to cDNA using the High-Capacity cDNA archive kit. From this, the cDNA was preamplified using the TaqMan PreAmp master mix (Applied Biosystems, USA). The cDNA, TaqMan universal PCR master mix, DA sample loading reagent, TaqMan gene primers and DA assay loading reagent were mixed accordingly and loaded into a stabilised integrated fluidic circuit controller following manufacturer’s instructions. The Q-RT-PCR reactions were read using the Biomark data collection machine (Fluidigm, USA). Data was analysed using the Biomark Real-Time PCR analysis software (Fluidigm, USA). Relative gene expression levels were obtained after normalisation to 18S rRNA. Relative quantification levels are obtained with respect to mock-infected sample at 0 hour time point.

**ELISAs**

After treatment of cells with either whole bacteria or bacterial secreted proteins, supernatants were harvested, centrifuged for 10 minutes at 3220 g and passed through 0.2 μm filters to remove the bacteria. PI3 and TNF-α measurements were performed according to instructions in the ELISA kits (R&D systems and Bio-Rad, USA).

**Neisserial secreted proteins**

*N. meningitidis* and *N. lactamica* were grown in RPMI media (Sigma, UK) supplemented with amino acids when necessary [66]. Crude neisserial secreted proteins were harvested as described in Robinson et al. [20] with some modifications. Briefly, the bacteria were grown at 37°C with shaking at 200 rpm. Bacterial supernatants were harvested at log phase and ultracentrifuged at 40,000 g for 1 hour. The sample was then concentrated 20 times with a 10 kDa molecular weight cut off using Vivaspin ultrafiltration spin columns (Sartorius Stedim, Germany). Endotoxin removal was carried out using the Detoxi-Gel Endotoxin Removing Gel consisting of immobilised polymyxin B (Pierce, Thermo Fisher Scientific, USA) until there was less than 0.1 EU/ml, as measured by LAL assay (Lonza, USA).

**Statistical analysis**

For the association and invasion assays, as well as the microarray validation by Q-RT-PCR, mean values were used. Significance of difference was determined using the Student’s T-test (assuming unequal variance), whereby $p$-values of less than 0.05 were considered to represent significance.

The expression of selected genes that were studied in detail over a time course at the transcript and protein level were analysed by SPSS11 (IBM, USA) software. Box plots representing the interquartile range, the median and the highest and lowest values among the biological replicates were drawn. The Mann–Whitney test was used to compare between conditions to determine if the expression of a gene was significantly different (with a $p$-value of less than 0.05).

**Supporting Information**

Table S1 Microarray and Q-RT-PCR results showing host gene expression fold changes from 5 to 7 hours in response to live WT *N. lactamica* and *N. meningitidis*.

(DOC)

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**Author Contributions**

Conceived and designed the experiments: HEEW M-SL MLH PRL. Performed the experiments: HEEW M-SL. Analyzed the data: HEEW M-SL MLH. Contributed reagents/materials/analysis tools: MLH PRL. Wrote the paper: HEEW M-SL JSK MLH PRL.

**References**


