Relative Contribution of P5 and Hap Surface Proteins to Nontypable Haemophilus influenzae Interplay with the Host Upper and Lower Airways

Euba, B., Moleres, J., Viadas, C., Ruiz de Los Mozos, I., Valle, J., Bengoechea, J. A., & Garmendia, J. (2015). Relative Contribution of P5 and Hap Surface Proteins to Nontypable Haemophilus influenzae Interplay with the Host Upper and Lower Airways. PloS one, 10(4), [0123154]. DOI: 10.1371/journal.pone.0123154

Published in: PloS one

Document Version: Publisher's PDF, also known as Version of record

Queen's University Belfast - Research Portal:
Link to publication record in Queen's University Belfast Research Portal

Publisher rights
© 2015 Euba et al.
This is an open access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

General rights
Copyright for the publications made accessible via the Queen's University Belfast Research Portal is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The Research Portal is Queen's institutional repository that provides access to Queen's research output. Every effort has been made to ensure that content in the Research Portal does not infringe any person's rights, or applicable UK laws. If you discover content in the Research Portal that you believe breaches copyright or violates any law, please contact openaccess@qub.ac.uk.
Relative Contribution of P5 and Hap Surface Proteins to Nontypable \textit{Haemophilus influenzae} Interplay with the Host Upper and Lower Airways

Begoña Euba\textsuperscript{1,2\*}, Javier Moleres\textsuperscript{1,2\*}, Cristina Viadas\textsuperscript{1,2,3}, Igor Ruiz de los Mozos\textsuperscript{2}, Jaione Valle\textsuperscript{2}, José Antonio Bengoechea\textsuperscript{1,3,4}, Junkal Garmendia\textsuperscript{1,2,3\*}

\textsuperscript{1}Centro de Investigación Biomédica en Red de Enfermedades Respiratorias (CIBERES), Madrid, Spain, \textsuperscript{2}Instituto de Agrobiotecnología, CSIC-Universidad Pública Navarra-Gobierno Navarra, Mutilva, Spain, \textsuperscript{3}Laboratory Microbial Pathogenesis, Fundación Investigación Sanitaria Illes Balears (FISIB), CSIC-Govern Illes Balears, Bunyola, Spain, \textsuperscript{4}Centre for Infection and Immunity, Queen’s University Belfast, Belfast, United Kingdom

\textsuperscript{\*}These authors contributed equally to this work.

\textsuperscript{\*}juncal.garmendia@unavarra.es

Abstract

Nontypable \textit{Haemophilus influenzae} (NTHi) is a major cause of opportunistic respiratory tract disease, and initiates infection by colonizing the nasopharynx. Bacterial surface proteins play determining roles in the NTHi-airways interplay, but their specific and relative contribution to colonization and infection of the respiratory tract has not been addressed comprehensively. In this study, we focused on the \textit{ompP5} and \textit{hap} genes, present in all \textit{H. influenzae} genome sequenced isolates, and encoding the P5 and Hap surface proteins, respectively. We employed isogenic single and double mutants of the \textit{ompP5} and \textit{hap} genes generated in the pathogenic strain NTHi375 to evaluate P5 and Hap contribution to biofilm growth under continuous flow, to NTHi adhesion, and invasion/phagocytosis on nasal, pharyngeal, bronchial, alveolar cultured epithelial cells and alveolar macrophages, and to NTHi murine pulmonary infection. We show that P5 is not required for bacterial biofilm growth, but it is involved in the NTHi-airways interplay, but their specific and relative contribution to colonization and infection of the respiratory tract has not been addressed comprehensively. In this study, we focused on the \textit{ompP5} and \textit{hap} genes generated in the pathogenic strain NTHi375 to evaluate P5 and Hap contribution to biofilm growth under continuous flow, to NTHi adhesion, and invasion/phagocytosis on nasal, pharyngeal, bronchial, alveolar cultured epithelial cells and alveolar macrophages, and to NTHi murine pulmonary infection. We show that P5 is not required for bacterial biofilm growth, but it is involved in NTHi interplay with respiratory cells and in mouse lung infection. Mechanistically, P5\textsubscript{NTHi375} is not a ligand for CEACAM1 or α5 integrin receptors. Hap involvement in NTHi375-host interaction was shown to be limited, despite promoting bacterial cell adhesion when expressed in \textit{H. influenzae} RdKW20. We also show that Hap does not contribute to bacterial biofilm growth, and that its absence partially restores the deficiency in lung infection observed for the Δ\textit{ompP5} mutant. Altogether, this work frames the relative importance of the P5 and Hap surface proteins in NTHi virulence.
Introduction

Nontypable (non-capsulated) *Haemophilus influenzae* (NTHi) is a Gram negative coccobacillus that is a common commensal in the nasopharynx of both children and adults, and also an important cause of localized respiratory tract infections such as acute otitis media, otitis media with effusion, community-acquired pneumonia, and exacerbations of chronic bronchitis and chronic obstructive pulmonary disease (COPD) [1]. Current evidence indicates that NTHi is highly adapted to the human airways [2]. NTHi interplay with host extracellular matrix (ECM) proteins and cell surfaces is facilitated by several proteinaceous adhesins, including the P5 and Hap outer membrane proteins (OMPs) [3]. The *ompP5* and *hap* genes are present in all *H. influenzae* isolates sequenced to date [4]. The *ompP5* gene encodes P5, a major outer membrane protein predicted to have hypervariable domains (Fig 1A) [5,6]. P5 has been shown to be an adhesin to human oropharyngeal cells [7], mucin [8], chinchilla eustachian tube mucus [9], and respiratory syncytial virus infected type II pneumocytes [10]. P5 levels seem to be preserved on NTHi biofilms compared to planktonically grown cells [11]. P5 is required for resistance of NTHi to the classical and alternative complement pathways [12], and belongs to a set of virulence genes required in nonvirally infected mice [13]. Depending on the strain, P5 may be important for optimal NTHi growth in rich medium [12], and it may be a bacterial ligand for the carcinoembryonic antigen-related cell adhesion molecule 1 (CEACAM1) [14], playing then a role in nasopharynx colonisation in the chinchilla model [15,16]. NTHi stimulates the expression of intercellular adhesion protein 1 (ICAM-1) on respiratory epithelial cells [17], and P5 has been shown to be a ligand for ICAM-1 [18].

Hap (*Haemophilus adherence and penetration*) is a monomeric self-associating autotransporter (SAAT) with homology to serine-type IgA1 proteases, identified as a bacterial factor promoting interaction with respiratory epithelial cells [3,19]. Hap contains a signal peptide (SP), a passenger domain (Hap,) and a β-barrel domain (Hapβ) [3]. Hap, harbours a C-terminal adhesive region that promotes adherence to human epithelial cells [20], to ECM proteins [21,22], and to neighbouring Hap-expressing bacteria [20], and a N-terminal serine protease domain that modulates Hap autoproteolytic cleavage [23] (Fig 1A). Hap autoproteolytic activity is inhibited by secretory leukocyte protease inhibitor (SLPI), a serine proteinase inhibitor found in human respiratory secretions whose activity results in enhanced Hap adhesive activity [24]. Hap seems to be associated with bacteria within the biofilm, and to be present in the biofilm extracellular matrix [25]. Mutation of a repertoire of genes encoding enzymes involved in the synthesis of the lipooligosaccharide (LOS) core results in decreased *hap* transcription [26].

Despite the wealth of evidence on P5 and Hap, there is not a systematic characterization of the relative contribution of these adhesins to NTHi virulence. In this study, we hypothesised that P5 and Hap may have a differential contribution to NTHi-host interplay through the human respiratory tract. We employed NTHi strain 375 (hereafter NTHi375) to generate single and double mutants defective in the *ompP5* and/or *hap* genes. We evaluated the effect of their disruption on NTHi biofilm growth, on bacterial interaction with a panel of cultured respiratory cells including nasal, pharyngeal, bronchial and alveolar cells, and on mouse respiratory infection. This work allowed us to assign specific roles to P5 and Hap OMPs, providing a context for their relative importance to a range of phenotypic traits and a greater understanding of their contribution to NTHi interplay with the host airways.

Methods

**Bacterial strains, media and growth conditions**

Strains used in this study are described in Table 1. NTHi strains were grown (overnight, 37°C, 5% CO₂) on Chocolate-agar (Biomérieux) or on brain-heart infusion (BHI)-agar plates.
Fig 1. Representation of predicted P5 and Hap structural domains, bacterial growth and biofilm formation. (A) Schematic representation of P5 and Hap surface proteins for NTHi375, predicted structural domains and amino acid polymorphisms. Multiple sequence alignments for NTHi375 P5 and Hap were performed in Muscle [49], by using all NCBI available NTHi orthologous proteins. Domain structures were predicted by using the conserved domain database by Marchler-Bauer and co-workers [50], and designated based on previous experimental findings and in agreement with the NCBI conserved domain database. P5 shows four hypervariable regions at the putative surface-exposed loops in the transmembrane domain [5,6,51]. Hap contains a N-
supplemented with 10 μg/ml hemin and 10 μg/ml nicotinamide adenine dinucleotide (NAD), referred to as sBHI. NTHi liquid cultures were grown in sBHI (37°C, 5% CO₂). Erythromycin 10 μg/ml (Erm₁₀), kanamycin 11 μg/ml (Km₁₁), or chloramphenicol 1 μg/ml (Cm₁) were used when required.

*Escherichia coli* was grown on Luria Bertani (LB) broth or LB-agar plates at 37°C, supplemented with ampicillin 100 μg/ml (Amp₁₀₀), erythromycin 150 μg/ml (Erm₁₅₀), kanamycin 50 μg/ml (Km₅₀), or chloramphenicol 30 μg/ml (Cm₃₀), when necessary.

To monitor growth, NTHi strains grown on chocolate agar for 24 h were inoculated (2 to 5 colonies) in 20 ml sBHI and incubated for 11 h under shaking. Cultures were then diluted 1:80 in sBHI and incubated for 9 h under the same conditions. OD₆₀₀ was recorded hourly.

NTHi strain 375 (NTHi375) is an OM isolate [4,27,28]; NTHi375Δhap has been described previously [29]. The ompP5 gene and its respective adjacent regions, was amplified by PCR with Taq polymerase (Promega) using NTHi375 genomic DNA as template and primers ompP5-F₁ (5´-AGCCAGACTTAATCTATCCGAATAATTGT) and ompP5-R₁ (5´-TTGCGGGTTTTATTTTTCCACTGTGATTAA). The gene-containing fragment was cloned into pGEM-T Easy (Promega), generating pGEMT/ompP5. Cloned PCR product was disrupted by inverse PCR with Vent polymerase (New England Biolabs), using primers ompP5-F₂ (5´-ACCAATGGCTAACTCGCGTAGGTAAATACC) and ompP5-R₂ (5´-CTGCGTATTCTGCACCTACTGCAAATAAAC). An internal 30-bp fragment (nucleotides 521 to 550 in the ompP5 coding sequence) was replaced by a blunt-ended (excised by *Sma*I) erythromycin resistance cassette from pBSLerm [30], terminal signal peptide, a passenger Hapₜ and a β-barrel Hapₚ domain. Hap alignment exhibits a hypervariable amino acid mosaic-like pattern [52]. Hapₜ contains a conserved serine protease domain with the catalytic triad site His₉₈, Asp₁₄₀ and Ser₂₄₄ (GDSGS₂₄₆), which mediates autoproteolysis. Predicted cleavage positions are denoted with arrows; dotted arrows show degenerated consensus cleavage sites. Similar residues are represented in white (>90% similarity); variable residues are shaded in gray (>80% similarity); hypervariable residues are shaded in black (<80% similarity).

![Growth curve of NTHi375 wild-type (circle), ΔompP5 (triangle), Δhap (square) and ΔompP5Δhap (inverted triangle) strains.](image1)

![Biofilm formation under continuous-flow conditions, in microfermenters containing glass slides where bacteria formed the biofilm. Images of a representative experiment for NTHi375 (a), ΔompP5 (b), Δhap (c) and ΔompP5Δhap (d) strains.](image2)

Table 1. Strains and plasmids used in this study.

| Strain | Description | Reference |
|--------|-------------|-----------|
| **H. influenzae** | | |
| NTHi375 | Wild-type, otitis media clinical isolate | [27] |
| 375Δhap | hap::ermC, ErmR | [29] |
| 375ΔompP5 | ompP5::ermC, ErmR | This study |
| 375ΔhapΔompP5 | hap::ermC, ompP5::km, ErmR KmR | This study |
| RdKW20 | Laboratory strain, capsule-deficient serotype d | [44] |
| RdKW20ΔompP5 | ompP5::ermC, ErmR | This study |
| **E. coli** | | |
| CC118 | Used for cloning assays | |
| **Plasmid** | | |
| pGEM-T Easy | Used for cloning assays | Promega |
| pGEM-T/ompP5 | ompP5 from NTHi375 and flanking regions cloned into pGEM-T Easy | This study |
| pBSLerm | Source of an ErmR cassette | [30] |
| pUC4K | Source of a KmR cassette | Addgene |
| pALG-2 | pGEM-T/ompP5 derivative where ompP5 is disrupted by an ErmR cassette | This study |
| pALG-3 | pGEM-T/ompP5 derivative where ompP5 is disrupted by an KmR cassette | This study |
| pSU20 | Cloning vector with a p15A replication origin, CmR | [32] |
| pSU20- Pr::hapNTHi375-HA | pSU20 derivative with a 4.6-kb insert containing hap from NTHi375 expressed under its own promoter | This study |

doi:10.1371/journal.pone.0123154.g001

doi:10.1371/journal.pone.0123154.t001

doi:10.1371/journal.pone.0123154.l001
generating plasmid pALG-2 (Table 1). This plasmid was digested with NotI to obtain a linear disruption cassette for ompP5, that was used to transform NTHi375 using the MIV method [31]. Transformants were screened by plating bacteria on sBHI-agar with Erm\(_{10}\), to obtain NTHi375\(\Delta\)ompP5. Same approach and disruption cassette was used to generate \(H.\) influenzae (Hi) RdKW20\(\Delta\)ompP5 mutant strain. The NTHi375\(\Delta\)ompP5\(\Delta\)hap double mutant was generated by two successive recombination events. First, strain NTHi375\(\Delta\)hap was generated as described previously [29], and it was then transformed with a disruption cassette for ompP5. This cassette was generated by cloning a blunt-ended kanamycin resistance gene released from pUC4K by digestion with HincII in a vector obtained by inverse PCR of pGEM-T/ompP5 with primers ompP5-F2/ompP5-R2. The resulting plasmid, pAGL-3, was digested with NotI, creating a 3.8 kb linear fragment containing the disruption cassette \(\text{ompP5:\text{KmR}}\), used to transform NTHi375\(\Delta\)hap. Double recombination events were selected on sBHI-agar with Erm\(_{10}\) and Km\(_{11}\).

Expression of Hap\(\text{NTHi375-HA}\) in \(H.\) influenzae RdKW20

The \(hap\) gene and its upstream 500 bp region, was amplified by PCR with Phusion High-Fidelity DNA Polymerase (Thermo Fisher Scientific) using NTHi375 genomic DNA as template, and primers hap-F1 (5´-ACTATCTGTCGTCATGGAACACAATCTTGAT)/hap-R1 (5´-TTAAGCGTACCGGACGCTGATTTTACCAACGATACCCAAATTTGACGCCCA). The hap-R1 reverse primer was used to introduce an HA tag coding sequence at the 3´end of the \(hap\) gene. This 4,677 bp blunt PCR product was phosphorylated with T4 kinase (New England Biolabs), and cloned into pSU20 [32], previously digested with HincII and dephosphorylated with antarctic phosphatase (New England Biolabs), generating pSU20-\(Pr::hap\)\(\text{NTHi375-HA}\). pSU20 and pSU20-\(Pr::hap\)\(\text{NTHi375-HA}\) were transformed into electrocompetent Hi RdKW20. Transformations were selected on sBHI-agar with Cm\(_1\). Hi RdKW20, RdKW20 (pSU20) and RdKW20 (pSU20-\(Pr::hap\)\(\text{NTHi375-HA}\)) whole cell extracts were prepared from cultures grown to OD\(_{600}\) = 0.9 in sBHI containing Cm\(_1\), when required. Hap\(\text{NTHi375-HA}\) expression was analyzed by western blot with a primary rabbit anti-HA antibody (Sigma) diluted 1:4000, and a secondary goat anti-rabbit IgG (whole molecule, Sigma) antibody conjugated to horseradish peroxidase, diluted 1:1000.

Biofilm formation

Biofilm formation under flow conditions was assessed using 60 ml microfermenters (Pasteur Institute, Laboratory of Fermentation) with a continuous flow of medium (40 ml/h) and constant aeration with sterile compressed air (0.3 bar). Submerged Pyrex glass slides served as the growth substratum. Three to four colonies of each strain grown on chocolate-agar were inoculated in 20 ml sBHI and incubated under shaking up to OD\(_{600}\) = 1. Approximately 5x10\(^8\) bacteria from this culture were used to inoculate each microfermenter, which was then run at 37°C for 16 h. The viability on each inoculated bacterial aliquot was tested by serial dilution and plating on sBHI-agar. Biofilm development was recorded with a Nikon Coolpix 950 digital camera. To quantify the biofilm formed, bacteria adhered to the Pyrex slides were resuspended in 10 ml PBS, and OD\(_{600}\) of the suspensions was determined. The viability on the bacteria recovered from the biofilm was tested by serial dilution and plating on sBHI-agar. Experiments were performed in duplicate on at least three independent occasions (n≥6).

Cell culture and bacterial infection

RPMI 2650 human nasal epithelial cells (ATCC CCL-30) were maintained in DMEM with HEPES 10mM, 10% heat inactivated foetal calf serum (FCS) and antibiotics (penicillin and streptomycin) in 75 cm\(^2\) tissue culture flasks at 37°C with 5% CO\(_2\). Cells were seeded to 5x10\(^4\)
cells/well in 24-well plates 24 h before infection. Detroit 562 human pharynx epithelium (ATCC CCL-138) was maintained as RPMI 2650 cells, and seeded to 2x10^5 cells/well 48 h before infection. NCI H-292 mucouspidermoid pulmonary human carcinoma epithelial cells (ATCC CR-L-1848) were maintained in RPMI 1640 with Hepes 10mM, 10% FCS and antibiotics, and seeded to 4x10^5 cells/well 24 h before infection. A549 human carcinomic alveolar basal epithelial cells (ATCC CCL-185) were maintained as described before [33]. Cells were seeded to 6x10^4 cells/well for 32 h, and serum-starved 16 h before infection. HeLa-BGP is a stably transfected HeLa cell line expressing hCEACAM1-4L receptor [34]. CEACAM1 expression has been previously tested in this cell line, under conditions identical to those used in this study [5]. HeLa-BGP cells were propagated as shown before [34], and seeded to 4x10^5 cells/well 24 h before infection. Murine alveolar macrophages MH-S (ATCC, CRL-2019) were grown on RPMI 1640 with Hepes 10mM, 10% FCS and antibiotics, and seeded to 7x10^5 cells/well 16 h before infection.

For NTHi infection, we used previously set up conditions [33,35,36]. RPMI 2650, Detroit 562, NCI H-292, A549, HeLa and HeLa-BGP cells were infected in 1 ml EBSS (Earle’s Balanced Salt Solution, Gibco) to get a multiplicity of infection (MOI) of ~100:1. MH-S cells were infected in 1 ml RPMI 1640 with Hepes 10mM and 10% FCS to get a MOI of ~100:1. To monitor adhesion, RPMI 2650, Detroit 562, NCI H-292, A549, HeLa and HeLa-BGP cells were infected for 30 min, and MH-S cells were infected for 1 h. Although this assay does not completely exclude a possible internalization of some bacteria, experimental conditions were previously set to monitor adhesion ([33,36], data not shown). Cells were then washed 5 times with PBS, lysed with 300 μl of PBS-saponin 0.025% for 10 min at room temperature, and serial dilutions were plated on sBHI-agar. For invasion, all epithelial cells were infected for 2 h, washed 3 times with PBS, incubated for 1 h with RPMI 1640 containing 10% FCS, Hepes 10mM and gentamicin 200 μg/ml. For phagocytosis, MH-S cells were infected for 1 h, washed 3 times with PBS, and incubated for 1 h with medium containing gentamicin 300 μg/ml. In all cases, cells were washed 3 times with PBS and lysed as described above. Bacterial adherence to the wells was excluded by monitoring infection of the panel of cell lines by microscopy (data not shown).

When indicated, A549 cells were pretreated for 16 h with 2 μg/ml tunicamycin (Sigma). This treatment did not induce cytotoxicity, verified by measuring the release of lactate dehydrogenase and microscopy (data not shown). Drug exposure was maintained during bacterial contact. Drug exposure had no effect on bacterial viability under the conditions tested (data not shown). α5 integrin analysis was performed by (i) cell incubation with anti-α5 P1D6 (20 μg/ml) function blocking antibody, for 1 h before infection. The antibody was kept during infection; (ii) cell incubation with RGD peptide (10 μM), added to the cells 1 h before infection, and removed before infection. All infections were carried out in triplicate at least three independent times (n>9).

**Mouse assays**

These assays were carried out as previously described [29]. CD1 female mice (4 to 5 weeks old) were purchased from Charles River Laboratories (France) and housed under pathogen-free conditions at the Institute of Agrobiotechnology of the Universidad Pública de Navarra (UPNA) facilities (registration number ES/31-2016-000002-CR-SU-US). Animal handling and procedures were in accordance with the current European (Directive 86/609/EEC) and National (Real Decreto 53/2013) legislations, following the FELASA and ARRIVE guidelines, and with the approval of the UPNA Animal Experimentation Committee (Comité de Ética, Experimentación Animal y Bioseguridad-CEEAB, http://www.unavarra.es/invest/comiteEtica.htm) and the local Government authorization. Animal’s condition was monitored daily. For
infection, bacteria were recovered with 1 ml PBS from chocolate-agar grown for 16 h, to obtain a suspension of \(\approx 5 \times 10^9\) c.f.u./ml. Before infection, mice were anesthetized intraperitoneally with a mixture of ketamine-xylazine (3:1). Each mouse received 20 μl of inoculum (\(\approx 10^8\) c.f.u.) intranasally. During infection, we did not observe signs of sickness or alterations in behavior, and the use of humane endpoints was not required. Groups of at least 5 mice were euthanized by cervical dislocation and necropsied at selected intervals to determine the number of c.f.u. per lung. Lungs were aseptically removed, individually weighed in sterile bags (Stomacher80, Seward Medical), homogenized, and serially ten-fold diluted in PBS. Each dilution was spread on sBHI-agar (detection limit <10 c.f.u./lung).

**Statistical analysis**

For biofilm growth, cell infection and mice infection assays, mean±SD were calculated and statistical comparison of means performed using the two-tail \(t\) test. In all cases, a \(p<0.05\) value was considered statistically significant. Analyses were performed using Prism software, version 4 for PC (GraphPad Software) statistical package.

**Results**

**Construction of NTHi375 mutant strains defective in the surface proteins P5 and Hap**

P5 and Hap are two NTHi OMPs encoded by the ompP5 and hap genes, respectively, present in all NTHi isolates analysed to date [4]. In this study, we employed NTHi375, an isolate used for previous studies on NTHi biology and infection [28,29,33], and whose genome sequence is available [37]. We analysed P5 and Hap amino acid sequence on NTHi375, and their level of conservation (Fig 1A, Fig A and Table A in S1 File).

NTHi375 was used to generate NTHi375ΔompP5, NTHi375Δhap [29] and ΔhapΔompP5 mutant strains (Table 1). The ompP5 and hap gene expression was monitored by RNA extraction and quantitative real-time PCR on those strains. As expected, we could not detect expression of the ompP5 gene in NTHi375ΔompP5 and NTHi375ΔhapΔompP5 strains, neither expression of the hap gene in NTHi375Δhap and NTHi375ΔhapΔompP5 strains (data not shown). These mutants did not exhibit growth defects compared to the wild-type strain when OD\(_{600}\) was monitored over time during growth in sBHI liquid culture (Fig 1B).

**P5 and Hap surface proteins are not required for NTHi biofilm growth**

Given that P5 levels have been shown to be preserved on NTHi biofilms [11], and that Hap can be detected in the biofilm extracellular matrix [25], we assessed P5 and Hap contribution to NTHi biofilm growth by using an *in vitro* model system based on the formation of biofilm communities by NTHi375 under continuous-flow culture conditions in microfermenters [29,38]. We separately inoculated comparable c.f.u. numbers for each strain, and monitored biofilm development in microfermenters by measuring the turbidity of bacterial suspensions detached from removable glass slides after 16 h. OD\(_{600}\) of bacterial suspensions detached from glass slide was similar for the four strains tested, wild-type (OD\(_{600}\) = 1.19±0.42), ΔompP5 (OD\(_{600}\) = 1.26±0.32), Δhap (OD\(_{600}\) = 1.24±0.35), and ΔompP5Δhap (OD\(_{600}\) = 1.49±0.35). Representative images are shown in Fig 1C. Biofilm viability was quantified by serial dilution plating of bacterial suspensions detached from the removable glass slides, which rendered similar numbers among strains (~10\(^{10}\) c.f.u./suspension, i.e. /biofilm).

In sum, under the conditions tested, NTHi375ΔompP5, Δhap and ΔompP5Δhap mutants formed biofilms similar to those observed for the wild-type strain.
Role of P5 and Hap in NTHi interplay with upper and lower airways epithelial cells

To assess the role of P5 and Hap in NTHi interaction with the human airways epithelia, we used nasal RPMI 2650, pharynx Detroit 562, bronchial NCI H-292 and A549 type II pneumocytes cultured epithelial cells. First, we established infection levels for NTHi375 wild-type strain in the four cell types. NTHi375 adhered to the four cell types with variable numbers. Adhesion to nasal cells was significantly lower than that observed for the other cell types tested (mean numbers for RPMI 2650 were lower than those obtained for Detroit 562 and H-292 (p<0.0001), and for A549 (p<0.05) cells); adhesion to Detroit 562 was significantly lower than to NCI H-292 (p<0.0001), but higher than to A549 (p<0.0001) cells; adhesion to NCI H-292 was higher than that observed for the other cell types tested (p<0.0001) (Fig 2A). In terms of NTHi375 internalization into the four cell types, the highest invasion was found for NCI H-292 (p<0.0001) cells, followed by pharynx, nasal and alveolar cells, respectively. Thus, NTHi375 invasion of RPMI 2650 cells was lower than that obtained for Detroit 562 (p<0.05) and H-292 (p<0.0001) cells, and higher than that shown by A549 (p<0.05) cells; invasion of Detroit 562 was lower than that obtained for H-292 (p<0.0001), and higher than the one measured for A549 (p<0.0005) cells (Fig 2A).

Next, we tested the ability of NTHi375ΔompP5, Δhap and ΔompP5Δhap mutants to infect the four cell types. NTHi375ΔompP5, Δhap and ΔompP5Δhap adhesion to RPMI 2650 cells was lower than that displayed by the wild-type strain, although the observed decrease was significant only for NTHi375ΔompP5Δhap (p<0.05). Invasion of the wild-type strain was significantly higher than that observed for ΔompP5 (p<0.0005), Δhap (p<0.05) and ΔompP5Δhap (p<0.0005) mutants (Fig 2B). NTHi375ΔompP5 and ΔompP5Δhap mutants adhered to Detroit 562 cells at a level similar to the wild-type strain. Conversely, adhesion of NTHi375Δhap to Detroit 562 cells was higher (p<0.05) than that displayed by the wild-type strain. Invasion of NTHi375ΔompP5 into Detroit 562 cells was significantly lower (p<0.05) than that displayed by the wild-type strain. NTHi375Δhap and ΔompP5Δhap mutants presented Detroit 562 cell invasion levels similar to the wild-type strain (Fig 2C). We also tested the role of P5 and Hap in NTHi375 interplay with NCI H-292 bronchial cells. Adhesion of NTHi375ΔompP5 and ΔompP5Δhap, but not NTHi375Δhap, to NCI H-292 cells was lower than that displayed by the wild-type strain (p<0.005 and p<0.0001, respectively). In agreement with adhesion data, NTHi375ΔompP5 and ΔompP5Δhap mutants invaded NCI H-292 cells to a lower extent than the wild-type strain (p<0.0001), but NTHi375Δhap entered NCI H-292 at the same level as the wild-type strain (Fig 2D). Finally, adhesion of NTHi375ΔompP5 and ΔompP5Δhap to A549 cells was lower than that displayed by the wild-type strain (p<0.001 and p<0.05, respectively). Similarly, single and double mutant strains lacking ompP5 invaded A549 cells significantly less than the wild-type strain (p<0.0001) (Fig 2E). To further confirm P5 involvement in H. influenzae epithelial infection, we compared cell infection by H. influenzae RdKW20 wild-type and ΔompP5 strains. We observed P5RdKW20 implication in RdKW20 infection of NCI H-292 (adhesion, p<0.05; invasion, p<0.001) and A549 (adhesion, p<0.005; invasion, p<0.01) cells (Fig 2F).

 Altogether, these results suggest a differential contribution of P5 and Hap to NTHi interface with the airways epithelial cells. P5 and Hap are involved in NTHi375 interplay with RPMI 2650 nasal cells. P5 seems to play a role in bacterial entry into Detroit 562 cells, and it is required for NTHi375 interplay with NCI H-292 bronchial and A549 alveolar cells, at both the adhesion and invasion levels. Hap does not participate in the NTHi-bronchial/alveolar cell interface, and the hap gene deficiency may favour NTHi375 interplay with Detroit 562 pharynx epithelial cells.
Fig 2. Infection of respiratory epithelial cells by NTHi375 mutant strains lacking the \textit{ompP5} and \textit{hap} genes. Bacterial adhesion is shown in the left and invasion in the right panels, in all sections of this figure. (A) NTHi375 interplay with upper and lower human epithelial airways. NTHi375 adhesion to- and invasion of nasal RPMI 2650, pharynx Detroit 562 and bronchial H-292 epithelia and type II pneumocytes A549. Effect of \textit{ompP5} and \textit{hap} gene disruption on NTHi interplay with RPMI 2650 nasal (B), Detroit 562 pharynx (C), NCI H-292 bronchial (D), and A549 type II alveolar (E) epithelial cells. NTHi375, \textit{ΔompP5}, \textit{Δhap} and \textit{ΔompP5Δhap} strains were used to assess bacterial adhesion and invasion. (F) Effect of \textit{ompP5} gene disruption on Hi RdKW20 interplay with NCI
P5 and Hap do not contribute to NTHi375 epithelial adhesion mediated by CEACAM1, α5 integrin or N-glycosylation

It has been previously reported that, depending on the strain, P5 could be a bacterial ligand for the eukaryotic receptor CEACAM1 [14]. NTHi375 adhesion assays were performed by using HeLa-BGP (biliary glycoprotein or CD66a, currently known as CEACAM1) cells [34], a HeLa derivative cell line stably expressing hCEACAM1-4L [5], and previously used to assess the impact of CEACAM1 on bacterial infections [5,39]. NTHi375 showed similar adhesion to HeLa control and HeLa-BGP cells, excluding a potential role for CEACAM1 in NTHi375 epithelial adhesion. Adhesion of NTHi375ΔompP5 and ΔompP5Δhap was lower than that displayed by the wild-type strain, to the same extent for both for HeLa (p<0.0001 and p<0.0005, respectively) and HeLa-BGP (p<0.0005) cells (Fig 3A). These results suggest that NTHi375 does not interact with CEACAM1, that P5NTHi375 is not likely to be a ligand for CEACAM1, and that P5, but not Hap, participates in NTHi375 adhesion to HeLa epithelial cells.

To further validate the use of HeLa-BGP cells, we took advantage of the previously shown interaction between P5RdKW20 and CEACAM1 [14]. As expected, Hi RdKW20 was shown to adhere better to HeLa-BGP than to HeLa cells (p<0.0001), and Hi RdKW20ΔompP5 showed lower adhesion to HeLa-BGP cells than the wild-type strain, (p<0.0005) (Fig 3B). Of note, Hi RdKW20ΔompP5 also showed to adhere better to HeLa-BGP than to HeLa cells (p<0.0001).

We have previously shown that α5β1 integrin subunit is implicated in NTHi A549 cell infection [35]. The bacterial ligand involved in this process is unknown, but it is likely to be a surface protein encompassing a RGD domain. Given that P5 and Hap do not contain a RGD domain in their respective amino acid sequences, these proteins are unlikely to be NTHi ligands for α5 integrin. As expected, NTHi375ΔompP5, Δhap and ΔompP5Δhap adhesion to A549 cells decreased in the presence of the anti-integrin α5 P1D6 blocking antibody, and of a synthetic peptide containing a RGD-sequence mimicking the physiological α5 integrin ligand fibronectin, compared to control untreated cells, same as observed for the wild-type strain (Fig 3C–3F). N-glycosylation of the α5β1 integrin receptor has been shown to be essential for association of the heterodimer subunits, and for its optimal binding to fibronectin [40]. Moreover, host cell surface glycoprotein structures may play a role in NTHi attachment to Chang epithelial cells [41]. We assessed the relevance of cell N-glycosylation on NTHi epithelial adhesion by A549 cell treatment with the N-linked glycosylation blocking agent tunicamycin. NTHi375 wild-type, ΔompP5, Δhap and ΔompP5Δhap showed a reduced adhesion to epithelial cells in the presence of tunicamycin, compared to control untreated cells. In all cases, bacterial adhesion to anti-α5 P1D6 antibody, RGD peptide or tunicamycin-treated cells was lower (p<0.0001) than to control untreated cells (Fig 3C–3F). These results suggest that NTHi375 infection of A549 cells may involve bacterial interaction with N-glycosylated residues on the host cell surface, but P5 and Hap are unlikely to participate in such interaction.

Role of P5 and Hap in NTHi interaction with alveolar macrophages

The lung contains alveolar macrophages which are both sentinels and the first line of defence against infection [42]. Clearance of NTHi from lungs depends on the efficiency of host phagocytes to recognise and destroy the pathogen, as we have previously described [36]. We next investigated the ability of MH-S alveolar macrophages to engulf NTHi375ΔompP5, Δhap and ΔompP5Δhap mutant strains. Adhesion of NTHi375ΔompP5 and ΔompP5Δhap to MH-S cells
was lower than that displayed by the wild-type strain (p<0.005 and p<0.05, respectively). Differently, adhesion of NTHi375Δhap to MH-S cells was similar to the wild-type strain (Fig 4A). In agreement with adhesion data, phagocytosis of NTHi375ΔompP5 and ΔompP5Δhap by MH-S cells was significantly lower than that displayed by the wild-type strain (p<0.01 and p<0.005, respectively), and NTHi375Δhap mutant was engulfed by MH-S cells at the same level as the wild-type strain (Fig 4B). These results suggest a relevant role for P5 in NTHi recognition and engulfment by alveolar macrophages, together with a differential contribution of P5 and Hap to NTHi375 interface with this cell type.
Heterologous expression of HapNTHi375 in Hi RdKW20 shows Hap involvement in bacterial adhesion to host cells

Hap involvement in NTHi colonisation of the human airways has been mostly concluded from gain of function studies, based on plasmid- or chromosome expression of the hap gene into Hi RdKW20 [20,21,43]. Under the conditions tested in this study, the hap gene did not show a significant role in NTHi375 interplay with respiratory cells. HapNTHi375 contains a SAAT domain (Fig A in S1 File), and shows sequence conservation on the autoproteolysis regions comprising the canonical catalytic amino acids triad and the previously established cleavage sites (Fig 1A). Therefore, this protein is likely to undergo cleavage of its passenger domain Hapβ. To determine Hap expression and autoproteolysis for NTHi375, the hap gene was HA-tagged, cloned into pSU20 together with its putative promoter region, and expressed in Hi RdKW20, a strain naturally lacking a functional Hap protein due to a stop codon in the hap gene [44]. As expected, the 155-kDa full-length HapNTHi375 protein and the 45-kDa Hapβ preferred cleavage product were detected in whole cell extracts by immunoblot with an anti-HA antibody (Fig 5A). Thus, according to its amino acid sequence, HapNTHi375 is expressed rendering a full length precursor protein and an outer membrane translocator domain Hapβ.
To further confirm HapNTHi375 functionality, we investigated its role in bacterial adhesion to epithelial cells and alveolar macrophages, by comparing the ability of strains Hi RdKW20, Hi RdKW20 (pSU20) and Hi RdKW20 (pSU20-Pr::hapNTHi375-HA) to adhere to NCI H-292, A549 and MH-S cells. Hi RdKW20 (pSU20-Pr::hapNTHi375-HA) adhesion to the three cell types was significantly higher than that displayed by Hi RdKW20 and RdKW20 (pSU20) strains (p < 0.0001, p < 0.0005, p < 0.0001, respectively) (Fig 5B–5D). In conclusion, the hap- NTHi375 gene is likely to express an adhesive protein, as shown by its heterologous expression in the naturally hap deficient strain Hi RdKW20.

Contribution of P5 and Hap to NTHi375 mouse pulmonary infection

We have previously assessed NTHi persistence on a mouse infection model by intranasal inoculation of CD1 mice with NTHi375 [29]. In this study, we sought to determine the impact of P5 and Hap deficiency in mouse lung infection. We quantified bacterial loads for wild-type and each mutant from lung homogenates of infected mice at 24 and 48 h post-infection (PI). We recovered comparable bacterial numbers for NTHi375 wild-type, ΔompP5, Δhap, and ΔompP5Δhap strains at 24 h PI (Fig 6A). Conversely, at 48 h PI, we recovered significantly fewer bacteria for NTHi375ΔompP5 (p < 0.001) and ΔompP5Δhap (p < 0.05) than for the wild-type strain (Fig 6B). NTHi375Δhap delivered counts indistinguishable from those obtained after infection with the wild-type strain. Of note, NTHi375ΔompP5Δhap rendered a significant increase in the number of bacteria recovered, when compared to NTHi375ΔompP5 (p < 0.05). These data indicate that P5 may delay the clearance of NTHi in mouse pulmonary infection.
Hap does not seem to contribute to NTHi persistence in the mouse lung, and its absence partially restores the deficiency in lung infection observed for the NTHi375ΔompP5 mutant strain.

**Discussion**

This study delineates novel roles for P5 and Hap surface proteins in the interplay between NTHi and the respiratory system. Our systematic use of microfermenters, respiratory cultured cells and mouse respiratory infection model systems has allowed for the first time to comprehensively compare single and double P5 and Hap defective mutants generated in a pathogenic genome sequenced strain, NTHi375 [37]. Previous studies separately explored P5, Hap and their roles in NTHi-host interface, by using mutant strains made in different pathogenic strain backgrounds. The ability of NTHi1128ΔompP5 mutant to adhere to epithelial cell surfaces has been evaluated on A549 type II pneumocytes [10], epithelial cells in the chinchilla middle ear [9], and CHO-ICAM-1 transfected cells [18]; the ability of Hi RdKW20ΔompP5 to adhere to CEACAM1 has been evaluated by using CEACAM1-Fc in a receptor overlay assay [14]. Also, high throughput sequencing of transposon insertion sites pointed the importance of ompP5 for *H. influenzae* RdKW20 survival in the mouse lung [13]. Although the ability of NTHiN187Δhap mutant to adhere to epithelial cells has been evaluated by using Chang cells [19], most studies devoted to decipher Hap adhesive properties have been performed by expressing the hap gene in strain DB117, a Hi RdKW20 derivative deficient in recI [20,43].

Our results indicate that P5 is involved in several aspects of the NTHi-host interplay. Although not required for biofilm growth and with a moderate role in NTHi interaction with upper airways cells, NTHi375ΔompP5 was impaired in terms of adhesion and entry into non-phagocytic bronchial and alveolar epithelial cells, and in adhesion to- and uptake by professional phagocytes. Our results suggest that P5 may be a bacterial ligand for host receptor(s) present on the surface of these cell types, although CEACAM1 and α5 integrin are unlikely to be P5 receptors for NTHi375. CEACAM1 has been previously identified as a receptor for several *H. influenzae* strains [45], targeted by P5 or by bacterial ligands alternative to P5 [14]. This was not the case for NTHi375, despite the presence or absence of the ompP5 gene. P5 extracellular loop domains display sequence variability (Table A in S1 File) [6,46], which could account...
for the observed phenotypic heterogeneity among isolates. Of note, our results confirmed that P5\textsubscript{RdKW20} is likely to bind CEACAM1, together with additional currently unknown bacterial ligands [14], and a panel of genome sequenced NTHi clinical isolates tested in terms of interaction with CEACAM1, has rendered a heterogeneous behavior among strains (B. Euba, personal communication), therefore validating the use of HeLa-BGP cells, and supporting the notion of a significant variability among isolates. Previous evidence suggests that P5 could be a bacterial ligand for ICAM-1 [18], and ICAM-1 is a heavily N-glycosylated transmembrane protein [47].

Given that tunicamycin reduced NTHi adhesion to alveolar epithelial cells for both the wild-type and the \textit{ΔompP5} mutant strains, NTHi375 may interact with N-glycosylated molecules at the host cell surface, but P5 is unlikely to be a ligand involved in such interaction. We speculate that the significant \textit{in vivo} clearance observed for NTHi375\textit{ΔompP5} in our respiratory infection model could be a consequence of this mutant impairment to firmly attach to host cell surfaces. The fast lung clearance displayed by NTHi375\textit{ΔompP5} is in agreement with previous observations for intranasal infection of Hi RdKW20 lacking the \textit{ompP5} gene [13].

Hap seemed to play a limited role in most aspects analysed in this work. Hap\textsubscript{NTHi375-HA} detection in whole-cell extracts when expressed in the \textit{hap} naturally lacking strain RdKW20, and the gain of cell adhesive function by RdKW20 when transformed with pSU20-\textit{Pr:\textit{hapNTHi375-HA}}, supported the functionality of the \textit{hap\textsubscript{NTHi375}} gene. However, NTHi375\textit{Δhap} infected bronchial and alveolar epithelial cells, and alveolar macrophages at similar levels than the wild-type strain. We speculate that \textit{hap} deficiency could be compensated by other NTHi surface molecules with potentially redundant function, therefore masking Hap-driven clear cut phenotypes. Hap autoproteolytic activity has been shown to be inhibited by SLPI, which protects the respiratory epithelium from injury due to neutrophil elastase and other proteases involved in acute inflammation [48]. During natural infection, inhibition of Hap autoproteolysis presumably facilitates \textit{H. influenzae} colonisation of the respiratory mucosa, while release of Hap, may result in dispersal and migration from the site of infection. The \textit{hap} gene deficiency would eliminate this predicted Hap involvement in colonisation facilitated by SLPI-dependent inhibition of Hap autoproteolysis. Based on this notion, NTHi375\textit{Δhap} would lack one of NTHi colonising predicted strategies. However, the results obtained in this study do not support such hypothesis, given that NTHi375\textit{Δhap} mouse lung infection was comparable to that observed for the wild-type strain. Of note, NTHi375\textit{ΔompP5Δhap} double mutant lung bacterial load after a 48 h infection was higher than that obtained for NTHi375\textit{ΔompP5}.

We acknowledge that extrapolation of the results obtained on cultured human cell lines to tissue locations \textit{in vivo} should be considered with caution. Also, some of the phenotypes obtained for the \textit{ompP5} gene agree with those previously shown for different strain backgrounds, supporting the results presented in this study; moreover, the gain of adhesive function by homologous expression of the \textit{hap} gene in the Rd KW20 naturally \textit{hap} deficient strain allowed us to assign an adhesive role to Hap\textsubscript{NTHi375}. In conclusion, this study presents a comparative analysis of NTHi-host interaction through cells representing different anatomical locations of the respiratory tract, frames the relative contribution of two major bacterial surface proteins, P5 and Hap, to this interplay, and extends our understanding of the colonisation and infection process by NTHi. We acknowledge that, considering the observed heterogeneity among NTHi strains [4], further systematic correlation between large repertoires of well characterised NTHi isolates and wide panels of relevant phenotypic traits, together with a detailed characterisation of cell surface receptors, will contribute to unravel key bacterial and host cell elements for the NTHi-host interplay. Interference strategies designed to abolish such interplay would limit NTHi adaptation to the human respiratory tract, therefore preventing colonisation and/or pathogenicity.
Supporting Information

S1 File. Table A, Level of conservation of P5NTHi375 and HapNTHi375. Length and percentages of identity, similarity and gaps for P5NTHi375 and HapNTHi375, compared to P5 and Hap orthologous proteins from NTHi strains listed below (BLASTp results). Fig A, Multiple sequence alignment for 311 amino acids at the C-terminal region of HapNTHi375 domain, corresponding to the SAAT domain. Alignment was performed in Muscle [49], by using all NCBI available NTHi orthologous proteins.

Acknowledgments

We thank Antonio López-Gómez for technical work, Rosemary Redfield (University British Columbia) for providing pSU20 cloning vector, and Francisco Sánchez Madrid (Centro Nacional Investigaciones Cardiovasculares) for providing antibody P1D6.

Author Contributions

Conceived and designed the experiments: BE JAB JG. Performed the experiments: BE CV JM JV. Analyzed the data: BE CV JM IRM JV JG. Contributed reagents/materials/analysis tools: IRM JV. Wrote the paper: JAB JG.

References

1. Agrawal A, Murphy TF (2011) Haemophilus influenzae infections in the H. influenzae type b conjugate vaccine era. J Clin Microbiol 49: 3728–3732. doi:10.1128/JCM.05476-11 PMID: 21900515
2. Garmendia J, Martí-Lliteras P, Moleres J, Puig C, Bengoechea JA (2012) Genotypic and phenotypic diversity of the noncapsulated Haemophilus influenzae: adaptation and pathogenesis in the human airways. Int Microbiol 15: 159~172. PMID: 23844475
3. Spahich NA, St Geme JW (2011) Structure and function of the Haemophilus influenzae autotransporters. Front Cell Infect Microbiol 1: 5. doi: 10.3389/fcimb.2011.00005 PMID: 22919571
4. De Chiara M, Hood D, Muzzi A, Pickard DJ, Perkins T, et al. (2014) Genome sequencing of disease and carriage isolates of nontypeable Haemophilus influenzae identifies discrete population structure. Proc Natl Acad Sci U S A 111: 5439–5444. doi:10.1073/pnas.1403353111 PMID: 24706866
5. Martí-Lliteras P, López-Gomez A, Mauro S, Hood DW, Viadas C, et al. (2011) Nontypable Haemophilus influenzae displays a prevalent surface structure molecular pattern in clinical isolates. PLoS One 6: e21133. doi:10.1371/journal.pone.0021133 PMID: 21698169
6. Webb DC, Cripps AW (1998) Secondary structure and molecular analysis of interstrain variability in the P5 outer-membrane protein of non-typable Haemophilus influenzae isolated from diverse anatomical sites. J Med Microbiol 47: 1059–1067. PMID: 9856641
7. Sirakova T, Kolattukudy PE, Murwin D, Billy J, Leake E, et al. (1994) Role of fimbriae expressed by non-typeable Haemophilus influenzae in pathogenesis of and protection against otitis media and relatedness of the fimbrin subunit to outer membrane protein A. Infect Immun 62: 2002–2020. PMID: 7909539
8. Reddy MS, Bernstein JM, Murphy TF, Faden HS (1996) Binding between outer membrane proteins of nontypeable Haemophilus influenzae and human nasopharyngeal mucin. Infect Immun 64: 1477–1479. PMID: 896123
9. Miyamoto N, Bakaletz LO (1996) Selective adherence of non-typeable Haemophilus influenzae (NTHi) to mucus or epithelial cells in the chinchilla eustachian tube and middle ear. Microb Pathog 21: 343–356. PMID: 8939642
10. Jiang Z, Nagata N, Molena L, Bakaletz LO, Hawkins H, et al. (1999) Fimbria-mediated enhanced attachment of nontypeable Haemophilus influenzae to respiratory syncytial virus-infected respiratory epithelial cells. Infect Immun 67: 187–192. PMID: 9864214
11. Murphy TF, Kirkham C (2002) Biofilm formation by nontypeable Haemophilus influenzae: strain variability, outer membrane antigen expression and role of pili. BMC Microbiol 2: 7. PMID: 11900553
12. Rosadini CV, Ram S, Akerley BJ (2014) Outer membrane protein P5 is required for resistance of nontypeable Haemophilus influenzae to both the classical and alternative complement pathways. Infect Immun 82: 640–649. doi: 10.1128/IAI.01224-13 PMID: 24478079

PLOS ONE | DOI:10.1371/journal.pone.0123154 April 20, 2015 16/18
13. Wong SM, Bernui M, Shen H, Akerley BJ (2013) Genome-wide fitness profiling reveals adaptations required by *Haemophilus* in coinfection with influenza A virus in the murine lung. Proc Natl Acad Sci U S A 110: 15413–15418. doi: 10.1073/pnas.1312117110 PMID: 24003154

14. Hill DJ, Toleman MA, Evans DJ, Villullas S, Van Alphen L, et al. (2001) The variable P5 proteins of typeable and non-typeable *Haemophilus influenzae* target human CEACAM1. Mol Microbiol 39: 850–862. PMID: 11251807

15. Bookwalter JE, Jurcisek JA, Gray-Owen SD, Fernandez S, McGillivary G, et al. (2008) A carcinoembryonic antigen-related cell adhesion molecule 1 homologue plays a pivotal role in nonotypeable *Haemophilus influenzae* colonization of the chinchilla nasopharynx via the outer membrane protein P5-homologous adhesin. Infect Immun 76: 48–55. PMID: 17938212

16. Novotny LA, Partida-Sanchez S, Munson RS Jr., Bakaletz LO (2008) Differential uptake and processing of a *Haemophilus influenzae* P5-derived immunogen by chinchilla dendritic cells. Infect Immun 76: 967–977. PMID: 18160476

17. Frick AG, Joseph TD, Pang L, Rabe AM, St Geme JW, et al. (2000) *Haemophilus influenzae* stimulates ICAM-1 expression on respiratory epithelial cells. J Immunol 164: 4185–4196. PMID: 10754314

18. Avadhanula V, Rodriguez CA, Ulett GC, Bakaletz LO, Adderson EE (2006) Nontypeable *Haemophilus influenzae* adheres to intercellular adherence molecule 1 (ICAM-1) on respiratory epithelial cells and upregulates ICAM-1 expression. Infect Immun 74: 830–838. PMID: 16428725

19. St Geme JW, de la Morena ML, Falkow S (1994) A *Haemophilus influenzae* IgA- protease-like protein promotes intimate interaction with human epithelial cells. Mol Microbiol 14: 217–233. PMID: 7830568

20. Fink DL, Buscher AZ, Green B, Fernsten P, St Geme JW (2003) The *Haemophilus influenzae* Hap autotransporter mediates microcolony formation and adherence to epithelial cells and extracellular matrix via binding regions in the C-terminal end of the passenger domain. Cell Microbiol 5: 175–186. PMID: 12614461

21. Fink DL, Green BA, St Geme JW (2002) The *Haemophilus influenzae* Hap autotransporter binds to fibronectin, laminin, and collagen IV. Infect Immun 70: 4902–4907. PMID: 12183535

22. Spahich NA, Kenjale R, McCann J, Meng G, Ohashi T, et al. (2014) Structural determinants of the interaction between the *Haemophilus influenzae* Hap autotransporter and fibronectin. Microbiology.

23. Fink DL, Cope LD, Hansen EJ, Geme JW (2001) The *Hemophilus influenzae* Hap autotransporter is a chymotrypsin clan serine protease and undergoes autoproteolysis via an intermolecular mechanism. J Biol Chem 276: 39492–39500. PMID: 11504735

24. Hendrixson DR, St Geme JW (1998) The *Haemophilus influenzae* Hap serine protease promotes adherence and microcolony formation, potentiated by a soluble host protein. Mol Cell 2: 841–850. PMID: 9885571

25. Webster P, Wu S, Gomez G, Apicella M, Plaut AG, et al. (2006) Distribution of bacterial proteins in biofilms formed by non-typeable *Haemophilus influenzae*. J Histochem Cytochem 54: 829–842. PMID: 16549506

26. Spahich NA, Hood DW, Moxon ER, St Geme JW (2012) Inactivation of *Haemophilus influenzae* lipopolysaccharide biosynthesis genes interferes with outer membrane localization of the Hap autotransporter. J Bacteriol 194: 1815–1822. doi: 10.1128/JB.06316-11 PMID: 22287523

27. Bouchet V, Hood DW, Li J, Brisson JR, Randle GA, et al. (2003) Host-derived sialic acid is incorporated into *Haemophilus influenzae* lipopolysaccharide and is a major virulence factor in experimental otitis media. Proc Natl Acad Sci U S A 100: 8898–8903. PMID: 12855765

28. Hood DW, Makepeace K, Deadman ME, Rest RF, Thibault P, et al. (1999) Sialic acid in the lipopolysaccharide of *Haemophilus influenzae*: strain distribution, influence on serum resistance and structural characterization. Mol Microbiol 33: 679–692. PMID: 10447878

29. Morey P, Viadas C, Euba B, Hood DW, Barberán M, et al. (2013) Relative contributions of lipooligosaccharide inner and outer core modifications to nontypeable *Haemophilus influenzae* pathogenesis. Infect Immun 81: 4100–4111. doi: 10.1128/IAI.00492-13 PMID: 23980106

30. Allen S, Zaleski A, Johnston JW, Gibson BW, Apicella MA (2005) Novel sialic acid transporter of *Hemophilus influenzae*. Infect Immun 73: 5291–5300. PMID: 16113244

31. Herrriott RM, Meyer EY, Vogt M, Modan M (1970) Defined medium for growth of *Haemophilus influenzae*. J Bacteriol 101: 513–516. PMID: 5308770

32. Sánchez R (1998) A medium-copy-number plasmid for insertional mutagenesis of *Streptococcus mutans*. Plasmid 40: 247–251. PMID: 9806683

33. Morey P, Cano V, Martí-Lliteras P, López-Gómez A, Reguiero V, et al. (2011) Evidence for a non-replicative intracellular stage of nontypeable *Haemophilus influenzae* in epithelial cells. Microbiology 157: 234–250. doi: 10.1099/mic.0.040451-0 PMID: 20929955
34. Gray-Owen SD, Dehio C, Haude A, Grunert F, Meyer TF (1997) CD66 carcinoembryonic antigens mediate interactions between Opa-expressing Neisseria gonorrhoeae and human polymorphonuclear phagocytes. EMBO J 16: 3435–3445. PMID: 9218786

35. López-Gómez A, Cano V, Moranta D, Morey P, García del Portillo F, et al. (2012) Host cell kinases, α5 and β1 integrins, and Rac1 signalling on the microtubule cytoskeleton are important for non-typable Haemophilus influenzae invasion of respiratory epithelial cells. Microbiology 158: 2384–2398. doi: 10.1099/mic.0.059972-0 PMID: 22723286

36. Marti-Lliteras P, Regueiro V, Morey P, Hood DW, Saus C, et al. (2009) Nontypeable Haemophilus influenzae clearance by alveolar macrophages is impaired by exposure to cigarette smoke. Infect Immnu 77: 4232–4242. doi: 10.1128/IAI.00305-09 PMID: 19620348

37. Mell JC, Sinha S, Balashov S, Viadas C, Grassa CJ, et al. (2014) Complete genome sequence of Haemophilus influenzae strain 375 from the middle ear of a pediatric patient with otitis media. Genome Announc 2(6). pii: e01245–14. doi: 10.1128/genomeA.01245-14 PMID: 25477405

38. Garmendia J, Viadas C, Calatayud L, Mell JC, Marti-Lliteras P, et al. (2014) Characterization of nontypeable Haemophilus influenzae isolates recovered from adult patients with underlying chronic lung disease reveals genotypic and phenotypic traits associated with persistent infection. PLoS One 9: e97020. doi: 10.1371/journal.pone.0097020 PMID: 24824990

39. Rougeaux C, Berger CN, Servin AL (2008) hCEACAM1-4L downregulates hDAF-associated signalling after being recognized by the Dr adhesin of diffusely adhering Escherichia coli. Cell Microbiol 10: 632–654. PMID: 17979980

40. Zheng M, Fang H, Hakomori S (1994) Functional role of N-glycosylation in α5β1 integrin receptor. De-N-glycosylation induces dissociation or altered association of α5 and β1 subunits and concomitant loss of fibronectin binding activity. J Biol Chem 269: 12325–12331. PMID: 7512965

41. St Geme JW (1994) The HMW1 adhesin of nontypeable Haemophilus influenzae recognizes sialylated glycoprotein receptors on cultured human epithelial cells. Infect Immnu 62: 3881–3889. PMID: 8063405

42. Russell T, Bell TJ (2014) Alveolar macrophages: plasticity in a tissue-specific context. Nat Rev Immunol 14: 81–93. doi: 10.1038/nri3600 PMID: 24445666

43. Fink DL, St Geme JW (2003) Chromosomal expression of the Haemophilus influenzae Hap autotransporter allows fine-tuned regulation of adhesive potential via inhibition of intermolecular autoproteolysis. J Bacteriol 185: 1608–1615. PMID: 12591878

44. Fleischmann RD, Adams MD, White O, Clayton RA, Kirkness EF, et al. (1995) Whole-genome random sequencing and assembly of Haemophilus influenzae Rd. Science 269: 496–512. PMID: 7542800

45. Virji M, Evans D, Griffith J, Hill D, Serino L, et al. (2000) Carcinoembryonic antigens are targeted by diverse strains of typable and non-typable Haemophilus influenzae. Mol Microbiol 36: 784–795. PMID: 10844687

46. Duim B, Bowler LD, Eijk PP, Jansen HM, Dankert J, et al. (1997) Molecular variation in the major outer membrane protein P5 gene of nonencapsulated Haemophilus influenzae during chronic infections. Infect Immnu 65: 1351–1356. PMID: 9119473

47. Otto VI, Damoc E, Cueni LN, Schurpf T, Frei R, et al. (2006) N-glycan structures and N-glycosylation sites of mouse soluble intercellular adhesion molecule-1 revealed by MALDI-TOF and FTICR mass spectrometry. Glycobiology 16: 1033–1044. PMID: 16877748

48. Gauthier F, Fryksmark U, Ohlsson K, Bieth JG (1982) Kinetics of the inhibition of leukocyte elastase by the bronchial inhibitor. Biochim Biophys Acta 700: 178–183. PMID: 6915782

49. Edgar RC (2004) MUSCLE: multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Res 32: 1792–1797. PMID: 15034147

50. Marchler-Bauer A, Zheng C, Chitsaz F, Derbyshire MK, Geer LY, et al. (2013) CDD: conserved domains and protein three-dimensional structure. Nucleic Acids Res 41: D348–352. doi: 10.1093/nar/gks1243 PMID: 23197659

51. Mullins MA, Register KB, Bayles DO, Loving CL, Nicholson TL, et al. (2009) Characterization and comparative analysis of the genes encoding Haemophilus parasuis outer membrane proteins P2 and P5. J Bacteriol 191: 5989–6002. doi: 10.1128/JB.00469-09 PMID: 19633080

52. Kilian M, Poulsen K, Lombolt H (2002) Evolution of the paralogous hap and iga genes in Haemophilus influenzae: evidence for a conserved hap pseudogene associated with microcolony formation in the recently diverged Haemophilus aegyptius and H. influenzae biogroup aegyptius. Mol Microbiol 46: 1367–1380. PMID: 12453222