Epitope-specific immune recognition of the nontypeable Haemophilus influenzae outer membrane protein 26

Author
Kunthalert, Duangkamol, A. Novotny, Laura, Massa, Helen, Ulett, Glen, O. Bakaletz, Lauren, M. Kyd, Jennelle, Cripps, Allan

Published
2013

Journal Title
Human Vaccines & Immunotherapeutics

DOI
https://doi.org/10.4161/hv.23255

Copyright Statement
Copyright remains with the authors 2013. This is the author-manuscript version of this paper. It is posted here with permission of the copyright owners for your personal use only. No further distribution permitted. For information about this journal please refer to the journal's website or contact the authors.

Downloaded from
http://hdl.handle.net/10072/57422
Epitope-Specific Immune Recognition of the
Nontypeable *Haemophilus influenzae* outer membrane protein 26

Kunthalert, Duangkamol. ¹, Novotny, Laura.A. ², Massa, Helen.M. ³, Ulett, Glen.C. ³, Bakaletz, Lauren.O. ², Kyd, Jennelle.M. ⁴, Cripps, Allan.W. ³*

¹ Faculty of Medical Science, Naresuan University, Thailand, ² The Research Institute at Nationwide Children's Hospital, Ohio, United States, ³ Griffith Health Institute, School of Medical Science, Griffith University, Gold Coast, Australia, ⁴ Swinburne University, Victoria, Australia.

**Running Title:** Immune Recognition of Nontypeable *Haemophilus influenzae* OMP26 epitopes

**Keywords:** Nontypeable *Haemophilus influenzae*, outer membrane protein, OMP26, B and T-cell epitopes, vaccine, peptide vaccine, immune response.

**Corresponding Author:** Allan W Cripps

Griffith Health Institute, School of Medical Science, Griffith University, Queensland 4222 Australia

Tel: +61 7 5678 0709 Fax: +61 7 5678 0795

Email: allan.cripps@griffith.edu.au
ACKNOWLEDGMENTS

The authors thank Penny Chapman for her editorial assistance in the preparation of the manuscript.
LIST OF ABBREVIATIONS

APC - Antigen presenting cell
BCA – Bicinchoninic acid
CPM – Counts per minute
ELISA – Enzyme linked immunosorbent assay
FBS – Fetal bovine serum
IPP/IT – Intra-Peyer’s patches/intra-tracheal
IP – Intra-peritoneal
MHC – Major histocompatibility complex
Ni-NTA - Nickel-nitriloacetic acid
NTHi – Nontypeable *Haemophilus influenzae*
OMP26 – Outer membrane protein 26
PBS – Phosphate buffered saline
PMSF - Phenylmethylsulfonyl fluoride
RU – Resonance units
SC – Subcutaneous
SDS-PAGE - Sodium dodecyl sulfate polyacrylamide gel electrophoresis
SPF – Specific-pathogen free
SPR – Surface plasmon resonance
ABSTRACT

Previous studies using rodent respiratory infection models of nontypeable Haemophilus influenzae (NTHi) infection have established the 26-kDa outer membrane protein of the bacterium, OMP26, as a potential vaccine antigen for NTHi. This study undertook a comprehensive immunological identification of OMP26 T- and B-cell epitopes. A series of OMP26 peptides were constructed and regions of the OMP26 antigen involved in recognition by lymphocyte receptors and induction of acquired immune responses were identified. The dominant T-cell epitopes for OMP26 were located towards the C-terminus between amino acid residues 95 and 197 (T3+T4 region) as mapped using antigen-specific lymphocyte proliferation assays. The newly identified T-cell epitopes exhibited strong capacity for efficient T-cell activation, suggesting that, compared to other OMP26 regions; epitopes within the T3+T4 region have the highest affinity for binding to major histocompatibility complex molecules. In contrast, the predominant B-cell epitopes of OMP26 were located more centrally within the molecule between amino acid residues 45 and 145 (T2+T3 region) as determined using enzyme linked immunosorbent assay and surface plasmon resonance assays. The T2+T3 region was immunodominant in several species including chinchilla, mice, and rats when assessed using both mucosal and parenteral immunization regimes. In addition, the antibodies directed against the T2+T3 region bound to intact NTHi cell surface, according to flow cytometry. Collectively, these results specifically locate the amino acid sequences containing the OMP26 T- and B-cell epitopes, which, as newly mapped antigenic epitopes for lymphocyte recognition, will be useful to improve existing NTHi vaccine strategies. Comprehensive definition of the minimum epitope length required for optimal B- and T-cell responses requires further study.
INTRODUCTION

Nontypeable *Haemophilus influenzae* (NTHi) is a significant human pathogen causing a wide range of respiratory infections. Several outer membrane proteins (OMPs) of NTHi and its oligosaccharide have been investigated as possible vaccine antigens against NTHi. One OMP that has shown promise as a potential vaccine candidate is OMP26. The amino acid sequence of this 26kDa OMP is conserved among NTHi isolates from various disease states.

Our laboratory has previously shown that immunization with OMP26 can stimulate enhanced pulmonary clearance of NTHi in a rat model in which animals were initially immunized via intra-Peyer’s patches followed by intra-tracheal boost (IPP/IT). Mucosal immunization with OMP26 protected animals against subsequent pulmonary challenge with both homologous and heterologous strains of NTHi and induced high levels of OMP26-specific IgA and IgG antibodies. Furthermore, parenteral immunization of chinchillas with OMP26 demonstrated good immunogenicity and enhanced the clearance of NTHi from the nasopharynx. Thus, OMP26 is appealing as an immunogen against NTHi and has demonstrated potential as a candidate vaccine antigen for this pathogen.

A high degree of antigenic heterogenicity between NTHi strains has led to vaccine approaches based on peptide formulations of immunodominant epitopes of the native protein. In one study, T-cell epitopes were included in a peptide-based approach to maximize induction of antibodies with higher affinity for the incorporated B-cell epitopes. This approach offers an additional advantage of accommodating multiple epitopes to cover a broader range of antigenically-distinct NTHi strains.

OMP26 is highly conserved among a large number of clinical NTHi isolates collected from a range of anatomical sites. Typically, vaccine formulations do not
favour the use of a single protein, however, a highly conserved protein such as OMP26 may provide the necessary broad-based protection against geographically-diverse and antigenically-distinct isolates of NTHi.

This study assessed epitope specificity of the immune responses to OMP26 by mapping the location of T- and B-cell epitopes within the protein to further characterize the immune response to OMP26. These results reveal unique T- and B-cell-targeting regions within OMP26 that will aid in the development of improved peptide-based vaccine strategies for NTHi.
RESULTS

Lymphoproliferative responses to OMP26 peptides

Lymphoproliferative response studies were conducted using splenocytes derived from rats and mice. Unfortunately, background cross reactivity against \textit{E. coli} proteins within the mouse samples masked any specific responses and thus only rat data are presented. To localize the immunologically important regions within OMP26 in this response, a series of overlapping OMP26 peptides spanning the entire sequence of full-length OMP26 was used as the \textit{in vitro} proliferation stimulus. Proliferation in response to Concanavalin A ranged from 85,000 to 110,000 counts per minute (CPM).

At a concentration of 1 µg/ml the OMP26 peptides stimulated little or no response from OMP26-primed lymphocytes with the exception of T3+T4 peptide and the whole OMP26 molecule itself where significant stimulation was observed (P<0.001) (data not shown). In contrast, compared to naive lymphocytes, a peptide concentration of 10 µg/ml stimulated significant increases in lymphocyte proliferation (Figure 1) in response to the T4 peptide (P<0.02; amino acid residues 140-197), T2+T3 peptide (P<0.05; amino acid residues 45-145) and T3+T4 peptide (P<0.001; amino acid residues 95-197). Lymphocyte proliferation stimulated by the individual peptides, T1 (amino acid residues 1-55), T2 (amino acid residues 45-100) or T3 (amino acid residues 95-145) and the T1a+T2 peptide (amino acid residues 24-100), did not differ significantly between the immunized versus non-immunized group. These data suggested that in terms of lymphocyte proliferation, peptides that represented the C-terminus of OMP26 were more stimulatory than peptides that represented the N-terminus of the protein.
Lymphoproliferative responses to OMP26 protein

The capacity of OMP26 peptides to prime T cells and induce lymphoproliferative responses against OMP26 protein was examined using rat lymphocytes obtained after immunization with OMP26 peptides. Lymphocytes from immune and non-immune rats were stimulated with whole OMP26 protein (10 µg/ml). Only lymphocytes from animals immunized with the T3+T4 peptide demonstrated a significant proliferative response to OMP26 protein stimulation (Figure 2), further implicating the C-terminus (AA 95-197) of OMP26 as a region of immunologic importance.

Immunogenicity of OMP26 peptides

Immunogenicity of each of the individual peptides was confirmed by immunization of the rats (IPP/IT) with the individual peptides and subsequent analysis of the anti-OMP26 sera using both enzyme linked immunosorbent assay (ELISA) and surface plasmon resonance (SPR). All antipeptide sera were reactive with their individual immunizing peptides, in addition to being reactive to the whole OMP26 protein, with the exception of the anti-T1 sera (Table 1). These sera were then used to determine whether the peptides were surface exposed on NTHi whole cells by flow cytometry.

Determination of surface-exposed regions of OMP26

Antibodies to the 50-mer peptides (T1, T2, T3, T4) all showed distinct and reproducible shifts of the curve to the right (Figure 3) which indicated binding of antibody to native structure on the bacterium. In addition, antisera to the 100-mer peptide T1a+T2 also showed a distinct shift to the right. In comparison, antisera to the
100 mer T2+T3 peptide demonstrated a lesser but consistent shift to the right whilst 
antisera against the T3+T4 peptide gave results similar to the non-immune serum. 

**Antibody responses assessed by ELISA and SPR analysis**

**Reactivity of mouse anti-OMP26 antibodies to OMP26 peptides**

B-cell epitope regions of OMP26 were examined in mice using pooled anti- 
OMP26 sera. The antibody responses against the OMP26 protein and the OMP26 
peptides in the mouse were measured using ELISA and SPR. For the ELISA results, 
animals were immunized either; IPP/IT (Figure 4A, 4C) or intra-peritoneal (IP) alone 
(Figure 4B, 4D). SPR assays to determine the antigen/antibody binding in real time 
are also shown for the IPP/IT and IP immunized groups (Figures 4C and 4D, 
respectively).

In mice, the predominant reactivity measured by ELISA was seen against the 
T2+T3 peptide, regardless of immunization regime (Figures 4A, 4B). Minor reactivity 
was also observed against the T1a+T2 peptide in the IPP/IT immunized group (Figure 
4A), whilst in the IP immunized group, low level reactivity was also observed in the 
T3 and T3+T4 peptides (Figure 4B).

Similarly, SPR analyses showed predominant OMP26 reactivity against the 
T2+T3 peptide regardless of immunization regime (Figures 4C and 4D). In both the 
IPP/IT and IP immunization groups, low levels of reactivity were also observed 
against the T2, T3, T4, T1a+T2 and T3+T4 peptides, whilst no reactivity was 
observed for the T1 peptide. These data indicated that immunization of mice with 
OMP26 protein yielded the production of antibodies with affinity primarily to the 
larger T2+T3 peptide, which represents the mid-region of OMP26 (AA 45-145).

**Reactivity of rat anti-OMP26 antibodies to OMP26 peptides**
The antibody responses against the OMP26 protein and the OMP26 peptides in the rat were measured using ELISA and SPR and the IPP/IT and IP immunization regimens (Figure 5).

In the rats immunized IPP/IT, a predominant response was observed against the T2+T3 peptide using ELISA, and minor reactivities were observed against the T2, T3, T1a+T2 and the T3+T4 peptides (Figure 5A). For animals immunized IP, the ELISA results again showed a substantial response against the T2+T3 peptide (Figure 5B). In addition, similar response was observed against the T1a+T2 peptide with substantially less reactivity stimulated in response to T3 and T3+T4 peptides.

In general, reactivity of rat serum assessed using SPR was greater and thus resulted in greater overall reactivity to the OMP26 peptides than observed with the ELISA.

Serum from rats immunized IPP/IT demonstrated substantial reactivity against T2+T3 peptide and lesser absolute responses were observed against the T2 and T3 peptides in comparison to the non-immune sera (Figure 5C). In animals immunized IP, the SPR reactivity provided even greater responses by the rat serum antibodies with the predominant response observed to the T2+T3 peptide. Substantial responses were also observed against the T2, T3, T1a+T2 and T3+T4 peptides (Figure 5D). Therefore, as observed with mice, the response induced in rats by immunization with OMP26 yielded antibodies with greater affinity to the mid-region of the protein.

**Reactivity of chinchilla anti-OMP26 antibodies to OMP26 peptides**

The antibody responses against the OMP26 protein and the OMP26 peptides in the chinchilla after subcutaneous (SC) immunization were also measured using SPR and the results are shown in Figure 6. Chinchilla anti-OMP26 antibodies exhibited the greatest overall reactivity to the T2+T3 peptide. Peptides T2, T3, and T4
showed lesser reactivity, whilst the T3+T4 peptide exhibited the lowest level of reactivity among the panel of peptides. Little antibody reactivity was observed to peptides T1 and T1a+T2.

**Relative reactivity score**

In order to summarise the data across species and immunization regimes, a relative reactivity score was calculated against the OMP26 response. The OMP26 response was assigned a relative reactivity score of 5 and the responses of the other OMP26 peptides were ranked against the OMP26 reactivity such that a response to an individual peptide that was 10-20% of the reactivity to OMP26 protein was scored as 1, 20-40% as 2, 40-60% as 3, 60-80% as 4, 80-100% as 5. No significant reactivity was detected against the T1 or T4 peptides regardless of species or immunization regime, although the chinchilla did demonstrate some reactivity to T2. Some reactivity was observed in response to all other peptides, with the T2+T3 peptide demonstrating the greatest activity (Figure 7). Therefore, in terms of identification of potential B-cell epitope(s) within OMP26, antibodies against the mid-region of the protein consistently yielded the greatest reactivity, regardless of rodent species or immunization regime.
DISCUSSION

NTHi is recognised as a significant human pathogen and has become the focus of attention towards the development of an effective vaccine against NTHi infections. Previous investigations have identified the efficacy of OMP26 as a potential vaccine antigen. The current study examined the immunological recognition of T- and B-cell epitopes within truncated OMP26 to further characterize this vaccine candidate.

A series of overlapping peptides was constructed to localize the immunologically important regions for the OMP26. These peptides utilized the amino acid sequence of OMP26 identified in the strain NTHi-289 and included the 23-amino acid leader peptide. Previous studies have shown that the inclusion of the leader peptide and signal sequence in the recombinant OMP26 increased efficacy in a rodent model of acute respiratory infection.

In the current study, each OMP26 peptide overlapped the preceding and/or subsequent peptide by 6-11 amino acid residues. The OMP26 peptide constructs consisted of four overlapping peptides (T1-T4) of approximately 50 amino acids in length that spanned the entire OMP26 sequence. In addition, combination peptides of approximately 100 amino acids in length (T1a+T2 to T3+T4) were designed to include the adjacent regions of each of the T1-T4 peptides. This design increased the opportunity for detection of all immunologically relevant regions within OMP26 in our analysis and thus provides the first detailed molecular insight into the basis of immune reactivity to OMP26.

In the rat, lymphocyte proliferative responses following priming with OMP26 protein showed strong antigen specific responses to both T4 and the T3+T4 peptides with the magnitude of the T3+T4 peptide exceeding that of the T4 alone. However,
individually, T2 and T3 failed to elicit a significant proliferative response. It is concluded that T-cell recognition is enhanced in the shared amino acid region between T3 and T4 and T-cell epitopes for OMP26 are located between amino acid residues 95 to 197, specifically the overlapping amino acid region between T3 and T4, upstream of amino acid residue 140. In addition, an OMP26-primed proliferative response was observed to the combined peptide T2+T3 but not to the individual T2 or T3 peptides suggesting that T-cell epitopes for OMP26 may also occur in the shared amino acid region between T2 and T3 (amino acid residues 94 to 100). When rats were immunized with each of the OMP26 peptides individually, the only lymphocyte proliferative response observed to the OMP26 protein was for the T3+T4 peptide. This supports the identification of a T-cell epitope within the T3+T4 peptide which is most likely located in the overlapping region and may be conformational although typically linear amino acid motifs are more readily recognized by the major histocompatibility complex (MHC) molecules.

Processing of the OMP26 protein in this study resulted in a peptide with similar flanking amino acid residues or similar conformation to that of the T3+T4 peptide, and presumably similar affinity for the MHC molecules. Lymphocytes primed with T4 and T2+T3 peptides did not prime any T-cell response, in contrast to the T3+T4 peptide-primed lymphocytes, despite proliferative responses to these peptides occurring in lymphocytes primed with the whole OMP26. Changes in the flanking amino acids and/or conformation of the T4 and the T2+T3 peptides may account for this failure and are of importance for subsequent vaccine designs utilizing epitopes.

Selection of T-cell epitopes may include competitive binding between the processed peptides contained within a single antigenic molecule for MHC molecules
Thus, only the peptides with the highest affinity to MHC molecules are ultimately presented to T-cells and the competition would be void when inoculation uses only a single peptide. The lack of apparent difference between the extent of proliferation generated by the T3+T4 peptide and intact OMP26 protein suggests that the T3+T4 region has highest affinity binding for MHC molecules in the current model.

Specificity of the epitope regions recognized by B- and T-cells vary and are dependent upon a number of factors including the host’s genetic background and route of immunization. Use of the two immunization regimes in this study has potentially presented the antigen to different sets of antigen presenting cells (APC)s for the primary encounter. Different APC populations have been postulated to process and present antigens differently and thus can account for differences in epitope recognition. Differences in B-cell epitope recognition following different routes of immunization may result from extracellular degradation at the site of immunization, although, the T2+T3 region may also be resistant to the extracellular medium encountered by these two immunization routes resulting in the identical B-cell recognition patterns. The route of immunization can also alter both the hierarchical organization of B- and T-cell epitopes, as well as, the titre and affinity of antibody responses. Use of multiple animal species (chinchilla, mice and rats) and immunization routes in the present study showed that the immunodominant B-cell epitopes of OMP26 are located towards the centre of the molecule between amino acid residues 45-145 (T2+T3).

In this study, the T2+T3 peptide exhibited the highest reactivity in mouse and rat anti-OMP26 sera regardless of immunization regime. For the development of a peptide based vaccine, the T2+T3 peptide would be a lead candidate for a genetically diverse human population.
Following SC immunization of chinchillas, anti-OMP26 sera also strongly recognized peptides T2, T3 and T4. Whilst antibody recognition levels to these peptide were not as high as those to the T2+T3 peptide. It does suggest additional B-cell epitopes within the T2, T3 and T4 regions.

Immune sera generated from rats immunized IP with OMP26 also demonstrated strong reactivity to the T1a+T2 peptide. Thus, apart from the T2+T3 region, additional B-cell epitopes may be located between amino acid residues 24 and 100.

The T1 peptide (amino acid residues 1-50) was not recognized by any immune sera from all three animal species tested and the T4 peptide (amino acid residues 140-197) was not recognized by the immune serum pools from both mice and rats. This lack of antibody reactivity suggests that the T1 and T4 regions were sequestered or inaccessible despite clear expression on the surface of NTHi cells being detected by flow cytometry. This finding is suggestive of less surface exposure of these regions and is consistent with evidence from previous studies showing a hydrophobic character exists at the N-terminal and C-terminal regions of OMP26. The lack of response to T1 and T4 peptides suggest that these peptide regions are not immunogenic, however, conformational diversion by T2+T3 dominance or assay bias cannot be excluded. Assay bias in this study is unlikely as antibody recognition patterns obtained from both ELISA and SPR were mostly similar despite potential differences in the orientation of the bound peptides on the microtiter plates and biosensor chips.

Anti-OMP26 serum pools from all three animal species strongly recognized the T2+T3 region, but reactivities to the T2 or T3 region were also detected by SPR. Reactivity to the individual T2 and T3 regions are interesting and do not approach the
level stimulated by the T2+T3 region. Whilst additional epitopes cannot be excluded, it is likely that the anti-OMP26 sera were reactive to the shared amino acid region between the T2 and T3 regions.

Since the majority of B-cell epitopes are discontinuous in nature, it is also possible that these identified B-cell epitopes are conformational. The sequence within the T2+T3 region may have brought amino acids into spatial proximity to form the conformation that resembles the OMP26 protein, and thus allowing antibodies raised against the whole protein to bind most efficiently.

Previous experiments involving immunolabeling of NTHi with antisera to OMP26 showed surface binding of gold-conjugated particles by transmission electron microscopy, suggesting at least one surface-exposed region on OMP26. Flow cytometry analysis revealed the binding of anti-T1, anti-T2, anti-T3 and anti-T4 to OMP26 on intact NTHi cells, indicating that within the T1 to T4, there were surface exposed regions. Interestingly, except for anti-T1a+T2 sera, antisera raised to T2+T3 and T3+T4 peptides did not bind to OMP26 expressed on NTHi cells. The T2+T3 and T3+T4 peptides were 100 and 102 acids in length respectively, whilst the T1a+T2 combination peptide was 76 amino acids in length and the other individual peptides were between 50 and 57 amino acids in length.

It is possible that the solubilized longer peptides (T2+T3, T3+T4) could be conformationally different to the native protein. As such, antibodies raised against these peptides could have recognized epitopes that were conformationally different from those of the native OMP26 protein, thus resulting in a failure to bind to NTHi cells. Again, this finding demonstrates the limitations of using long peptides as immunizing antigens. It is important to note that binding of antisera to the shorter individual peptides (T2, T3, T4) to OMP26 on the surface of NTHi cells occurred and
further analysis will be needed to determine which particular location within the long peptide regions (T2+T3 and T3+T4) contributes to conformational changes.

Key target regions of the acquired immune responses for the NTHi vaccine candidate, OMP26, have been identified in this study. The regions covering amino acid residues 45-145 (T2+T3 region), and 95-197 (T3+T4 region) were the dominant B- and T-cell epitopes for OMP26, respectively. At least two potential T-cell epitopes for OMP26 were located within the T3+T4 region, one was within the T4 region, and another one was within the shared amino acids upstream of residue 140. The T-cell epitopes within the T3+T4 region potentially had the highest affinity binding for the MHC molecules compared with other OMP26 region, and contained a full-length sequence required for efficient T-cell activation. The identified B-cell epitopes were immunodominant across all animal species tested, and with all immunization regimens used. Additionally, these particular epitopes were expressed on the surface of NTHi cells. The significant outcomes from the present study provide a better understanding of the specificity of the host immune responses, both humoral and cell-mediated, to the NTHi vaccine candidate, OMP26. Significantly, knowledge of the key target regions identified will be of great benefit for the future design of a better whole protein, peptide and/or epitope based vaccine against NTHi infections.

Future studies are still needed to define the minimum epitope length required for B- and T-cell responses identified in this study. The significance of these responses in immune protection against NTHi infection also requires further investigation.
MATERIALS AND METHODS

Bacterial strains, plasmids and growth conditions

This study used NTHi-289, a biotype I strain isolated from the sputum of a patient with chronic bronchitis. Bacteria were grown at 37°C in 5% CO₂ on brain heart infusion agar (Oxoid, UK) supplemented with 5% (v/v) defibrinated horse blood (BioMerleux, Australia).

Escherchia coli M15 [pREP4] (Qiagen, USA) was used as a host strain for the recombinant plasmid DNA. Transformed E. coli was propagated on Luria-Bertani (LB) (Life Technologies, USA) or in liquid LB supplemented with 100 μg/ml of ampicillin and 50 μg/ml of kanamycin. Bacteria were grown at 37°C and liquid cultures were aerated with shaking at 220-250 rpm in a BioLine Shaking Incubator (Edwards Instrument Company, Australia) unless stated otherwise. Plasmid pQE30 was purchased from Qiagen (GmbH, Germany) and purified using QIAprep Spin Miniprep columns (Qiagen) according to the manufacturer’s instruction.

OMP26 protein and peptides

OMP26 protein. The OMP26 protein used in this study was purified from a OMP26VTAL clone of NTHi-289. The 6x His-tagged recombinant protein contains all but the first 6 amino acids (MKNIAK-negative) of the complete OMP26 protein, and performs in a similar manner to the complete protein in animal immunization studies, as previously reported. The leader peptide of the recombinant protein is less efficiently cleaved by E. coli signal peptidases, therefore, use of the MKNIAK-negative clone results in higher yields of purified protein and thus, is an improvement on the original OMP26 (McGrath, unpublished data).
OMP26 peptides. A series of overlapping peptides was designed and constructed to span the entire sequence of OMP26, based on the amino acid sequence of the full length OMP26 from NTHi-289. The locations of the peptide regions are shown in Figure 8.

T1 peptide. The T1 peptide, representing the first 50 amino acids of full-length OMP26, was synthesized by Auspep Pty., Australia. The synthesized peptide was purified by high performance liquid chromatography on a Superspher®, 250-4, LiChroCART, 100 RP-18 column (Merck Serono, Australia) using an acetonitrile gradient of 0 to 70% and 0.1% trifluoroacetic acid as the mobile phase. The molecular weight, confirmed by mass spectral analysis, was 5344 Da.

Inclusion of the hydrophobic 23-amino acid signal sequence of OMP26 resulted in difficulty in the expression and purification of the T1 peptide, regardless of the E. coli expression system used. This phenomenon has been previously reported for short hydrophobic proteins34. Exclusion of this hydrophobic sequence in the peptide design resulted in the successful expression and purification of the T1a+T2 peptide (results not shown). To ensure that the immunological studies were not compromised, we did not attempt to improve solubility of the protein and utilized a commercially synthesized T1 peptide for the immunological studies.

T2, T3, T4, T1a+T2, T2+T3 and T3+T4 peptides. Peptides were constructed by cloning various OMP26 gene segments into a pQE30 expression vector. Gene segments were obtained by PCR amplification of genomic DNA of NTHi-289 using primer pairs as listed in Table 2. Recombinant plasmids were transformed into E. coli M15 [pREP4] and expression and purification of recombinant OMP26 peptides was performed in accordance with the manufacturer’s instructions (Qiagen). An overnight culture of E. coli M15 harboring recombinant pQE30 was diluted 1:50 in 400 ml of
fresh, pre-warmed LB containing ampicillin and kanamycin and grown at 30°C with shaking until the OD<sub>600</sub> reached 0.6-1.0. Expression of recombinant protein was then induced with 1 mM isopropylthio-β-galactoside (Invitrogen™; Life Technologies) for 2 hours at 30°C. Bacteria were harvested (5,000g, 20 minutes at 4°C: Hettich Zentrifugen Universal 32 Centrifuge; Hettich, Germany) and the pellet was stored at -80°C until protein purification. For purification, the cell pellet was thawed on ice and resuspended with lysis buffer (50 mM NaH<sub>2</sub>PO<sub>4</sub>, 300 mM NaCl, 10 mM imidazole, pH 8.0) to give 25x concentration factor (e.g. 4 ml/100 ml culture). To prevent protein degradation, phenylmethylsulfonyl fluoride (PMSF) (Sigma-Aldrich, USA) was added at a final concentration of 0.1 mM. The cells were then treated with lysozyme (final concentration of 1 mg/ml; Sigma-Aldrich) for 30 minutes on ice, and subsequently sonicated (Branson Sonifier 250 micro tip; 20-40 watt output: Branson Ultrasonics, USA) on ice in 6x10 second bursts, with 20 second cooling periods between each burst. After centrifugation (10,000g, 30 minutes at 4°C), 4 ml of the cleared lysate was transferred to a 10 ml centrifuge tube containing 1 ml of 50% (w/v) nickel-nitriloacetic acid (Ni-NTA) slurry (Qiagen). The lysate-Ni-NTA mixture was rocked on ice for 60 minutes, followed by centrifugation at 1000g for 3 minutes. The supernatant was discarded and the Ni-NTA agarose pellet was washed twice with 4 ml of wash buffer (50 mM NaH<sub>2</sub>PO<sub>4</sub>, 300 mM NaCl, 50 mM imidazole, pH 8.0). Recombinant proteins bound to Ni-NTA matrices were eluted 4 times with 0.5 ml of elution buffer (50 mM NaH<sub>2</sub>PO<sub>4</sub>, 300 mM NaCl, 250 mM imidazole, pH 8.0) and stored at 4°C prior to analysis. OMP26 peptide purity were determined using sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) (16% w/v). Removal of imidazole in the elution buffer was performed by dialyzing the pooled eluates (containing purified OMP26 peptides) against 1 litre of wash buffer for 3
hours, dialysis buffer (50 mM NaH$_2$PO$_4$, 300 mM NaCl, 10% (v/v) glycerol) for 1 hour and then phosphate buffered saline (PBS) overnight at 4°C. Following dialysis, PMSF (final concentration of 0.1 mM) was added to the dialyzed protein solutions. The concentration of purified OMP26 peptides was determined by a microbicinchnonic acid (BCA) kit (Thermo Scientific, USA) as per the manufacturer’s instructions. Amino acid sequences for the OMP26 peptides are shown in Table 3.

**Immunization**

**Animals.** Specific-pathogen free (SPF) male DA rats aged 8-10 weeks or BALB/c mice aged 6-8 weeks were purchased from Animal Resources Centre, Western Australia, Australia and maintained under SPF conditions with food and water accessible *ad libitum*. Monitoring of rats and mice in experimental infection assays was performed according to protocols approved by the University of Canberra Animal Ethics Committee, Canberra Australia, until harvest, 21 days post immunization. Adult chinchillas (*Chinchilla lanigera*) were also included in this study and were examined using otoscopy or tympanometry prior to inclusion to confirm the absence of clinically recognizable signs of middle ear infection. Pre-immune serum samples, collected by cardiac puncture, were tested to determine the presence of pre-existing antibodies to NTHi-OMP $^{18}$ and only sero-negative animals were used in the study. All chinchilla experiments were conducted at Nationwide Children’s Hospital Research Institute, Columbus, Ohio. Chinchilla studies were approved by Nationwide Children's Hospital Research Institute’s Animal Care and Use Committee.

**IPP/IT Immunization.** Mucosal immunizations were performed for the mice and rats using an IPP/IT approach, as previously described $^9,35,36$. Twenty one days after the first immunization, animals were euthanized using pentobarbital sodium overdose and tissue harvest and blood collection were performed. Blood volumes of
0.5-1 ml (mice) or 8-10 ml (rats) were collected by cardiac puncture into plain 1 ml or 10 ml blood collection tubes (Sarstedt, Germany). Sera were pooled by cohort, separated by centrifugation, then frozen and stored at -20°C until analysis.

**SC immunization.** Six adult chinchillas were immunized SC with 10 μg of OMP26 plus 10 μg of SBAS-4 (GSK, Belgium) in 100 μl along the rear flank three times at 30 day intervals. Ten days after receipt of the third dose, the animals were bled by cardiac puncture, serum separated, pooled and frozen at -80°C until subsequent analysis.

**IP immunization.** The immunization antigen was prepared by emulsifying a required amount of antigen with Incomplete Freund’s Adjuvant (Sigma-Aldrich, USA). Animals were anaesthetized via halothane inhalation and antigen was administered IP (40 μg per dose to rats; mice, received 10 μg per dose). Antigens were injected on days 0 and 14. Sera were collected 21 days post-immunization, at euthanasia, and frozen at -20°C for subsequent analysis.

**Antigen-specific lymphocyte proliferation assay**

Spleens from euthanized rats and mice were aseptically removed, homogenized, and suspended in cold sterile buffer [PBS containing 1mM CaCl₂, 1mM MgCl₂, 5% (v/v) fetal bovine serum (FBS) (JR Scientific, USA), 100 U of penicillin per ml, 100 μg of streptomycin per ml and 0.25 μg of amphotericin B per ml (Fungizone; Life Technologies)]. A single cell preparation was then obtained by passage of the suspension through a nylon sieve (Becton Dickinson, USA), after which, the suspension was centrifuged at 100g for 10 minutes and the supernatant discarded.

Lysis of red blood cells was performed by addition of 0.5 ml 0.17M NH₄Cl, pH 7.65 for 1 minute and then diluted with 10 ml of the above buffer. Centrifugation was repeated (100g, 10 minutes at room temperature) and cells were washed 3 times in the
buffer. The final cell concentration was $10^6$ cells/ml in the same buffer, with viability determined by 0.1% trypan blue exclusion using a haemocytometer. Cells were cultured in RPMI medium containing 0.01 M HEPES, pH 7.4 (Life Technologies); 5x$10^{-5}$ M β-mercaptoethanol (BDH-Prolabo Chemicals, Australia) 2 mM L-glutamine (Life Technologies); 5% (v/v) FBS (JR Scientific) and penicillin-streptomycin-amphotericin B (Fungizone; Life Technologies). Ten-fold dilutions of antigens, peptides or Concanavalin A (1 µg/ml; Sigma-Aldrich) were dispersed into the wells of flat-bottom 96-well culture plates (Nunc; Thermo Scientific, USA). Blank control wells (without antigens) were also included in each run. A total of 2x$10^5$ cells were added to each well and plates were incubated at 37°C in 5% CO₂. On day 3, cells were pulsed with 25 µl/well with [methyl-³H]-thymidine (aqueous solution, specific activity of 20 Ci/mmol; PerkinElmer, Australia), which was diluted with culture media at a ratio of 20 µl/ml. Cells were harvested onto a printed filter mat A 1450-421 at 18-20 hours post-pulsing using a Tomtec Harvester 96 (Tomtec, USA). Radioactivity in the cells was measured in CPM via liquid scintillation (Wallac MicroBeta 1450 TriLux liquid scintillation counter; PerkinElmer). An average count for each specimen was calculated using triplicate cultures stimulated with each antigen or peptide. Similarly, blank control samples, lacking antigens were performed in triplicate. The experiments were repeated on four occasions. Cell proliferation was expressed as mean CPM ± standard deviation of antigen-stimulated cultures minus the CPM of antigen-free cultures. An additional antigen of E. coli protein components from a sham purification extraction was also included to control for the potential presence of trace components of E. coli proteins in the purified OMP26 peptide preparations. The concentration of E. coli proteins in these controls represented the maximum amount that may potentially exist in any purified OMP26 peptide
preparation. For each antigen the proliferative response for this control was also
deducted from the total lymphoproliferative response observed for each peptide to
determine the antigen specific response due solely to the recombinant OMP26
peptides. Data were analyzed for statistical significance by a fully factorial analysis
of variance (Macintosh Systat; Systat Software, USA) using log_{10}-transformed data.

**Antigen-specific ELISA**

Immunogenicity of OMP26 peptides to induce antibody responses was examined by
testing rat antipeptide sera, generated by immunization using individual OMP26
peptides, against the immunizing peptide and the whole OMP26 protein\(^{16}\). Antigen-
specific ELISA were performed and standardized as previously described by Kyd,
Cripps, Novotny, Bakaletz \(^{18}\). There was minimal background reactivity against an
*E.coli* antigen control for both immune and non-immune animals.

**Surface plasmon resonance analysis**

Serum and OMP26 protein and peptide interactions were assessed by SPR using the
Biacore 3000 instrument (GE Healthcare Life Sciences, USA) as previously
described\(^ {37}\). The Biacore system measures label-free binding of an analyte from a
continuous flow to an immobilized ligand in real time\(^ {38}\). Changes in refractive index
due to variation of the mass on the sensor chip, which results from the interaction of
the analyte and ligand, is expressed in relative Resonance Units (RU)\(^ {39}\). In brief,
reagent grade carboxymethylated dextran matrix (CM5) reagent grade sensor chips
(GE Healthcare Life Sciences) were activated with 35μl 400mM N-ethyl-N’-(3-
diethlyaminopropyl carbodiimide hydrochloride-100mM N-hydroxsuccinimide
solution at a flow rate of 5μl/min. OMP26 protein or peptides, suspended in 10 mM
sodium acetate, pH 4.5 were injected to immobilize approximately 0.1-0.2ng of
OMP26 protein or peptides per mm\(^2\) chip surface followed by deactivation of
unbound sites with 1.0 M ethanolamine-HCl, pH 8.5. 10 µl of each serum sample
diluted 1:5 with HBS-EP buffer (0.01 M HEPES, pH 7.4, 0.15 M NaCl, 3 mM EDTA,
0.005% [v/v] Surfactant P20; GE Healthcare Life Sciences), was exposed to the
immobilized protein or peptides at a flow rate of 5 µl/min. Between samples, the
sensor surface chip was regenerated with 10 µl injection of 50 mM NaOH. The
relative amount of antibody bound to each peptide was determined by comparison of
the change of resonance units between sample injection cycles.

Flow cytometry
Flow cytometry was used to determine whether antibodies induced by immunization
with OMP26 or peptides could bind to the surface of NTHi. To remove any potential
E. coli antibodies that may have cross-reacted with NTHi, sera were pre-absorbed
with E. coli prior to flow cytometry as previously described \(^\text{40}\).

NTHi-289 was grown as previously described \(^\text{16}\), harvested and washed twice
with sterile PBS by centrifugation (400g, 10 minutes at room temperature). The
bacterial cell pellets were subsequently incubated for 1 hour at 37°C with 200 µl of
pre-adsorbed sera from the immunized rats, which had been diluted 1:50 in PBS.
Following incubation, the bacterial cells were washed twice with sterile PBS as
before. Cells were then incubated with 200 µl of Alexa Fluor® 488 goat anti-rat IgG
(Life Technologies) diluted 1:50 in PBS, pH 7.4, for 30 minutes at 37°C. After
washing twice with sterile PBS, cells were re-suspended in 500 µl of PBS, pH 7.4,
and analyzed on a Flow Epic® XL-MCL Flow Cytometer (Beckman Coulter,
Australia).

Identification of surface exposed epitopes was determined by incubation of
NTHi-289 with seven antisera developed by immunizing rats (see above) with
individual OMP26 . As a negative control, PBS replaced the antibody and a bacterial
population of $8 \times 10^4$ bacteria was analyzed. Intact bacteria were incubated with a 1:50 dilution of antisera, revealed with Alexa Fluor® 488 goat anti-rat IgG and analyzed for fluorescent intensity by flow cytometry.
REFERENCES


8. Webb DC, Cripps AW. A P5 peptide that is homologous to peptide 10 of OprF from Pseudomonas aeruginosa enhances clearance of nontypeable Haemophilus


**TABLE 1.** Antibody reactivity of rat anti-OMP26 peptide against OMP26 protein and peptides.

<table>
<thead>
<tr>
<th>Immunogen</th>
<th>Homologous peptide assayed by</th>
<th>OMP26 protein assayed by</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ELISA&lt;sup&gt;a&lt;/sup&gt;</td>
<td>SPR&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>T1</td>
<td>440.3</td>
<td>502.2</td>
</tr>
<tr>
<td>T2</td>
<td>25.5</td>
<td>730.8</td>
</tr>
<tr>
<td>T3</td>
<td>209.1</td>
<td>1410</td>
</tr>
<tr>
<td>T4</td>
<td>2350.8</td>
<td>1896.6</td>
</tr>
<tr>
<td>T1a+T2</td>
<td>3915.7</td>
<td>1783.5</td>
</tr>
<tr>
<td>T2+T3</td>
<td>1119.5</td>
<td>1812.6</td>
</tr>
<tr>
<td>T3+T4</td>
<td>5290.9</td>
<td>807.4</td>
</tr>
</tbody>
</table>

<sup>a</sup> Antibody reactivities are expressed as IgG concentration in μg/ml. The assay was performed in duplicate, and the values represent the mean reactivity of immune sera from 3-5 rats. Non-immune sera did not react to any of the OMP26 peptides, and only slight reactivity to the OMP26 protein was detected at the dilution 1:25 (results not shown).
TABLE 2. Oligonucleotide primers used in PCR to construct OMP26 peptides

<table>
<thead>
<tr>
<th>Primer</th>
<th>Nucleotide position</th>
<th>Sequence $^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1F</td>
<td>1-21</td>
<td>5’GGC GGA TCC ATG AAA AAC ATC GCA AAA GTA 3’</td>
</tr>
<tr>
<td>T1aF</td>
<td>70-90</td>
<td>5’ GGC GGA TCC GAA GAA AAA ATT GCT TTC ATT 3’</td>
</tr>
<tr>
<td>T1HindR</td>
<td>145-165</td>
<td>5’ GGC AAG CTT AGG TTT AAA TTC AGC ATC AAG 3’</td>
</tr>
<tr>
<td>T2F</td>
<td>133-153</td>
<td>5’ GGC GGA TCC GTA GCA GAT AAA CTT GAT GCT 3’</td>
</tr>
<tr>
<td>T2HindR</td>
<td>280-300</td>
<td>5’ GGC AAG CTT GCG TTT TTG AAT ATC AGC TTG 3’</td>
</tr>
<tr>
<td>T3F</td>
<td>283-303</td>
<td>5’ GGC GGA TCC GCT GAT ATT CAA AAA CGC CAA 3’</td>
</tr>
<tr>
<td>T3HindR</td>
<td>415-435</td>
<td>5’ GGC AAG CTT TAA TAA TTT ACC ACG TTC TTC 3’</td>
</tr>
<tr>
<td>T4F</td>
<td>418-438</td>
<td>5’ GGC GGA TCC GAA CGT GGT AAA TTA TTA GAT 3’</td>
</tr>
<tr>
<td>T4HindR</td>
<td>571-591</td>
<td>5’ GGC AAG CTT TTT TTT CTC TTG TGC TTT TTC 3’</td>
</tr>
</tbody>
</table>

$^a$ Nucleotide position in the *omp26* gene from NTHi-289

$^b$ Recognition sites for restriction endonucleases *Bam*HI and *Hind*III are underlined
### TABLE 3. Amino acid sequences of the OMP26 peptides

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Amino acid sequence&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1&lt;sup&gt;b&lt;/sup&gt;</td>
<td>MKNIAKVTALALGIALASGYASAEEKIAFINAGYIFQPHDRQAVADKLD</td>
</tr>
<tr>
<td>T2</td>
<td>VADKLDAEFPVAEKLASKKEVDDKIAAARKVKEAKVAALKDAPRLQRQDIQKR</td>
</tr>
<tr>
<td>T3</td>
<td>ADIQKRQEENKLGAEDAELQKLMQEQDDKQVEFQAQNEKRQAEERKLL</td>
</tr>
<tr>
<td>T4</td>
<td>ERGKLLDSIQTATNLLARAKGYTYVLDANSVVFAVEGKDITEEVLKSIKASEKAFK</td>
</tr>
<tr>
<td>T1a+T2</td>
<td>EEKIIFAGYIFQHPDRQAVADKLDAEFPVAEKLASKKEVDDKIAAARKVKEAKVAALKDAPRLQRQDIQKR</td>
</tr>
<tr>
<td>T2+T3</td>
<td>VADKLDAEFPVAEKLASKKEVDDKIAAARKVKEAKVAALKDAPRLQRQDIQKRQEEINKLGAEDAELQKLMQEQDDKQVEFQAQNEKRQAEERKLL</td>
</tr>
<tr>
<td>T3+T4</td>
<td>ADIQKRQEENKLGAEDAELQKLMQEQDDKQVEFQAQNEKRQAEERKLLDSIQTATNLARAKGYTYVLDANSVVFAVEGKDITEEVLKSIKASEKAEFQ</td>
</tr>
</tbody>
</table>

<sup>a</sup> amino acid sequence from NTHi-289<sup>17</sup>

<sup>b</sup> synthetic peptide
FIGURE 1.

[Diagram showing [³H]-Thymidine incorporation (CPM) x 10^3 for immune and non-immune groups across different peptides: OMP26, T1, T2, T3, T4, T1a+T2, T2+T3, T3+T4. Bars indicate statistical significance with * for p < 0.05, ** for p < 0.01, and *** for p < 0.001.免疫 and non-immune groups are represented by open and filled squares, respectively.]
FIGURE 2.

[Diagram showing the incorporation of [3H]-thymidine (CPM) for different peptides, with bars for immune and non-immune conditions.]

immune  non-immune
FIGURE 3.
FIGURE 4.

(A) Antigen specific IgG (µg/ml)

(B) Peptide

(C) Resonance units (RU)

(D) Peptide

immune  non-immune
FIGURE 5.
FIGURE 6.

![Bar graph showing resonance units (RU) for different peptides (OMP26, T1, T2, T3, T4, T1a+T2, T2+T3, T3+T4). The graph compares immune and non-immune responses.]

- □ immune
- ☑ non-immune
FIGURE 8.

OMP26 with leader peptide

1  T1
1  55  T2
45  100  T3
95  145  T4

1  100  T1

24  100  T1a+T2

45  145  T2+T3

95  197  T3+T4
FIGURE LEGEND

Figure 1:
Antigen-specific proliferative responses to T1, T2, T3, T4, T1a+T2, T2+T3 and T3+T4 OMP26 peptides from OMP26-primed and naive DA rats. Splenic lymphocytes from immunized and non-immunized rats were cultured with individual OMP26 peptides or Concanavalin A (positive control) at concentrations of 10 μg/ml and 1 μg/ml for 3 days. Proliferative responses to peptides at 10 μg/ml are shown and values presented are expressed as mean counts per minute (CPM) ± standard deviation. Background lymphoproliferative responses to the negative E. coli protein control have been subtracted. Significance shown as ***P<0.001, **P<0.02, * P<0.05 compared to naïve lymphocytes stimulated with the same peptide on log10 transformed data.

Figure 2:
Lymphoproliferative responses induced by OMP26 protein by rat lymphocytes obtained after immunization with OMP26 peptides. Lymphocytes from immune and non-immune rats were stimulated with whole OMP26 protein (10μg/ml). Background lymphoproliferative responses to the negative E. coli control have been subtracted. Only lymphocytes from animals immunized with the T3+T4 peptide demonstrated a significant response to OMP26 protein stimulation compared with non-immunized rats (***P<0.001, **P<0.02). The OMP26 control showed a significant response (***P<0.001). Values presented are expressed as mean counts per minute (CPM) ± standard deviation. NI = non immune.
Figure 3:  
Binding of anti-OMP26 peptide sera to intact NTHi-289 cells. The x axes represent the levels of fluorescence and the y axes represent the number of cells counted. Intact bacteria were incubated with a 1:50 dilution of antisera, stained with Alexa Fluor® 488 goat anti-rat IgG and analyzed for intensity of green fluorescence by Flow Cytometry. The gray areas represent fluorescence in the absence of antibody. Blue lines represent the negative control of bacterial cells incubated with non-immune sera. The results with each immune sera are indicated in red. The total population analyzed was 8x10^4 bacteria.

Figure 4:  
Antibody reactivities to the OMP26 peptides of sera from mice immunized with OMP26 protein using the IPP/IT (Panel A, C) and IP (Panel B, D) immunization regimens and assayed by ELISA (Panel A, B) and SPR (Panel C, D). Immune and non-immune sera collected from 3-5 mice were pooled and assayed in duplicate for antibody binding against the whole OMP26 protein and peptides. Antibody reactivities are expressed as IgG concentration in µg/ml in ELISA and Resonance Units (RU) in SPR. No specific antibodies to OMP26 peptides were detected in non-immune sera at the lowest sample dilution (1:25) by ELISA.

Figure 5:  
Antibody reactivities to the OMP26 peptides of sera from rats immunized with OMP26 protein and non-immune DA rats, using the IPP/IT (Panel A, C) and IP (Panel B, D) immunization regimens and assayed by ELISA (6A, B) and SPR (Panel
Immune and non-immune sera collected from 3-5 rats were pooled and assayed in duplicate for antibody binding against the whole OMP26 protein and peptides. Antibody reactivities are expressed as IgG concentration in μg/ml in ELISA and Resonance Units (RU) in SPR. No specific antibodies to OMP26 peptides were detected in non-immune sera at the lowest sample dilution (1:25) by ELISA.

Figure 6:
Antibody reactivities to the OMP26 peptides of sera from chinchillas immunized with OMP26 protein. Pooled immune and non-immune sera were tested against all OMP26 peptides immobilized on the biosensor chips and assayed by SPR. Antibody reactivities are expressed as Resonance Units (RU).

Figure 7:
Summary of relative reactivity score was calculated against the OMP26 response. The OMP26 response was assigned a relative reactivity score of 5 and the responses of the other OMP26 peptides were scored against the OMP26 reactivity such that a response of 10-20% was scored as 1, 20-40% as 2, 40-60% as 3, 60-80% as 4, 80-100% as 5.

Figure 8:
Schematic diagram of OMP26 peptides. Numbering is based on the translated amino acid sequence of the full-length OMP26 from NTHi-289.