Mucosal immunization with the Moraxella catarrhalis porin M35 induces enhanced bacterial clearance from the lung: a possible role for opsonophagocytosis

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Moraxella catarrhalis is a significant cause of respiratory tract infection against which a vaccine is sought. Several outer membrane proteins are currently under investigation as potential vaccine antigens, including the porin M35. We have previously shown that the third external loop of M35 was immunodominant over the remainder of the protein for antibody produced in mice against the refolded recombinant protein. However, as this loop is predicted to fold inside the porin channel we also predicted that it would not be accessible to these antibodies when M35 is expressed on the surface of the bacteria in its native conformation. This study investigated the functional activity of antibodies against M35 and those specific for the loop 3 region of M35 in vitro and in vivo. Antisera from mice immunized with M35 or the loop 3-deletion, M35loop3−, recombinant proteins were not bactericidal but did have enhanced opsonic activity, whereas antibodies raised against the loop 3 peptide were not opsonizing indicating that the immunodominant loop 3 of M35 was not accessible to antibody as we had previously predicted. Mucosal immunization with M35, M35 that had an antigenically altered loop 3 [M35(ID78)] and M35loop3− enhanced the clearance of M. catarrhalis from the lungs of mice challenged with live M. catarrhalis. The in vivo clearance of bacteria in the mice with the M35-derived protein constructs correlated significantly (p < 0.001) with the opsonic activity assessed in an in vitro opsonophagocytosis assay. This study has demonstrated that the immunodominant B-cell epitope to loop 3 of the M. catarrhalis outer membrane protein M35 is not associated with immune protection and that M35-specific antibodies are not bactericidal but are opsonizing. The opsonizing activity correlated with in vivo clearance of the bacteria suggesting that opsonizing antibody may be a good correlate of immune protection.

Keywords: Moraxella catarrhalis, porin, mucosal immunization, opsonophagocytosis, respiratory tract

INTRODUCTION

Moraxella catarrhalis is a Gram-negative bacterium that is often the causal organism for respiratory tract infections such as otitis media, sinusitis, and exacerbations of chronic obstructive pulmonary disease (COPD; Murphy, 1996; Karalus and Campagnari, 2000; Verduin et al., 2002). On rare occasions this bacterium can cause more serious diseases such as meningitis and septicaemia (Meyer et al., 1995; Daoud et al., 1996). Additionally, the presence of M. catarrhalis appears to influence the pathogenicity of other respiratory pathogens, such as Haemophilus influenzae (Armbruster et al., 2010) and Streptococcus pneumoniae (Krishnamurthy et al., 2009). A vaccine against M. catarrhalis is sought and a number of outer membrane proteins are currently being characterized and evaluated for their potential as vaccine antigens (Murphy, 2005, 2009; Massa et al., 2009; Mawas et al., 2009). We have previously reported the characterization of an OMP from M. catarrhalis, designated M35, that we are investigating for its functional role and potential as a vaccine antigen.

Our initial characterization of M35 suggested that it possesses the desired characteristics of a vaccine antigen in that it is highly conserved, surface exposed, and immunogenic in an animal model (Easton et al., 2005). Subsequent studies confirmed a high level of conservation of M35 across strains, as well as demonstrating an important role for M35 in resistance to aminopenicillins (Jetter et al., 2009) and involvement in several stress responses. M35 shares homology with the family of outer membrane porins and is likely to conform to the usual tertiary structure of porins in which the third of eight external loops folds into the lumen of a channel formed by 16 anti-parallel β-sheets (Seshadri et al., 1998). In the predicted structure of M35 the third external loop is one of three loops that are much longer than the other loops (Figure 1). M35 contains an immunodominant epitope in the third external loop.
and we hypothesized that if loop 3 folded into the porin channel then it would not be accessible to antibody (Easton et al., 2005). By directing the majority of the antibody response against an inaccessible epitope, the bacteria could protect this protein from the host’s immune system and subsequent immunological selective pressure. This could potentially explain why M35 is so highly conserved, while other similar OMPs tend to have variable surface regions in response to selective pressure from the host’s immune system (Haase et al., 1991; Bash et al., 1995; Schirmer, 1998; Mes and van Putten, 2007).

M35 was also shown to have epitopes that are accessible to antibody binding on the surface of the bacterium (Easton et al., 2005). If the above hypothesis is correct, these surface epitopes would not include loop 3. As part of identifying the specificity of antibody to loop 3 it was also found that a single amino acid substitution in loop 3 (in the isolate ID78LN266) was sufficient to significantly reduce recognition of M35 by antibody. In this context, both the conformational significance of the loop and single point mutations within immunodominant loop regions have been identified as affecting antibody binding for the P2 porin of non-typeable Haemophilus influenzae (NTHi; Duim et al., 1993; Yi and Murphy, 1997). The lack of M35-specific antibody binding to the surface of the variant isolate implied that the majority of antibodies raised in mice against M35 were specific for loop 3 (where the mutation is located) and that loop 3 might be surface exposed. Such a result would suggest that either the predicted folding of M35 was incorrect or that loop 3 is either partially or fully outside the channel in M35, which would be highly unusual for this type of an outer membrane porin.

The aims of this study were threefold. Firstly, to further investigate the surface accessibility of antibodies directed against M35, specifically the loop 3 region. Secondly, to determine the bactericidal and opsonizing activity of M35-specific and loop 3-specific antibodies in vitro. Thirdly, to determine the efficacy of M35 as a protective antigen in a mouse mucosal immunization pulmonary clearance model (Kyd et al., 1998, 1999; Murphy et al., 1998).

MATERIALS AND METHODS

CLONING AND EXPRESSION OF M35-DERIVED PROTEIN CONSTRUCTS

Three recombinant proteins based on M35 were constructed: (1) M35(ID78), the full M35 protein based on the gene from the variant isolate ID78LN266; (2) L3DHFR, loop 3 with dihydrofolate reductase (DHFR) as a carrier protein; and (3) DHFR alone as a control. M35(ID78) contains two amino acid substitutions in comparison to the consensus M35 sequence, one of which was found previously to be antigenically significant (Easton et al., 2005). In addition to these newly constructed recombinant proteins, the consensus M35, the mature M35 (M35mat), and M35 without the loop 3 region (M35loop3−) were also used in this study (Easton et al., 2005). The relative structures of the recombinant proteins used in this study are represented diagrammatically in Figure 1. All recombinant proteins expressed a 6×His tag sequence at the N-terminal and were produced by expression in M15 [pRep4] Escherichia coli (Qiagen) with purification by nickel-nitritriacetic acid column chromatography according the manufacturer’s protocol for this expression system under denaturing conditions (8 M urea). M35 and the variant forms were refolded as previously described (Watanabe, 2002; Easton et al., 2005).

M35(ID78) was cloned by amplifying the M35 gene sequence from genomic DNA extracted from the ID78LN266 isolate of M. catarrhalis, as described previously for the consensus M35 using the pQE30 vector (Qiagen) (Easton et al., 2005). L3DHFR was produced by amplifying the gene sequence coding for the loop 3 region of the M35 gene from genomic DNA extracted from the 4223 isolate of M. catarrhalis using primers DEM35L3R (antisense) GCCCTGCAGATTGTTGGCACG and DEM35L3F (sense) GCCAGATCTATTGATGACAGTGTTG. These primers introduced SphI and PstI restriction enzyme recognition sites at either end of the PCR product that enabled directional insertion into the vector. The pQE40 vector (Qiagen) was used because this vector contains the coding sequence for DHFR immediately preceding the multiple cloning site and the recombinant protein produced therefore contains the carrier at the N-terminal end of the construct.

PRODUCTION OF HYPERIMMUNE MOUSE SERUM

Antisera was raised for each of the M35-derived proteins and whole killed M. catarrhalis in male BALB/c mice. Mice were immunized intraperitoneally three times at weekly intervals with 10 μg of each of the M35-derived protein constructs or 10⁸ whole killed M. catarrhalis emulsified with an equal volume of incomplete Freund’s adjuvant (IFA). The antibody concentration (total IgG) was measured by enzyme linked immunosorbent assay (ELISA)
against the specific antigen. The immunization experiments were approved by the University of Canberra Committee for Ethics in Animal Experimentation and the Central Queensland University Animal Ethics Committee.

ENZYME LINKED IMMUNOSORBENT ASSAY

Enzyme linked immunosorbent assay was used to measure specific IgG and IgA as described previously (Kyd et al., 1999), however the volumes were reduced to 50 μL and the assay was developed using Zymed™ 3,3′,5,5′-tetramethylbenzidine (TMB) single solution (Invitrogen). The ELISA plates were coated with 0.5 μg of the relevant antigen for each group. The whole killed 4223 cells were sonicated prior to coating for the WKC group. The serum and bronchoalveolar lavage (BAL) were serially diluted starting at 1/5 and 1/2, respectively. The antibody concentration was measured for each mouse individually against the same antigen with which the mouse was immunized and the samples from the non-immunized mice were tested for antibody against both M35 and the whole M. catarrhalis cells. The limit of detection was approximately 0.2 μg/mL for IgG and 0.125 μg/mL for IgA.

FLOW CYTOMETRY

Binding of antibodies raised against the M35-derived protein constructs to the surface of M. catarrhalis cells was analyzed by flow cytometry. M. catarrhalis isolates 4223 and ID78LN266 were grown overnight on brain heart infusion agar (Oxoid) supplemented with 5% defibrinated horse blood, washed in sterile PBS with 0.5% BSA and adjusted to a concentration of 1 x 10^8 CFU/mL using the optical density (OD) at 400 nm. One hundred microliter aliquots of the cell suspensions were centrifuged and resuspended in the mouse anti-serum diluted to approximately 20 μg/mL with sterile PBS with 0.5% BSA and incubated for 1 h at 37°C to allow the antibodies to bind. The bacteria were centrifuged and resuspended in a 1/200 dilution of goat anti-mouse IgG AlexaFluor488 (Molecular Probes, Invitrogen) and incubated for 1 h at 37°C. The labeled cells were washed twice in sterile PBS before analysis with a flow cytometer (Coulter XL-MCL, Beckman Coulter). About 100,000 events were counted and binding of antibodies was assessed by observation of the curve shift in comparison to bacteria exposed to serum from non-immunized mice.

EVALUATION OF COMPLEMENT SENSITIVITY

The isolate 4223 was not used for the bactericidal and opsonophagocytosis assays because, like many other M. catarrhalis isolates, it is complement resistant. Five isolates from our collection that had originally been collected by nasopharyngeal swab from healthy children with asymptomatic carriage were tested because these isolates are more likely to be serum sensitive than isolates taken from infection (Hol et al., 1995). To assess the complement sensitivity of these isolates the culture and spot test described by Verduin et al. (1995) was used. The isolates were grown overnight with shaking at 37°C in Mueller–Hinton broth (Oxoid). The bacteria were pelleted by centrifugation and resuspended in veronal buffered saline (VBS2+; 5 mM barbitone, 145 mM NaCl, 0.15 mM CaCl2, 0.5 mM MgCl2, pH 7.4). The concentration was adjusted to approximately 1.5 x 10^8 CFU/mL using the OD at 400 nm. One hundred microliter of this suspension was then spread on 5% sheep agar plates (Oxoid) and allowed to absorb. Fifty microliter of guinea pig serum, both active and heat-inactivated (30 min at 56°C), was spotted in duplicate on to each plate and the plates were immediately incubated at 37°C. After overnight incubation the bacterial growth within the spots was evaluated. This assay was repeated twice and the 2:03KAL isolate was chosen because of consistency in the active serum inhibiting growth and the heat-inactivated serum not inhibiting its growth to any detectable degree. M35 is very well conserved across diverse isolates (Easton et al., 2005), however as this isolate had not been included in our previous study, the expression of M35 and its antigenic conservation with 4223 was confirmed by western blot.

BACTERICIDAL ASSAY

The bactericidal activity of antibodies raised against the M35-derived proteins was assessed using the method described by Murphy et al. (1998). Serum samples from intraperitoneally immunized mice were heat-inactivated by incubation at 56°C for 30 min before use. The serum was also diluted with PBS to give an equivalent concentration of antibody as determined by ELISA, which was necessary because the purpose of the experiment was to compare the bactericidal activity of the antibodies raised against the different proteins rather than the efficacy of immunization with these proteins. The final antibody concentration of each sample was approximately 20 μg/mL.

Moraxella catarrhalis isolate 2:03KAL was grown in brain heart infusion (BHI) broth at 37°C with shaking at 220 rpm until the OD at 600 nm was approximately 0.2, indicative of early log-phase growth. The early log-phase broth culture was diluted 1/4000 with sterile PCGM saline (3.26 mM NaH2PO4, 12.8 mM NaHPO4, 125 mM NaCl, 1.25 mM CaCl2, 0.5 mM MgCl2, 1% gelatin). Twenty microliter of this cell suspension was added to each well of the microtiter plate with 10 μL of sterile PBS. Ten microliter of heat-inactivated serum sample was added to each well, with each serum sample being tested in triplicate. Serum from mice immunized with whole formalin-killed M. catarrhalis 4223 cells (WKC) was included as a positive control. Control wells without serum received sterile PBS instead of serum. The samples were mixed gently by pipetting and incubated for 30 min at room temperature to allow the antibodies present in the mouse serum to bind to the bacteria. Ten microliter of guinea pig serum was then added to each well as the complement source and the cells were incubated for a further 30 min at 37°C to allow complement mediated killing of the bacteria. No complement control wells received sterile PBS instead of the guinea pig serum. Two hundred microliter of sterile PCGM saline was added to each well and 50 μL aliquots were spotted on to CBA plates in duplicate to enable quantification of live bacteria. The plates were incubated overnight and the number of colony forming units in each spot was counted.

The bacterial counts were compared with the negative control wells that did not have mouse serum, complement source or both. Serum from non-immunized mice was also included as a negative control. A viability control was included that was titrated at the beginning of the incubation period so that survival during the incubation time could be assessed. A positive result is considered...
killing that reduces recovery to \(\leq 50\%\) of the recovery from the negative control (Murphy et al., 1998).

**OPSONOPHAGOCYTOSIS ASSAY**

An assay to measure the ability of antibodies raised against the M35-derived proteins to opsonize *M. catarrhalis* and thereby promote phagocytosis of the bacteria was developed with reference to similar assays used to analyze opsonization of other species of bacteria (Aase et al., 1995; Delvig et al., 1997a; Martinez et al., 1999; Jomaa et al., 2005a). *M. catarrhalis* 2:03KAL was grown overnight on CBA plates, harvested, washed twice in sterile PBS and fluorescently dyed with 5-chloromethylfluorescein diacetate (CMFDA) using the method suggested by the manufacturer (Molecular Probes). Approximately 1 × 10^10 CFU was harvested from a CBA plate, washed in PBS, and resuspended in 400 μL of 5 μM CellTracker Green CMFDA diluted in BHI media. The bacteria were incubated for 30 min at 37°C, pelleted again by centrifugation, resuspended in 1 mL BHI media, and again incubated for 30 min at 37°C. The bacteria were washed five times in Hank’s buffered salt solution (HBSS) with 0.2% BSA and after the final wash their concentration was adjusted to 2 × 10^6 CFU/mL by estimation from the OD at 400 nm. Twenty microliter of the bacterial suspension was mixed with 10 μL of mouse serum and incubated for 30 min at 37°C with gentle shaking (150 rpm). The serum samples were diluted to approximately the same relative antibody concentration as determined by ELISA (approximately 20 μg/mL) so that the results would reflect the opsonizing ability of the antibodies rather than to differences associated with titer to the specific antigens.

Ten microliter of guinea pig serum was added as an exogenous complement source and the mixture was incubated for a further 15 min at 37°C with gentle shaking. Mouse macrophages (RAW 264.7) were washed twice with HBSS and once with HBSS containing Ca^{2+} and Mg^{2+} and the concentration of cells was adjusted to 2.5 × 10^6 cells/mL. Forty microliter of this macrophage suspension was added to the bacterial suspension. The mixture was incubated for 30 min at 37°C with gentle shaking to allow phagocytosis of the bacteria by the macrophages. The cells were diluted with 200 μL of HBSS with Ca^{2+} and Mg^{2+} with 0.2% BSA, 50 μL of 0.5 mg/mL trypan blue was added to quench the fluorescence of extracellular bacteria and the cells were kept on ice before analysis using a flow cytometer (Beckman Coulter XL-MCL).

The relative fluorescence of the macrophages was used to assess the degree of phagocytosis in each sample. A higher percentage of fluorescent macrophages indicated a higher percentage of these cells had phagocytosed bacteria. Controls were included that received buffer instead of the mouse serum and serum from either non-immunized mice or mice immunized with whole killed *M. catarrhalis* 4223 (WKC) were included as negative and positive controls, respectively.

**MOUSE LUNG CHALLENGE MODEL WITH MUCOSAL IMMUNIZATION**

The mucosal immunization regime consisted of an intra-Peyer’s patch (IPP) immunization followed 14 days later by an intratracheal (IT) boost, following the method developed by Kyd et al. (1999). Each mouse received 10 μg of recombinant protein antigen or 10^8 CFU of formalin-killed bacteria in each immunization. The mice were immunized with one of the following antigens: (1) whole killed *M. catarrhalis* 4223; (2) M35; (3) M35mat; (4) M35loop3−; (5) L3DHFR; and (6) DHFR. Five male BALB/c mice were included in each immunization group and a non-immunized control group was included in each experiment. Protein concentrations were determined with a micro BCA assay kit (Pierce) and adjusted to 2 mg/mL. Proteins were concentrated using Amicon Ultra-15 Centrifugal Filter Units (Millipore) with a molecular weight limit of 5 kDa and final adjustments were made by dilution with 10 mM Tris–Cl, pH 7.4. As the M35-derived proteins precipitate without the presence of a denaturing agent or detergent, urea was added to final concentration of urea of 2.5 M.

Pulmonary challenge with 1 × 10^7 CFU live *M. catarrhalis* 4223 was carried out 21 days after the IPP immunization, following the protocol described by Kyd et al. (1999). This experiment was undertaken with the approval of the Central Queensland University Animal Ethics Committee.

**STATISTICAL ANALYSIS**

For the opsonophagocytosis assay triplicate values for the percentage of macrophages with fluorescence greater than the background for each sample were averaged and compared by one-way analysis of variance (ANOVA) with Tukey’s multiple comparison test. Values for bacterial recovery and antibody concentrations for each immunization group were compared to the non-immunized control group using a one-way ANOVA followed by Bonferroni’s multiple comparison test. Correlation of results was assessed using a Pearson’s test. All analyses were performed using GraphPad Prism 4.0b.

**RESULTS**

**ANTIBODY-BINDING TO SURFACE EPITOPES OF M35**

Surface accessibility of the recombinant M35 proteins and loop 3 of M35 was investigated by determining the ability of antibody to bind to surface epitopes of M35 in *M. catarrhalis* isolates 4223 and ID78LN266 by flow cytometry (Figure 2). Antibodies raised against M35 bound to the cell surface of *M. catarrhalis* as demonstrated by a small right-shift in the relative fluorescence compared to the control non-immune serum. While this shift was small, it was reproducible and consistent with our previous results (Easton et al., 2005). Antibodies raised against M35(ID78) and M35loop3−, respectively, caused similar sized right-shifts to that seen with anti-M35 serum. Antibodies raised against L3DHFR and DHFR caused smaller right-shifts than those seen with the anti-M35 serum. The lack of difference between the right-shifts seen with L3DHFR and DHFR antisera indicates that any binding was not due to loop 3-specific recognition by the serum antibodies.

The binding of antibodies to *M. catarrhalis* differed between the strains of bacteria used. Antibodies raised in mice immunized with isolate 4223 demonstrated a greater ability to bind to whole killed cells of strain 4223 than that seen when tested against whole killed cells from strain ID78LN266. This may indicate that these two isolates differ substantially in the availability of outer membrane protein epitopes that are targeted by the antibodies. It was also noted that the background staining with naïve serum was higher.
with isolate ID78LN266 than isolate 4223 (data not shown), suggesting that isolate ID78LN266 may express surface components with epitope sequences homologous to the normal flora of the mice. This difference between the strains may partly explain the lack of significant binding between the naïve sera and M35-specific antibodies to isolate ID78LN266 in our previous study (Easton et al., 2005).

**BACTERICIDAL ACTIVITY OF M35-SPECIFIC ANTIBODIES**

In an *in vitro* bactericidal assay, a greater than 50% killing of bacteria was achieved using anti-serum raised against the whole formalin-killed *M. catarrhalis* cells (WKC; Figure 3). None of the antisera raised against loop 3 or the M35-derived proteins were bactericidal (Figure 3).

**ENHANCEMENT OF OPSONOPHAGOCYTOSIS BY M35-SPECIFIC ANTIBODIES**

The ability of antibodies raised against each of the recombinant proteins to opsonize and therefore enhance phagocytosis of live *M. catarrhalis* cells was tested by determining the percentage of macrophages that had phagocytosed clearly labeled live *M. catarrhalis*. Using flow cytometric analysis the fluorescent intensity of the macrophages was assessed following incubation with fluorescently labeled bacteria in the presence or absence of antisera (Figure 4). Bacteria exposed to the 4223 WKC anti-serum as a positive control were readily phagocytosed with nearly 50% (48 ± 1.3%) of macrophages showing greater than background fluorescence (p < 0.001). An average of 9.8 ± 0.2% of macrophages phagocytosed bacteria that were exposed to serum from non-immunized mice and this was a similar percentage to the control sample exposed to sterile PBS instead of serum (10.8 ± 0.84%). Antibodies against L3DHFR and DHFR both failed to enhance phagocytosis, with results of 11.8 ± 0.6 and 10.7 ± 0.4%, respectively. Antibodies raised against M35, M35(ID78), and M35loop3− all enhanced phagocytosis with a small significant increase in fluorescence of 15.6 ± 0.8% (p < 0.05), 17.2 ± 1.6% (p < 0.01), and 16.4 ± 2.3% (p < 0.01), respectively (Figure 4).

**BACTERIAL RECOVERY FROM MOUSE LUNGS AFTER CHALLENGE WITH LIVE M. CATARRHALIS**

The effect of mucosal immunization with each M35-derived construct was assessed by quantification of *M. catarrhalis* recovered from the lungs of mice 4 h after challenge by intra-tracheal inoculation with live bacteria. The quantity of live bacteria recovered from the BAL fluid, lung homogenate and in total from mice mucosally immunized with each antigen is shown in Figure 5. When compared to the non-immunized mice the number of bacteria recovered from mice mucosally immunized with the formalin-killed *M. catarrhalis* 4223 cells (WKC) were significantly lower in both the BAL (p < 0.001) and lung homogenate (p < 0.01).
The bacterial recovery in the lung homogenate from mice mucosally immunized with M35, M35(ID78), and M35loop3− was also significantly lower than from the non-immunized mice (p < 0.01), however in the BAL fluid, enhancement of bacterial clearance only reached significance when M35(ID78) was the immunizing antigen (p < 0.01). When BAL and lung values were added to give total bacterial recovery (Figure 5A) all three groups [M35, M35(ID78), and M35loop3−] had significantly lower numbers of bacteria recovered in total than in the non-immunized control group (p < 0.01). Bacterial recovery from the mice immunized with M35mat, L3DHFR, and DHFR did not differ significantly from the non-immunized control groups in either the BAL fluid, lung homogenate, or total recovery (p > 0.05).

**ANTIBODY TITER AFTER MUCOSAL IMMUNIZATION**

IgG and IgA concentrations in the BAL fluid of immunized mice were very low and were undetectable in many samples (Table 1). The results from individual mice were found to be variable within the immunization groups, as indicated by the large SDs shown in Table 1. The concentration of IgG in the BAL tended to reflect the concentration in the serum for each individual animal. The non-immunized mice did not have any detectable anti-M. catarrhalis antibody. Serum IgG antibody to loop 3 was very low when detected and was undetectable in most of the mice immunized with L3DHFR and none of those immunized with DHFR (Table 1).

**WHITE BLOOD CELL RECOVERY FROM MUCOSALLY IMMUNIZED MICE**

The recruitment of white blood cells to the lungs is associated with both innate and acquired immune responses to infection. In general, the total white blood cells recovered at 4 h post challenge is significantly higher in the WKC immunized groups when compared with the non-immune groups and this pattern was replicated in this study (Figure 6A). The total number of white blood cells recruited to the lungs and counted in the BAL was highest in the L3DHFR group (p < 0.001; Figure 6A). The most abundant cell type were neutrophils (Figure 6B) followed by monocytes/macrophages (Figure 6C) and in general the proportions followed the same trend as the total white blood cell counts for each group (Figure 6A), with the exception that there was significantly more monocyte/macrophage cells recovered from the WKC group compared to the non-immunized mice (p < 0.01).
Table 1 | Specific antibody levels in serum and BAL after mucosal immunization.

<table>
<thead>
<tr>
<th>Immunization group</th>
<th>Serum IgG (μg/mL)b</th>
<th>BAL IgG (μg/mL)</th>
<th>BAL IgA (μg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-immune</td>
<td>n.d. c</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>WKc</td>
<td>2.25 ± 1.07</td>
<td>0.34 ± 0.34</td>
<td>n.d.</td>
</tr>
<tr>
<td>M35</td>
<td>16.4 ± 25.6</td>
<td>0.25 ± 0.75</td>
<td>0.01 ± 0.04</td>
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<tr>
<td>M35(mat)</td>
<td>54.4 ± 42.6</td>
<td>0.49 ± 0.45</td>
<td>0.01 ± 0.03</td>
</tr>
<tr>
<td>M35(ID78)</td>
<td>75.8 ± 47.8</td>
<td>1.23 ± 0.95</td>
<td>0.02 ± 0.04</td>
</tr>
<tr>
<td>M35(loop3−)</td>
<td>52.9 ± 62.1</td>
<td>0.60 ± 0.65</td>
<td>n.d.</td>
</tr>
<tr>
<td>L3DHFR</td>
<td>0.064 ± 0.14</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>DHFR</td>
<td>n.d.</td>
<td>n.d.</td>
<td>n.d.</td>
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*a The antibody concentration was measured for each mouse individually against the same antigen with which the mouse was immunized and the samples from the non-immunized mice were tested for antibody against both M35 and the whole M. catarrhalis cells. DHFR was assayed against L3DHFR.

*b Values are mean and SD.

*c Not detected.

**CORRELATION BETWEEN BACTERIAL CLEARANCE AND OPSONOPHAGOCYTOSIS**

To determine if there was a direct relationship between the ability of the animals to clear the infection challenge and the opsonophagocytosis results, a Pearson’s test was conducted. There was a direct and significant correlation (Figure 7) between the total bacterial recovery from mice in each immunization group (Figure 5A) and the enhancement of opsonophagocytosis mediated by serum antibody raised against the same immunization antigens (Figure 4; \( p = 0.001 \)).

**DISCUSSION**

M35 is an OMP from M. catarrhalis that we have been investigating for its functional role and potential as a vaccine antigen. Our initial characterization of M35 suggested that it possesses the desired characteristics of a vaccine antigen in that it is highly conserved, surface exposed, and immunogenic in an animal model (Easton et al., 2005). Our previous study also indicated that the loop 3 region of M35 was immunodominant over the remainder of the protein when mice were immunized with recombinant refolded M35 protein and raised the possibility that this loop may not be accessible to antibodies in the native protein expressed on the surface of the bacteria. Thus a situation may exist where the majority of antibodies produced against M35 are not effective due to an inability to bind to the protein on the cell surface. This would also be also consistent with previous observations relating to other porin proteins such as the loop 3 of the E. coli OMP F porin protein which has a major B cell region within the loop 3 domain (Williams et al., 2000; Yamashita et al., 2008). To investigate this hypothesis we have assessed the functional activity of antibodies specific to M35 and its loop 3 region.

This study has established that whilst at least some of the antibodies raised against the full M35 protein are able to bind to the protein on the cell surface, those specific for loop 3 do not bind a surface accessible epitope. The functional assays have demonstrated that while M35-specific antibodies were not bactericidal, antibodies to intact M35 constructed proteins were opsonizing in vitro. Loop 3-specific antibodies were neither opsonizing nor bactericidal, providing further evidence that antibodies specific for this immunodominant epitope are unlikely to be effective against infection. While this is the longest of the external loop regions, the lack of surface recognition, or opsonizing activity by loop 3-specific antibody supports the proposed structural model that this loop folds into the channel formed by the protein (Easton et al., 2005). Bactericidal activity has been used to assess functional activity for other M. catarrhalis antigens. Human antibodies to MID,
Epstein et al., 2003) and mice antibodies dLOS, CD, and UspA (Hu et al., 2004) have been shown to be bactericidal in vitro. Therefore in the current study, antibody to antigens other than M35 were responsible for the bacterical killing by WKC antisera.

The efficacy of potential vaccine antigens against M. catarrhalis has been modeled using the murine mucosal immunization – pulmonary challenge model, in which effective immunization enhances the clearance of live bacteria from the lungs of mice (Murphy et al., 1998; Kyd et al., 1999). At present there is no correlate of protection for M. catarrhalis in this model and it is unknown whether antigens found to be effective in this model will also be effective in humans (Murphy, 2005, 2009). Attempts to determine the mechanism of protection in the mouse model have investigated the quantity, type and in vitro bactericidal capacity of antibodies produced by the mice, however none of these parameters has been shown to be definitive correlates of protection (Murphy et al., 1998; Kyd et al., 1999). For example, Murphy et al. (1998) found that while mucosal immunization led to the production of bactericidal antibody, intramuscular immunization did not, but both routes were similarly effective at enhancing clearance of the bacteria, suggesting that the bactericidal capacity of the antibodies was not necessarily a correlate of protection in an acute infection mode identical to that described in the current study. Differences between the naive and immune-induced infiltration of phagocytic cells can be an indicator of antigen specific upregulation of the host’s immune response to the infection. Therefore we hypothesized that immunization would enhance Fc-mediated phagocytic removal of live bacteria via enhanced opsonization. Sera from WKC immunized mice demonstrated the most significant opsonizing activity and these mice also showed the greatest level of enhanced bacterial clearance in the lung infection study. In contrast, the L3DHFR immunization was the least effective and the antisera was not opsonizing. Analysis of the correlation of the opsonic capacity of M35-specific antibodies with bacterial clearance in vivo showed a linear relationship, indicating a direct correlation between the two measures. This is an important finding in that it raises the question of whether opsonizing antibody would also be protective in humans, or whether it only appears to be protective in the mice due to the short time-frame of the challenge model and perhaps even just for this antigen. Also of interest is that there does not appear to be a relationship between the number cells recruited to the lung following challenge and clearance. However, if opsonizing antibodies are not induced a large cellular influx does not result in enhanced bacterial clearance from the lung. This is most striking for the L3DHFR data.

A hallmark of respiratory infections is the rapid recruitment of white cells to the bronchial space following infection. This recruitment into the lung is in response to bacterial lipopolysaccharide and a range of inflammatory mediators that are released by resident phagocytes and endothelial cells of the bronchus (Dunkley et al., 1995). Immunization with antigens from NTHi, Streptococcus pneumoniae and Pseudomonas aeruginosa have previously been reported to enhance the recruitment of white cells (both neutrophils and monocytes) when the lung is challenged with live microbes (Buret et al., 1994; Foxwell et al., 2001; Jomaa et al., 2005b). Furthermore, this recruitment is antigen dependent with immunization by WKC normally being the most effective (Jomaa et al., 2005b). However, in a previous study of M. catarrhalis, we did observe that immunization with purified OMP CD was as effective as WKC in enhancing the recruitment of white cells to the lung (Murphy et al., 1998). In the current study we observed that whilst WKC immunization was effective in enhancing the recruitment of monocytes to the lung, L3DHFR was actually more effective and enhanced the recruitment of neutrophils as well. The mechanism by which immunization is able to enhance the recruitment of white cells to the lung is likely T cell dependent (Khader et al., 2009). It is thought that antigen specific T cells are generated in the gut associated lymphoid tissue following immunization and home to the lung via the common mucosal immune system (Otczyk and Dunkley, 2011) where they secrete a range of cytokines and chemokines that result in the recruitment of further neutrophils (Clancy and Dunkley, 2011). In this scenario a plausible explanation for the significantly enhanced recruitment of cells after immunization with L3DHFR would be the presence of T cell epitopes within the construct.

Although the role of opsonizing antibody has been speculated upon (Murphy et al., 2003), the only study demonstrating the development of opsonizing antibody showed that antisera from 11 children with M. catarrhalis otitis media had increased opsonic activity in convalescent sera (34.6 ± 27.1) compared to acute sera (15.5 ± 6.7; p < 0.05; Faden et al., 1994). We are not aware of any other studies that have shown a correlation between antibody responses to a M. catarrhalis antigen and opsonic activity in the absence of bactericidal activity. It will be important to test in future studies whether this is specific to immunization with the M35-derived proteins or whether enhancement of in vitro phagocytosis is also correlated with protection after immunization with other M. catarrhalis antigens. A similar observation was reported for N. meningitidis, where the majority of PorB-specific IgG was not bactericidal but was strongly opsonic (Delvig et al., 1995, 1996, 1997b).

Anti-M35 serum demonstrated weak binding to M35loop3”, no binding to DHFR, weak binding to M35(ID78) but strong
binding to both M35 and L3DdFr. (data not shown). Anti-L3 sera showed strong recognition of the M35 proteins, demonstrating that while there are conformational differences between the L3 regions of the whole M35 and the L3DdFr construct, there is cross recognition of the epitopes in common between the peptide and the recombinant whole protein. Antibodies specific for M35, M35(ID78), and M35loop3− were all able to bind to the surface of the bacterium, meaning that the presence, absence, or alteration of loop 3 did not affect the recognition of surface epitopes of M35. However, loop 3-specific antibodies were unable to bind to M35 on the surface of M. catarrhalis, supporting the prediction that the loop 3 region would not be accessible to antibodies, as it was predicted that it folded into the porin channel (Easton et al., 2005). Analysis of the in vitro opsonophagocytic activity of the M35-specific antibodies revealed that antibodies specific for the loop 3 region were not able to enhance phagocytosis, presumably due to their inability to bind on the cell surface. Antibodies raised against M35, M35(ID78), and M35loop3− were all able to enhance in vitro phagocytosis, demonstrating that it was antibody directed at the surface region that was responsible for the increase in phagocytosis and also that deletion of loop 3 did not create alterations to the protein that affected production of antibody with opsonizing activity. It therefore appears that, unlike the N. meningitidis PorB where the immunodominant region was in loop 1 (Delvig et al., 1995, 1996, 1997b) and surface exposed, the majority of antibodies raised against M. catarrhalis M35 are against loop 3 and are not opsonizing due to specificity for an apparently hidden loop region. Preliminary studies with the native protein had demonstrated the efficacy of M35 as a protective antigen when used in mucosally immunized mice prior to pulmonary challenge with live M. catarrhalis (unpublished data). Since our subsequent investigations had suggested that the majority of antibodies produced following immunization were directed at loop 3, it was hypothesized that immunization with loop 3 alone would not be protective. Mice immunized with M35, M35(ID78), and M35loop3− cleared significantly more of the bacteria from their lungs than the non-immunized control mice, indicating that deletion of the loop 3 region from the protein did not reduce, or enhance, the efficacy of the M35-specific response to infection. Mice mucosally immunized with the loop 3 peptide were not able to enhance clearance of the bacteria, however, these mice produced very low levels of L3DdFr-specific IgG detectable only in the sera and no IgG or IgA was detected in the BAL. However, mice immunized with M35 that contains an antigenically different loop 3 M35(ID78) (Easton et al., 2005) or had the loop 3 region deleted (M35loop3−) had similar immune efficacy against infection, indicating that antibody generated against loop 3 from whole protein immunization was not involved in M35-specific bacterial clearance. Therefore, while clearance of M. catarrhalis in the acute mouse lung challenge model could be enhanced by mucosal immunization with M35, this was independent of loop 3 as predicted. Studies with NTHi OMP P5 found that the fourth surface exposed region of the P5 antigen from NTHi was immunodominant over the remainder of the protein and not protective, however, immunization with the third surface exposed region of P5 was protective (Novotny and Bakaletz, 2003). It is possible that a similar pattern could exist with M35, suggesting further investigation of the antigenic roles of specific surface loops of M35 is warranted.

CONCLUSION
This study has confirmed our prediction that the majority of antibodies raised against M35 that were specific for loop 3 are ineffective and unlikely to be associated with immune protection against infection. The study has found that there was a correlation between opsonizing antibodies associated with enhanced phagocytosis and in vivo enhancement of bacterial clearance in mucosally immunized mice. Immunization with M35 with or without loop 3 was effective in enhancing clearance in the mouse pulmonary challenge model, suggesting that the enhancement of opsonophagocytosis due to antibodies to other surface epitopes contributed to this response. The study suggests that opsonic antibodies are potentially important for enhanced clearance and the role of opsonic antibody to M. catarrhalis should be further investigated for other antigens.

AUTHORIZATION FOR THE USE OF EXPERIMENTAL ANIMALS
Animal studies were undertaken with the approval of the Central Queensland University Animal Ethics Committee and the University of Canberra Committee for Ethics in Animal Experimentation.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest. The porin protein known as M35 has been patented for vaccine use with the authors as named inventors.

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