Brief Definitive Report

Eliminating a Region of Respiratory Syncytial Virus Attachment Protein Allows Induction of Protective Immunity without Vaccine-enhanced Lung Eosinophilia

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Summary

In a murine model of respiratory syncytial virus disease, prior sensitization to the attachment glycoprotein (G) leads to pulmonary eosinophilia and enhanced illness. Three different approaches were taken to dissect the region of G responsible for enhanced disease and protection against challenge. First, mutant viruses, containing frameshifts that altered the COOH terminus of the G protein, were used to challenge mice sensitized by scarification with recombinant vaccinia virus (rVV) expressing wild-type G. Second, cDNA expressing these mutated G proteins were expressed by rVV and used to vaccinate mice before challenge with wild-type respiratory syncytial virus (RSV). These studies identified residues 193–205 to be responsible for G-induced weight loss and lung eosinophilia and showed that this region was not necessary for induction of protective immunity. Third, mice were sensitized using an rVV that expressed only amino acids 124–203 of the G protein. Upon RSV challenge, mice sensitized with this rVV developed enhanced weight loss and eosinophilia. This is the first time that a region within RSV (amino acids 193–203) has been shown to be responsible for induction of lung eosinophilia and disease enhancement. Moreover, we now show that it is possible to induce protective immunity with an altered G protein without inducing a pathological response.

Key words: respiratory syncytial virus • G protein • eosinophilia • vaccine • T helper cell

R espiratory syncytial virus (RSV) is the leading cause of bronchiolitis and viral pneumonia among infants and young children (1, 2). Primary infection with RSV usually occurs within the first or second year of life with children between the ages of 6 wk and 9 mo of age at the greatest risk of developing serious illness. RSV bronchiolitis is associated with the development of atopy and asthma in later years (3). Because bronchiolitis is the leading cause of hospital admissions for infants in the western world, development of a safe and effective vaccine is a high priority.

A cautious approach to RSV vaccine development has been taken since the failure of vaccine trials in the 1960’s. In these trials children were inoculated intramuscularly with formalin-inactivated RSV. After subsequent natural exposure to RSV, children who had been given the RSV vaccine had greater morbidity and mortality than control vaccinees (4–7). Postmortem examination of these children showed peribronchiolar infiltration and excess eosinophils in the lungs and blood (4, 7).

In a BALB/c mouse model, this type of enhanced pathology can be reproduced by scarifying mice with recombinant vaccinia virus (rVV) expressing only the G protein of RSV followed by intranasal inoculation of infectious RSV (8). The G protein is a highly glycosylated membrane protein thought to be responsible for viral attachment to the host cell. It is produced in both a secreted and membrane-bound form; the function of the secreted form is unknown, but it may function as a decoy for neutralizing antibodies. The COOH terminus of the G protein varies amongst the different strains of RSV, although in human RSV there is a conserved region (amino acids 164–176) that is believed to be the receptor attachment site (9).

In the BALB/c mouse model, vaccination with individual RSV proteins expressed in VVs generates different immunological and pathological responses upon intranasal exposure to RSV (2). Vaccination with rVV expressing the fusion protein (F) of RSV leads to the generation of CTLs and CD4+ cells with a Th1 type of phenotype (i.e., high IFN-γ and no IL-4 or IL-5), whereas rVV expressing the matrix protein (M2) generates a CTL response with little or no T helper response. By contrast, vaccination with rVV...
expressing the G protein leads to the generation of a Th2 type of response (less IFN-γ and some IL-4 and IL-5) with an eosinophilic influx into the alveolar space (8, 10, 11). This eosinophilic influx and vaccine-enhanced illness resembles the pathology found in the children from the 1960’s vaccine trials.

To aid the development of a safe and effective vaccine, this murine model of vaccine-enhanced illness has been used to investigate immunopathological mechanisms involved in RSV illness (2). Since eosinophilia depends on CD4+ T cell recognition of G (12, 13), we mapped G protein regions in BALB/c mice using RSV frameshift mutants and rVV expressing partial or mutant G proteins. These studies identified an 11-amino acid portion of the G protein that is involved in the generation of pulmonary eosinophilia and vaccine-enhanced weight loss, but is not essential for the induction of protective immunity.

Materials and Methods

Viruses. RSV and rVV were grown on HEp-2 cells and titrated as previously described (8). Frameshift mutants were isolated by selecting escape variants in vitro with anti-G monoclonal antibody (63G) (14). All stocks were free of mycoplasma contamination as determined by DNA hybridization (Genprobe Inc., San Diego, CA).

Construction of rVV Expressing G Protein Frameshift Mutants. pGEM-4 derived plasmids encoding the G proteins of frameshift mutants R63/1/2/3 and R63/2/4/8 have been described previously (15). The G protein inserts of the above plasmids were isolated by digestion with HindIII and E.coli, blunt-ended with Klenow polymerase, and ligated to site-directed mutagenesis as described (15). The G protein inserts of the above plasmids were isolated by digestion with HindIII and EcoRI, blunt-ended with Klenow polymerase, and ligated to Smal linearized pSC11 vector. CV-1 cells were infected with the WR strain of VV and immediately transfected with the pSC11-derived plasmids. Progeny virus was plaque purified three times on HUTK-124B cells in the presence of 15 μg/ml of bromodeoxyuridine to select for thymidine kinase-negative recombinants. The rVVG27 (expressing amino acids 124–203) and rVVTer (empty vector) vaccinia viruses were a gift of Dr. M. Trudel (Centre de Recherche en Virologie, Institut Armand-Frappier, Québec, Canada) and have been described previously (16). rVV contains the insert for the whole G protein and rVVβgal contains the LacZ gene inserted into the vaccinia vector as a negative control in all experiments (17, 18).

Mouse Infections. Anesthetized 8–10-wk-old female BALB/c mice (Harlan Olac, Bicester, UK) were scarified with 2 × 106 PFU recombinant vaccinia at the base of the tail. After 2 wk, mice were intranasally inoculated with 1.5 × 106 PFU/mouse of Long and RSV mutants or 3 × 106 PFU/mouse of RSV A strain.

Measurement of Eosinophilia in the Bronchoalveolar Lavage. 7 d after RSV infection, bronchoalveolar lavage (BAL) fluid was collected as previously described (19). In brief, individual mice were terminally anesthetized with pentobarbitone and bled via the femoral artery. Lungs were perfused six times with 1 ml of lignocaine in Eagle’s media (Sigma Chemical Co., Poole, UK). 200 μl of BAL fluid was cytocentrifuged onto glass slides and stained with Giemsa’s reagent for cytological analysis. Eosinophils were counted by flow cytometry as the proportion of the granulocytes compared with total cells and confirmed with microscopic examination of cytospin slides to distinguish between polymorphonuclear cells and eosinophils.

Titration of RSV from Mouse Lungs. Mice were scarified with recombinant vaccinia 14 d before challenge with whole RSV. 4 d after challenge with RSV, whole lungs were disrupted using glass homogenizers (Jencons, Leighton Buzzard, UK) in 1.3 ml RPMI (Sigma Chemical Co.) supplemented with 2 mM glutamine (GIBCO BRL, Paisley, Scotland), 100 U/ml penicillin, and 100 μg/ml streptomycin on ice. Homogenates were clarified at 10,000 g for 1 min. 50 μl of supernatant and twofold serial dilutions thereof were titrated on HEp-2 monolayers in 96-well plates and plaques were assayed as previously described (8). The theoretical limit of detection for this assay was 0.86 PFU/lung.

Statistical Analysis. Kruskal-Wallis tests were used to test for effects between groups and Mann-Whitney U tests were used to perform comparisons between the experimental and control groups. Data analysis was performed using SPSS statistical software.

Results

Mapping of the Eosinophilic Induction Using RSV Mutants. A series of RSV escape mutants were generated using a monoclonal antibody against the G protein (14). These mutants contain frameshift mutations generating G proteins with truncations and/or alterations in the COOH terminus of the protein (Fig. 1). Mice were scarified with rVV or rVVβgal (control construct) and 2 wk were later intranasally challenged with the parental Long strain or one of the different mutant viruses. 7 d after infection, BAL fluid was collected and the percentage of eosinophils in the BAL was assessed. As in primary infection, mice scarified with rVVβgal and challenged with either Long or mutant viruses showed no eosinophilia. However, mice scarified with rVVβgal and challenged with whole RSV showed marked pulmonary eosinophilia except for one mutant. Mutant 63/1/2/3 failed to induce eosinophilia. Mice scarified with rVVβgal and challenged with mutant 63/1/2/3 generated a low level of eosinophilia similar to mice scarified with rVVβgal (P = 0.86) (Fig. 1). The low level of eosinophilia observed in mice scarified with rVVβgal after intranasal challenge with 63/1/2/3 was significantly different from mice scarified with rVVβgal followed by either Long, 63/2/4/1, or 63/2/4/8 (P ≤ 0.01, 0.01, and 0.01, respectively). The differences between mutant 63/1/2/3 and 63/2/4/8 lie between amino acids 193 and 205 (Fig. 1). From these data, the portion of the G protein responsible for eosinophil induction can be localized to this region.

A possible explanation for this finding is that the large deletion in 63/1/2/3 altered viral infectivity and subsequent eosinophilia generation. Viral lung titers on days 4 and 7 after infection showed no differences between the parental Long strain virus and viral mutants (data not shown). Despite the large alteration of the COOH terminus of the G protein, all mutant viruses infected the lungs efficiently and replicated as well as Long strain virus and all were cleared by day 7.

Mapping of the Eosinophilic Antigen Using rVV. To complement the studies described above and ensure that the
Discussion

This is the first study to show that a single region within the G protein is responsible for enhanced illness and lung eosinophilia and that in its absence it is possible to induce protective immunity without pathology. The critical importance of this region for lung eosinophilia was evident both at the priming (rVV63/1/2/3) and challenge (63/1/2/4).

Eosinophilia was due only to sensitization to the attachment protein G, rVVVs expressing mutant G proteins were used to sensitize mice. Mice were scarified with recombinant vaccinia and challenged intranasally with wild-type RSV (A2 strain) after 14 d. The G protein of the A2 strain is 95% homologous to Long strain and is identical within region 193–205. Mirroring the first set of studies, no eosinophilia was observed in the BAL of mice scarified with rVV63/1/2/3 (Fig. 3 A). The level of eosinophilia in rVV63/1/2/3-vaccinated mice was not significantly different from rVVGβgal-primed mice (P=0.556) and was significantly reduced compared with that seen in mice primed with rVVG, rVV63/2/4/1, or rVV63/2/4/8 (P ≤0.02, 0.02, and 0.05, respectively). To confirm that the lack of eosinophilia observed in mutant rVV63/1/2/3 mice was not due to a loss of stability or secondary structures, mice were scarified with recombinant vaccinias and challenged intranasally with wild-type RSV. All mice vaccinated with either of the frameshift mutants (rVV63/2/4/8 or rVV63/1/2/3) or with wild-type G (rVVG) were protected from RSV infection at day 4 (Fig. 5) and no virus was recovered from the lungs of any mice at day 7 (data not shown). These results confirm that it is possible to protect mice from subsequent RSV infection without inducing lung eosinophilia.

Protection against RSV challenge. To examine whether rVV63/1/2/3 could still provide protection even though it does not induce eosinophilia, mice were scarified with recombinant vaccinias and challenged intranasally with wild-type RSV. Data shown is representative of four experiments and each bar represents the mean percentage of individual eosinophil counts in each group (n=4). The significance levels between rVVG and rVVβgal using Mann-Whitney paired comparisons test are indicated by **. The significance levels between rVVG and rVVβgal vaccinated mice are as follows: Long, P=0.03; 63/2/4/1, P=0.03; 63/2/4/8, P=0.03; 63/1/2/3, P=0.86.

Figure 1. The primary structure of the G protein from RSV (Long strain) and the mutants used in this study (14). The stippled region indicates the transmembrane domain and hashed boxes denote amino acids changed due to frameshift mutations. The location of the intracellular (IC), transmembrane (TM), mucinoid I, mucinoid II, and conserved domains are indicated.

Figure 2. Eosinophilic influx in the BAL fluid following scarification with rVVβgal (filled bars) or rVVG (dashed bars) after intranasal inoculation with mutant RSV. Data shown is representative of four experiments and each bar represents the mean percentage of individual eosinophil counts in each group (n=4). Significant differences (P=0.03) between rVVG and rVVβgal using Mann-Whitney paired comparisons test are indicated by **. The significance levels between rVVG and rVVβgal vaccinated mice are as follows: Long, P=0.03; 63/2/4/1, P=0.03; 63/2/4/8, P=0.03; 63/1/2/3, P=0.86.
which would be disrupted in the 63/1/2/3 mutant. These potential epitopes are located at amino acids 185-193 (ICKRIPKK) and 189–197 (IKPGKK) (bold type indicates anchor and subanchor residues).

In other viral systems, single epitopes have been shown to be responsible for Th2-like CD4+ responses. Proteins from mouse hepatitis virus (JHM; reference 21), hepatitis B virus (22), and retroviruses (23) have been shown to generate Th2 responses and have been mapped using peptides and T cell clones and lines. Unlike our study, in none of these systems has it been possible to examine the effects of altering these epitopes upon the host immune response and induction of pathology. Interestingly, the critical epitope in JHM virus is also I-E<sub>d</sub>-restricted (ILNPKPRQKR) and contains at least four identical or similar residues to the proposed critical motifs of RSV G. It is possible that these homologous epitopes have similar binding affinities for MHC class II molecules, thus skewing towards a Th2-type response (24).

In this and in previous studies, induction of eosinophilia is indicative of a Th2-type response (10, 11, 25). The recruitment of eosinophils into the alveolar space could be due to IL-5 production (26) or the induction of chemokines such as eotaxin (27) or RANTES (regulated on activation, T cell expressed and secreted; reference 28). In our study, this same region is also responsible for the increased weight loss that is observed upon challenge. Weight loss, however, is not a specific indicator of Th2-type, eosinophilic pathology since mice lose weight late in a primary
infection and may show disease enhancement due to sensitization to other RSV proteins (10). Hussell et al. have shown that IL-12 treatment of mice primed with rVV G induces a Th1-type response and diminishes eosinophilia but weight loss is unaffected or in some cases enhanced (29). Our data indicate that region 193–203 of the G protein not only induces eosinophilia (probably via the Th2 pathway), but also induces an immune response (such as TNF-α; reference 30) that is responsible for increased weight loss.

These findings provide a framework which may allow the development of an RSV vaccine without vaccine-enhanced illness. In the mouse model of RSV, vaccine-enhanced illness presents two possible pathways for eosinophil induction: first, a CD4+ response to eosinophilic epitopes within the primary structure of the G protein such as described here, or second, the elimination of the CD8+ response that controls the Th2 response (31). We have recently shown that it is possible to generate eosinophilia to the F protein of RSV when CD8+ cells are deleted. Normally, priming with the F protein of RSV generates a Th1-type response and virus-specific CTLs that are an abundant source of IFN-γ. When CD8+ T cells are deleted, the response is switched to a Th2-type response that generates lung eosinophilia. It appears that in the F response, CD8+ cells are responsible for controlling the Th2-type response, most likely through the production of IFN-γ (13). Similarly, the failure of the formalin-inactivated RSV vaccine trials could have been due to the lack of CD8+ T cell induction since soluble antigen is processed and presented via the MHC class II pathway. Both aspects of the immune response need to be considered for the development of future RSV vaccines.

Our results suggest that it is possible to induce a protective immune response with the G protein without inducing a detrimental eosinophilic response. Since the G protein does not induce a CTL response in BALB/c mice (32), eosinophil induction could in part be due to the lack of a CD8+ response. In spite of this, we have shown that it is possible to induce protective immunity with the G protein if the eosinophilic portion has been altered. Vaccines need to induce a CD4+ response to promote B cell production of protective antibody, but IFN-γ production by CD8+ cells may also be essential to protect against augmentation of illness. However, induction of a strong and unopposed CD4+ response can also be detrimental. Sensitization with the M2 protein alone leads to enhanced illness, and transfer of isolated CD4+ T cells causes augmented disease as well (10, 33). An ideal vaccine should strike a balance between inducing CD4+ and CD8+ T cell responses. Vaccination that stimulates either a CD4+ or CD8+ T cell response alone might lead to pathological consequences (34). Whether these tenets for vaccine development will hold true in an outbred human population remains to be tested.

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References


Figure 5. RSV titers from the lungs of mice vaccinated with rVV s. Mice were scarified with rVV s expressing either βgal, wild-type G, or mutated G proteins (rVV 63/2/4/8 or rVV 63/1/2/3) and challenged intranasally with whole RSV. 4 d after challenge, lungs were homogenized and assayed for RSV. Titers from individual mice are shown from one of two separate experiments which gave similar results.


