Intranasal immunization with W805EC adjuvanted recombinant RSV rF-ptn enhances clearance of respiratory syncytial virus in a mouse model

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Abbreviations: used in this article: BAL, bronchial alveolar lavage; CMI, cell-mediated immune responses; GAPDH, glyceraldehyde-3-phosphate dehydrogenase; IN, intranasal; NE, nanoemulsion; NU, neutralizing units; PAS, Periodic Acid Schiff; rF-ptn, recombinant RSV F protein; RSV, respiratory syncytial virus; SMLN, submandibular lymph node

Respiratory Syncytial Virus (RSV) is a ubiquitous virus that infects almost all people by age two and is a major source of respiratory illness in infants, the elderly and others with compromised immune systems. Currently there is no available vaccine. Prior efforts using formalin-inactivated RSV (FI-RSV) were associated with enhanced respiratory disease upon viral exposure following clinical vaccine trials. Several researchers and pharmaceutical companies have utilized vector-associated live attenuated RSV vaccines in pre-clinical and clinical studies. Another attractive approach, however, is a subunit vaccine which would be easier to produce and quality control. Our group has previously demonstrated in a murine model of infection that intranasal immunization with nanoemulsion-inactivated and adjuvanted RSV induces humoral and cellular immune responses, resulting in protection against RSV infection. The present studies characterize the immune responses elicited by intranasal RSV F protein adjuvanted with nanoemulsion. Intranasal application of nanoemulsion adjuvanted F protein induced a rapid and robust systemic and mucosal antibody response, as well as protection against subsequent RSV challenge. Importantly, RSV challenge in immunized animals did not elicit airway hyper-reactivity, a Th2-skewed immune response or immunopathology associated with hypersensitivity reactions with formalin-inactivated vaccine. These results suggest that RSV F protein adjuvanted with nanoemulsion may be a good mucosal vaccine candidate. Formulating RSV F protein in nanoemulsion creates a well-defined and well-controlled vaccine that can be delivered intranasally to induce T cell mediated immunity without inducing enhanced disease associated with the mouse model of FI-RSV vaccination and infection.

Introduction

RSV infects almost all people by two years of age and is the most common cause of hospitalizations and viral deaths in children under age five.1 It is a worldwide problem causing approximately 34 million cases of lung infection and 199,000 deaths in young children and infants.2 RSV infection can result in bronchitis and is positively correlated with the development of childhood asthma.3,4 Additionally, severe RSV infections occur in the elderly and in people with compromised immune systems and can exacerbate chronic obstructive pulmonary disease (COPD) causing significant mortality.5-7

Despite the enormous medical burden of RSV, to date there is no approved RSV vaccine available. A previous attempt at generating a vaccine using formalin-inactivated virus (FI-RSV) in the 1960s resulted in a vaccine-enhanced disease upon natural infection.8,9 This problem has stilled progress in inactivated RSV vaccine development. An alternative approach is to use live attenuated RSV; however, it has been difficult to achieve the appropriate level of attenuation for infants.10 Previous attempts at attenuated RSV vaccines have resulted in vaccines that do not induce protective immunity or under-attenuated viruses that cause nasal congestion in infants.11,12 As a result, investigators have turned to RSV subunit vaccines adjuvanted with new immunostimulants
Potential vaccine antigens include two of the major RSV capsid glycoproteins, attachment (G) and fusion (F) protein. F protein has been shown to confer Th-1-associated protective immunity against RSV and has been the preferred target for RSV vaccine development.

Vaccination with recombinant F protein and NE induces a rapid and robust humoral immune response

We first determined whether IN immunization with recombinant F protein (rF-ptn) mixed with NE adjuvant is capable of inducing an antibody response. Mice were treated IN with rF-ptn alone or rF-ptn mixed with NE adjuvant (NE + rF-ptn). Treated mice received 2 doses; the first dose of vaccine (day 0) contained 4.45 µg rF-ptn/mouse. The second dose of vaccine (four weeks later) contained 10µg rF-ptn/mouse. Control mice received PBS IN. As shown in Figure 1A, mice treated with rF-ptn alone did not develop detectable levels of anti-F IgG antibodies, and the response level was similar to mice treated with PBS alone. In contrast, animals immunized with NE + rF-ptn produced anti-F IgG endpoint titer of 1 × 10^4 after only one immunization and approximately 1 × 10^5 at week 6, two weeks after the second immunization.

We further characterized the immune response elicited by NE + rF-ptn by comparing it to that induced by immunization with viral RSV particles inactivated in NE adjuvant (NE + RSV). For these studies, mice were immunized either with rF-ptn (2.5 µg/animal) or RSV (1.3 × 10^5 plaque forming units (pfu)/animal) and NE adjuvant at day 0 and four weeks later. Control mice received PBS IN. As shown in Figure 1B, both groups immunized with NE combined with either rF-ptn or RSV demonstrated the production of anti-F IgG antibodies; however, anti-F IgG was significantly higher in mice immunized with NE + rF-ptn as compared with NE + RSV at 6 wk (P ≤ 0.001). To confirm that the resulting antibodies recognized epitopes present on RSV and not just the recombinant protein, we evaluated antibody response against rF-ptn and purified, inactivated RSV at the week 6 timepoint (Fig. 1C). There was a significant increase in endpoint titers in the serum of NE + rF-ptn vaccinated mice (10^5) as compared with those vaccinated with NE + RSV (10^3) (P < 0.001).

It has been suggested that mucosal IgA is critical in protection of the host against respiratory pathogens such as influenza A. We therefore determined IgA endpoint titers in the BAL of mice immunized with either NE + rF-ptn or NE + RSV. As shown in Figure 1D, immunization with either NE + rF-ptn or NE + RSV significantly increased the anti-F IgA endpoint titer as compared with the sham-immunized mice (P = 0.008 and 0.032, respectively), suggesting that both vaccines induced secretion of F protein specific antibodies in the mucosa.

High levels of serum neutralizing antibodies have been associated with decreased risk of severe RSV infection.
Neutralization activity was assessed in mice vaccinated with NE + rF-ptn or NE + RSV. Serum samples from week 8 terminal bleeds were pooled for each group and run in a neutralization assay. NE + rF-ptn had a pooled Neutralization Unit (NU) of 593 whereas NE + RSV had a pooled NU of 25 (Fig. 2).

Cell-mediated immune response of mice immunized with NE + rF-ptn or NE + RSV

To further characterize the quality of the immune response generated by NE + rF-ptn and the polarity of the T cell response to this antigen, we evaluated cytokine recall response from the submandibular lymph node-associated lymphocytes of immunized animals that were not challenged with live virus. Various cytokines were assessed (IFN-γ, IL-2, IL-4, IL-5, IL-10, IL-17), but there were no significant differences in these cytokines between the vaccinated or sham control animals (Fig. 3). Only the Th1 cytokine IL-2 was significantly increased in mice vaccinated with NE + RSV compared with NE + rF-ptn vaccinated mice. There was no change in the Th1/Th2 skewing of the cell-mediated cytokine response in any groups compared with sham vaccinated animals.

Immunization with NE + rF-ptn or NE + RSV leads to enhanced viral clearance

Mice immunized intranasally with NE + rF-ptn or NE + RSV were challenged with 1 × 10⁵ pfu RSV two weeks after the second immunization. Mice were euthanized 8 d post-challenge to assess the impact of the vaccine on viral clearance and to characterize the immune response. Immunization with either NE + rF-ptn or NE + RSV resulted in immune characterized by enhanced viral clearance (Fig. 4A), as demonstrated by a significant decrease in lung mRNA expression of RSV proteins F and G when compared with sham vaccinated animals ($P < 0.01$). In the NE + rF-ptn group, there was an overall 8- and 10-fold decrease in F and G mRNA, respectively, with 60% of the animals having no detectable viral mRNA. All of the animals in the NE + RSV group had undetectable viral mRNA. Intranasal inoculation of mice with L19 RSV leads to an infection that is associated with mucus hypersecretion and inflammation. At day 8, post-challenge, NE + rF-ptn vaccinated mice exhibited similar mucus secretion and goblet cell infiltration compared with challenged sham-immunized mice, as assessed via histologic analysis ($P > 0.05$) (Fig. 4B and C). Cytokine mRNA levels in the lungs were also determined (Fig. 4D). There was no significant increase in Th2 cytokine mRNA (IL-4, -5, -13) in lungs of animals vaccinated with NE + rF-ptn or NE + RSV. There was, however, a significant increase in IL-17 in NE + RSV and NE + rF vaccinated groups compared with sham vaccinated mice ($P < 0.05$, $P < 0.01$, respectively). In addition, mice challenged after vaccination with NE + rF-ptn showed no increase in airway hyper-reactivity as compared with challenged sham vaccinated animals (Fig. 4E).

Discussion

There have been many RSV vaccine candidates since the failure of the FI-RSV vaccine in the 1960s. Among these are killed virus, attenuated virus, virus-like particles, recombinant viral vaccines, and subunit vaccines.²⁶ While live-attenuated viral vaccines are better at inducing immune memory, it is difficult to achieve the appropriate level of attenuation in infants,¹⁰ so there is continued interest in developing subunit vaccines for RSV. F and G proteins have each been studied as subunit vaccines, with F-protein being the preferred candidate because its structure is conserved across RSV strains, and anti-F antibodies are sufficient to induce protection.¹⁴ A vaccine consisting of RSV F-protein vs. whole RSV also has the advantage of containing only one component and therefore may be more consistent from lot to lot. Being only one component, the manufacturing of a NE-formulated F-protein vaccine can be well defined, well controlled and reproducible.

Pre-clinical and clinical evaluations of RSV F protein subunit vaccines have demonstrated safety and suggest efficacy against viral challenge.¹³ However, most reports using recombinant F protein have administered the experimental vaccines via parenteral delivery routes. Given the mucosal transmission of the virus, a mucosal vaccine may be preferred. Purified F and G protein vaccines have been administered to rodents with a variety of adjuvants, including alum, monophosphoryl lipid A, Freund’s adjuvant, and cholera toxin B. A preclinical study comparing an F and G protein subunit vaccine found that while both intranasal and intramuscular vaccination induced antibody responses, only the intranasal vaccine protected against both upper and lower respiratory tract RSV infections.²⁷

Of the experimental purified F protein vaccines that were tested in clinical trials, most were well-tolerated and induced neutralizing antibodies but did not induce sufficient protection against RSV infection in the lower respiratory tract, suggesting the importance of innate immune activation for protection against viral infection.

The development of a mucosally delivered RSV subunit vaccine is potentially attractive in regards to safety and efficacy given the nature of the mucosal transmission of the virus and the potential of improved protection from mucosal immunity. In this regard, nanoemulsion adjuvant has been previously demonstrated to be an effective mucosal adjuvant for intranasal delivery of inactivated viruses and recombinant proteins including RSV.²²⁻²⁴ In this experiment, the nanoemulsion adjuvant generated F-protein specific IgA secreted in the BAL. The intranasal
route of administration of this NE-rF-ptn vaccine may also be advantageous, as it should induce the mucosal response appropriate for a mucosal infection.

The mechanism of NE-induced immunity following intranasal administration has previously been studied.29 Antigen formulation with NE leads to antigen uptake in nasal epithelial cells. Antigen is then taken up in the local lymphoid tissue or trafficked to draining lymph nodes. NE is associated with immunogenic apoptosis of epithelial cells and increased MHC class II gene expression. Intranasal delivery of NE also induces heterogeneous cytokine production (Th1, Th2, and Th17) in lymphoid cells in various tissues including the nasal septum, spleen, and bone marrow. IL-17 has been indicated for its role in autoimmunity and asthma.31,32 However, it has also been discovered to play a key role in the clearance of many respiratory pathogens, including *Mucocutaneous candidiasis* and *Mycobacterium tuberculosis*.33,34 IL-17 has been shown to increase pathogenesis of RSV infections in mice, leading to increased IL-13 and mucus secretion in lungs.35 However, in this study, we show increased lung IL-17 without an accompanying increase of IL-13 or mucus. Our lab has previously shown NE-induced IL-17 and its association with Th1 cytokine IFNγ and not IL-4.36 In this study, immunization with NE + rF-ptn led to CMI-associated responses (IFN-γ and IL-17) without skewing toward a Th2 response.

We have previously demonstrated that NE-based vaccines induce both innate and adaptive immune responses, and it is clear that NE is a very potent activator of innate immunity.22,28,30,37-40 It has been suggested that vaccine-enhanced disease induced by FI-RSV may be caused by low antibody affinity due to poor Toll-like receptor stimulation.41 We have evidence that NE activates TLR-2 and -4 pathways which would enhance the affinity of the induced antibodies, making this less of a problem for NE than other adjuvants.36 When combined with NE, the F protein subunit vaccine induced a rapid and robust neutralizing antibody response. As previously observed with a NE + RSV vaccine, intranasal immunization with NE + rF-ptn vaccine also resulted in enhanced viral clearance after RSV challenge, and this activity was not associated with an increase in airway hyper-reactivity.22 Although we did not directly compare with animals immunized with FI-RSV in these studies, mice vaccinated with FI-RSV have demonstrated at least a 10-fold increase in AHR compared with control mice; a finding indicative of a hyper-immune response resulting from immunization.42 One shortcoming in this current study was the lack of comparison to immunity induced by live non-inactivated RSV. This comparison is planned for future studies and will be essential to evaluate the potential utility of the approach.

While there were differences between antibody responses in mice vaccinated with NE + rF-ptn and NE + RSV, both treatments induced protection against viral challenge. The amount of RSV in the vaccine was based on pfu, not F protein content, and T cell epitopes on other components of the virus may have contributed to protection. Alternatively, both vaccines may have induced antibody levels above a threshold needed for protection.
NE is a technology that potentially allows for multiple approaches for prevention of RSV infection. NE can be formulated with either whole split virus or recombinant protein. Each of these vaccine formulations enhances immune responses and leads to protection in a mouse model without evidence of vaccine-enhanced disease. An intranasal NE-based vaccine is additionally interesting because it offers needle-free administration and lacks the need for cold-chain storage, as it is stable at room-temperature. Taken together, this highlights the potential utility of a NE-based RSV mucosal vaccine in infants, children, and adults.

**Materials and Methods**

RSV, recombinant F protein, adjuvant, and vaccine

RSV Line 19 (L19) strain and W805EC nanoemulsion (NE) were provided by NanoBio Corp. W805EC is an oil-in-water emulsion (~500nm droplet size) made from ingredients that are Generally Recognized As Safe (GRAS) or are approved pharmaceutical excipients by the Unites States Food and Drug Administration. W805EC NE is composed of Tween 80 (5%), ethanol (8%), cetyl pyridinium chloride (CPC, 1%), soybean oil (64%) and water and is emulsified into droplets with a mean diameter of 500 nm using a high speed emulsifier. Recombinant F protein (derived from human respiratory syncytial virus A [Strain A2]) produced in Baculovirus was purchased from Sino Biologicals, Inc. (Cat # 11049-V08B). 60% W805EC nanoemulsion adjuvant was provided by NanoBio Corp. Different concentrations of recombinant F protein (rF-pton) were mixed with NE at a final concentration of 20% (v/v). RSV L19 virus was co-incubation with NE at a final concentration of 20% NE (v/v) for 3 h at room temperature resulting in an inactivated, split virus RSV/NE. After incubation, RSV/NE was used as an intranasal vaccine. RSV vaccine contained 1.3 x 10^5 pfu/mouse. F protein vaccine contained 2.5 – 10µg rF-pton/mouse.

Mice intranasal (IN) immunization and challenge

Specific pathogen-free BALB/c mice (females 6–8 wk old) were purchased from the Jackson Laboratory (Bar Harbor, ME).
Whole blood samples were centrifuged at 7000x g for 9 min to separate the serum. Bronchoalveolar lavage (BAL) was obtained from freshly euthanized mice and single cell suspensions were prepared by disruption and filtration through a 100 μm nylon mesh screen, followed by NH₄Cl lysis of erythrocytes. Cells were washed twice with PBS and resuspended in RPMI containing 50 μM β-mercaptoethanol, 5% FBS, and 1% of Penicillin/Streptomycin, L-glutamine, non-essential amino acids, sodium pyruvate, and MOPS and plated in duplicate in a 96 well plate at 80000 cells per well in 200 μL media. Cells were stimulated with L19 RSV at an MOI of 0.5 or were untreated (control). Cells were incubated at 37 °C for 48 h. Supernatants were harvested and frozen at –20 °C before being analyzed by mouse Multiplex kits and LumineX software (Millipore, Cat. # MPXMCYTO-70k-08) as described previously. Data are expressed as [RSV] – [Control] = Total (pg/ml) for each cytokine (mean of duplicate determinations).

Neutralization assay

Serum samples were pooled by group in order to have sufficient quantities for the neutralization assay. Serum sample dilutions were prepared in EMEM containing 2% FBS. Eight hundred pfu of RSV were added to one milliliter of serum test dilutions. The serum/virus mixture was incubated at 37 °C for one hour. As a positive control, virus was mixed with EMEM containing 2% FBS and incubated along with test samples. The serum/virus mixture was applied on Vero cells and plates were incubated for 3 h at 37 °C. After incubation the antibody/virus mixture was removed and replaced with 0.9% methylcellulose pre-warmed at 37 °C. The serum/virus mixture was incubated at 37 °C for 48 h. Supernatants were harvested and frozen at –20 °C before being analyzed by mouse Multiplex kits and LumineX software (Millipore, Cat. # MPXMCYTO-70k-08) as described previously. Data are expressed as [RSV] – [Control] = Total (pg/ml) for each cytokine (mean of duplicate determinations).

Real-time Taqman PCR

Eight days after RSV challenge, mice were euthanized and a lung lobe was removed and homogenized in 1 ml of Triozol reagent (Invitrogen, Cat. #15596-026). RNA was isolated, and 5 μg was reverse-transcribed to assess gene expression. Detection of cytokine mRNA in lung samples was determined using pre-developed primer/probe sets, as described previously. Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was used as an internal control, and gene expression was normalized to GAPDH. Fold changes in gene expression levels were calculated by comparison of the gene expression in uninfected mice, which were assigned an arbitrary value of 1.

Histology

Lung lobes of infected mice were perfused with 10% formalin, removed, and placed in fresh formalin for an additional 24–48 h. Formalin-fixed lung tissues were embedded in paraffin and then cut into 5μm sections prior to staining with Periodic Acid Schiff (PAS) to detect mucus production. Mucus was graded in a blinded fashion on a scale of 0–4 as follows: 0—no goblet cell staining; 1—some goblet cell staining with PAS in multiple airways; 2—extensive goblet cell staining in multiple airways; 3—extensive goblet cell staining with mucus in the airways; and 4—extensive cell staining with mucus plugging in multiple airways.

Airway response

Airway hyper-reactivity was determined using whole body plethysmography with ventilation of anesthetized mice. Briefly, mice were anesthetized with sodium pentobarbital and intubated via cannulation of the trachea with an 18-gauge metal tube. Mice were ventilated with a Harvard pump ventilator (tidal volume, 0.4 mL; frequency, 120 breaths/minute; positive end-expiratory pressure, 2.5 to 3.0 cm H₂O; Harvard Apparatus) and challenged intravenously with 0.2 mg of methacholine. Resistance was calculated by dividing the change in pressure (Ptp) by the change in flow (F) (Ptp/F; units = cm H₂O/mL/second). The peak airway resistance was recorded as a measure of airway hyper-reactivity.

Statistical analysis

Statistical comparisons were assessed by Mann-Whitney test using GraphPad Prism version 5.00 (GraphPad Software). A P value < 0.05 was considered significant.

Disclosure of Potential Conflicts of Interest

The authors of the manuscript do not have any financial conflicts of interest that might be construed to influence the results.
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References

1. Hall CR, Weinberg GA, Iwane MK, Blamkin AK, Edwards KM, Staat MA, Aunger P, Griffin MR, Poelfling KA, Ermann D, et al. The burden of respiratory syncytial virus infection in young children. N Engl J Med 2009; 360:588-98; PMID:19196675; http://dx.doi.org/10.1056/NEJMoA0804877

2. Nair H, Nokes DJ, Gesner BD, Dherani M, Madhi SA, Singleton RJ, O’Brien KL, Roca A, Wright PF, Bruce N, et al. Global burden of acute lower respiratory infections due to respiratory syncytial virus in young children: a systematic review and meta-analysis. Lancet 2010; 375:1545-55; PMID:20399433; http://dx.doi.org/10.1016/S0140-6736(10)60286-1

3. Sabin AB, Murphy BR, Muller M, Piddick JA, Johnston SL. Respiratory syncytial virus persistence in chronic obstructive pulmonary disease. Pediatr Infect Dis J 2008; 27(Suppl):S63-70; PMID:18820581; http://dx.doi.org/10.1097/INF.0b013e3181646d47

4. Wu P, Dupont WD, Griffin MR, Carroll KN, Mitchell EF, Gebretsadik T, Herttuv TEV. Evidence of a causal role of winter virus infection during infancy in early childhood asthma. Am J Respir Crit Care Med 2008; 178:1123-9; PMID:18776531; http://dx.doi.org/10.1164/rcrm.20080804.579OC

5. Foley AR. Respiratory syncytial virus infection in adults. Semin Respir Crit Care Med 2007; 28:171-81; PMID:17458771; http://dx.doi.org/10.1055/s-2007-976489

6. De Serres G, Lampton N, La Forge J, Rouleau I, Bourbeau J, Weiss K, Barret B, Boivin G. Importance of viral and bacterial infections in chronic obstructive pulmonary disease exacerbations. J Clin Virol 2009; 46:129-33; PMID:19665843; http://dx.doi.org/10.1016/j.jcv.2009.07.006

7. Thompson WJ, Shuk WP, Weintraub E, Bramer L, Cox N, Anderson LJ, Fukuda K. Morbidity associated with influenza and respiratory syncytial virus in the United States. JAMA 2003; 289:179-86; PMID:12517228; http://dx.doi.org/10.1001/jama.289.2.179

8. Chanock RM, Parrott RH. Acute Respiratory Disease in Infancy and Childhood: Present Understanding and Prospects for Prevention. Pediatrics 1965; 36:21-39; PMID:14313563

9. Fulgentis VA, Eller JF, Sieber OF, Joyner JW, Minamimizu M, Meiklejohn G. Respiratory virus immunization. I. A field trial of two inactivated respiratory virus vaccines; an aqueous trivalent parainfluenza virus and an alum-precipitated influenza virus FG subunit vaccine. J Infect Dis 1994; 170:425-8; PMID:8035830; http://dx.doi.org/10.1086/313919

10. Schickli JH, Dubovsky F, Tang RS. Challenges in developing a pediatric RSV vaccine. Hum Vaccin 2009; 5:582-91; PMID:19556888

11. Karron RA, Wright PF, Crowe JE Jr., Clements-Moore JL, Thompson J, Makhlouf M, Casey B, Murphy BR. Evaluation of two live, cold-passaged, temperature-sensitive respiratory syncytial virus vaccines in chimpanzees and in human adults, infants, and children. J Infect Dis 1997; 176:1428-36; PMID:9395551; http://dx.doi.org/10.1086/314138

12. Wright PF, Karron RA, Belseck RB, Thompson J, Crowe JE Jr., Boyley TG, Haliburton LL, Reed GW, Whitehead SS, Anderson EL, et al. Evaluation of a live, cold-passaged, temperature-sensitive, respiratory syncytial virus vaccine candidate in infancy. J Infect Dis 2000; 182:1351-42; PMID:11008386; http://dx.doi.org/10.1086/313859

13. Cherukuri A, Stokes KL, Patton K, Hsu K, Sakamoto K, Lambert S, Stillman E, Moore ML, Lee S. An adjuvanted respiratory syncytial virus fusion protein induces protection in aged BALB/c mice. Immun Ageing 2012; 9:21; PMID:23036100; http://dx.doi.org/10.1186/1743-4933-9-21

14. Kneyber MC, Kimpen JL. Advances in respiratory syncytial virus vaccine development. Curr Opin Investig Drugs 2004; 5:163-70; PMID:15434990

15. Singh SR, Dennis VA, Carter GL, Pillai SR, Moore FG. Respiratory syncytial virus recombinant F protein (residues 255-278) induces a helper T cell type 1 immune response in mice. Viral Immunol 2007; 20:261-75; PMID:17663843; http://dx.doi.org/10.1089/vim.2007.0088

16. Felts TF, Sondheimer HM, Tulloch RM, Harris BS, Jensen KM, Losonsky GA, Griffin MP, Matzevitz Cardio Study Group. A randomized controlled trial of motavizumab versus palivizumab for the prophylaxis of serious respiratory virus disease in children with hemodynamically significant congenital heart disease. Pediatr Res 2011; 70:186-91; PMID:21522037; http://dx.doi.org/10.1203/PDR.0b013e3182665253

17. Robinson KACO, Odedola OA, Saldanha DJ, McKoy NA, Palivizumab for prophylaxis against respiratory syncytial virus infection in children with cystic fibrosis. Cochrane Database Syst Rev 2012; 2:CD007743; PMID:22358632

18. Neusil KM, Johnson JE, TangYW, Pietsch JP, Slani M, Gar N, Hoagland P. Adjuvant influenza vaccine: clinical trial of a novel adjuvanted respiratory syncytial virus vaccine. J Fam Pract 2012; 54:905-12; PMID:22238168; http://dx.doi.org/10.1093/jfip/ctx055

19. Hurwitz JL. Respiratory syncytial virus vaccine development. Expert Rev Vaccines 2011; 10:1415-33; PMID:21988307; http://dx.doi.org/10.1586/erv.11.120

20. Oen NL, Bradeau RJ, Welch EE, Withrow MW. Induction of local and systemic immunity against human respiratory syncytial virus using a chimeric FG glycoprotein and cholera toxin B subunit. Vaccine 1994; 12:731-3; PMID:8091852; http://dx.doi.org/10.1016/0264-410X(94)90224-0

21. Makidon PE, Bielinska AU, Ngavekar SS, Janczak KW, Knowlton J, Scott AJ, Mank N, Cao Z, Rathinavelu S, Beer MR, et al. Pre-clinical evaluation of a novel nanoemulsion-based hepatitis B mucosal vaccine. PLoS One 2008; 3:e2994; PMID:18698426; http://dx.doi.org/10.1371/journal.pone.0002994

22. Makidon PE, Belyakov IM, Blanco LP, Janczak KW, Landers J, Bielinska AU, Groom JV 2*, Baker JR Jr. Nanoemulsion mucosal adjuvant uniquely activates cytokine production by nasal ciliated epithelium and induces dendritic cell trafficking. Eur J Immunol 2012; 42:2073-86; PMID:22653620; http://dx.doi.org/10.1002/eji.201142346

23. Myc A, Kukowska-Latalla JF, Bielinska AU, Cao P, Myc PP, Janczak K, Sturm TR, Grabinski MS, Landers JY, Young KS, et al. Development of immune responses that protect against ethanol-induced liver fibrosis after a single intranasal immunization with influenza A virus and nanoemulsion. Vaccine 2003; 21:3801-14; PMID:12922114; http://dx.doi.org/10.1016/S0264-410X(03)00381-5

24. Bertelli E, Oukka M, Kuchnio VK. T(H)17 cell scientists in the circle of immunity and autoimmunity. Nat Immune 2007; 8:345-50; PMID:17375096; http://dx.doi.org/10.1038/nri0407-345

25. Wang YH, You KS, Liu B, Chen CY, Uygur-Bimpool B, Spodolski W, Brunner JA, Huston DP, Liu YJ. A novel subset of CD4(+) T(H)2 memory/effector cells that produce inflammatory IL-17 cytokine and promote the exacerbation of chronic allergic asthma. J Exp Med 2010; 210:2479-91; PMID:20921287; http://dx.doi.org/10.1084/jem.20101576

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33. Dominguez-Villar M, Hafler DA. Immunology. An innate role for IL-17. Science 2011; 332:47–8; PMID:21454778; http://dx.doi.org/10.1126/science.1205311

34. Khader SA, Bell GK, Pearl JE, Fountain JJ, Rangel-Moreno J, Cilley GE, Shen F, Eaton SM, Gaffin SL, Swain SL, et al. IL-23 and IL-17 in the establishment of protective pulmonary CD4+ T cell responses after vaccination and during Mycobacterium tuberculosis challenge. Nat Immunol 2007; 8:369-77; PMID:17351619; http://dx.doi.org/10.1038/ni1449

35. Mukherjee S, Lindell DM, Berlin AA, Morris SB, Shanley TP, Hershenson MB, Lukacs NW. IL-17-induced pulmonary pathogenesis during respiratory viral infection and exacerbation of allergic disease. Am J Pathol 2011; 179:248-58; PMID:21703407; http://dx.doi.org/10.1016/j.ajpath.2011.03.003

36. Bielinska AU, Gerber M, Blanco LP, Makidon PE, Janczak KW, Beer M, Swanson B, Baker JR Jr. Induction of Th17 cellular immunity with a novel nanoemulsion adjuvant. Crit Rev Immunol 2010; 30:189-99; PMID:20370629; http://dx.doi.org/10.1615/CritRevImmunol.v30.i2.60

37. Das SC, Hatta M, Wilker PR, Myc A, Hamouda T, Neumann G, Baker JR Jr., Kawaoka Y. Nanoemulsion W805EC improves immune responses upon intranasal delivery of an inactivated pandemic H1N1 influenza vaccine. Vaccine 2012; 30:6871-7; PMID:22898689; http://dx.doi.org/10.1016/j.vaccine.2012.09.007

38. Stanberry LR, Simon JK, Johnson C, Robinson PL, Morry J, Flack MR, Gracon S, Myc A, Hamouda T, Baker JR Jr. Safety and immunogenicity of a novel nanoemulsion mucosal adjuvant W805EC combined with approved seasonal influenza antigens. Vaccine 2012; 30:107-16; PMID:22079079; http://dx.doi.org/10.1016/j.vaccine.2011.10.094

39. Hamouda T, Sutcliffe JA, Castronova, Baker JR Jr. Intranasal immunization of ferrets with commercial trivalent influenza vaccines formulated in a nanoemulsion-based adjuvant. Clin Vaccine Immunol 2011; 18:1167-75; PMID:21543588; http://dx.doi.org/10.1128/CVI.00035-11

40. Bielinska AU, Gerber M, Blanco LP, Makidon PE, Janczak KW, Beer M, Swanson B, Baker JR Jr. Induction of Th17 cellular immunity with a novel nanoemulsion adjuvant. Crit Rev Immunol 2010; 30:189-99; PMID:20370629; http://dx.doi.org/10.1615/CritRevImmunol.v30.i2.60

41. Delgado MF, Coviello S, Monsalvo AC, Melendi GA, Hernandez JZ, Batalle JP, Diaz L, Trento A, Chang HY, Mitra S, et al. Lack of antibody affinity maturation due to poor Toll-like receptor stimulation leads to enhanced respiratory syncytial virus disease. Nat Med 2009; 15:34-41; PMID:19079256; http://dx.doi.org/10.1038/nm.1894

42. Polack FP, Teng MN, Collins PL, Prince GA, Exner M, Regele H, Lirman DD, Rabold R, Hoffman SJ, Karp CL, et al. A role for immune complexes in enhanced respiratory syncytial virus disease. J Exp Med 2002; 196:859-65; PMID:12235218; http://dx.doi.org/10.1084/jem.20020781

43. Hamouda T, Chepurnov A, Mank N, Knowlton J, Chepurnova T, Myc A, Sutcliffe J, Baker JR. Efficacy, immunogenicity and stability of a novel intranasal nanoemulsion-adjuvanted influenza vaccine in a murine model. Hum Vaccin 2010; 6:585-94; PMID:20421727; http://dx.doi.org/10.4161/hv.6.7.11818

44. Tekkanat KK, Maassab H, Berlin AA, Lincoln PM, Evansoff HL, Kaplan MH, Lukacs NW. Role of interleukin-12 and stat-4 in the regulation of airway inflammation and hyperreactivity in respiratory syncytial virus infection. Am J Pathol 2001; 159:631-8; PMID:11485921; http://dx.doi.org/10.1016/S0002-9440(10)61734-8