

Article

Flexible RSV prefusogenic fusion glycoprotein exposes multiple neutralizing epitopes that may collectively contribute to protective immunity

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Abstract: Human respiratory syncytial virus (RSV) is a significant cause of lower respiratory tract infection in infants, young children, and older adults. There is no licensed vaccine and prophylactic treatment options are limited and not widely available in developing countries with the greatest disease burden. The RSV fusion (F) glycoprotein is a primary target of host immunity and thus a major focus for vaccine development. The native F glycoprotein structure is flexible and undergoes significant rearrangements from the metastable prefusion to a stable postfusion structure with neutralizing epitopes on intermediate structures. We hypothesize trimeric vaccine strategies that recapitulate the breathable F quaternary structure, and provide accessibility of B-cells to epitopes on intermediate conformations, may collectively contribute to protective immunity, while ridge prefusion F structures restrict access to key protective epitopes. To test this hypothesis, we used the near full-length native prefusogenic F as a backbone to construct three prefusion F variants with substitutions in the hydrophobic head cavity: 1) disulfide bond double mutant (DS), 2) space filling hydrophobic amino acid substitutions (Cav1), and 3) DS plus Cav1 substitutions (DS-Cav1). In this study, we compared the immunogenicity of prefusogenic F to the immunogenicity of the prefusion F variants in two animal models. Native prefusogenic F was significantly more immunogenic producing high titer antibodies to prefusogenic, prefusion, and postfusion F structures compared to animals immunized with prefusion F DS or DS-Cav1. Importantly, native prefusogenic F elicited antibodies that targeted neutralizing epitopes including prefusion-specific site zero (\emptyset) and V as well as conformation-independent neutralizing sites II and IV. Immunization with prefusion F DS or DS-Cav1 elicited antibodies primarily targeting antigenic sites \emptyset and V with little or no detectable antibodies to other key neutralizing sites. Animals immunized with native prefusogenic F also had significantly higher neutralizing antibodies that cross-neutralized RSV A and B subtypes while immunization with DS or DS-Cav1 elicited neutralizing antibodies primarily to the A subtype. We conclude that breathable trimeric vaccines that closely mimic the native F-structure, and incorporate strategies for B-cell accessibility to protective epitopes, are important considerations for vaccine design. F structures locked in a single conformation restrict B-cell access to neutralizing epitopes that may collectively contribute to destabilizing F-trimers important for broad protection. These results also have implications for vaccine strategies targeting other type 1 integral membrane proteins.

Keywords: respiratory syncytial virus; fusion glycoprotein; prefusogenic RSV F; cotton rat

1. Introduction

Human respiratory syncytial virus (RSV) is a cause of lower respiratory infection (LRTI) in young children and older adult populations. The disease burden is particularly high in developing countries with over 3 million hospitalizations and 50,000-70,000 deaths in young children. Children under 5 years of age are the most susceptible and account for 45% of RSV-related deaths with the vast majority (>90%) in developing countries [1]. RSV disease burden is also substantial in older adults with over 1 million infections, 300,000 hospitalizations, and over 10,000 in-hospital deaths [2]. Although RSV is the cause of significant worldwide disease burden, there are no licensed vaccines and palivizumab (Synagis®) is the only licensed prophylaxis for prevention of RSV in high-risk newborn. There is no established correlate of protective immunity [3-5].

RSV is a negative-strained, enveloped RNA virus in the *Pneumoviridae* family. The fusion (F) glycoprotein is a major component of the virus envelope. RSV F protein is conserved between human RSV A and B subtypes (>90% amino acid identity) with shared neutralizing epitopes, is essential for infection, and a major target of host immune defense [6-10]. RSV F is a type 1 integral membrane protein produced as a 70 kDa inactive precursor (F0). Unique to RSV, the F glycoprotein has two cleavage sites at positions R109 (site I) and R136 (site II) that are processed by host cell furin-like proteases. Removal of the intervening 27 amino acid peptide (p27) generates a small F2 subunit and a larger F1 subunit [11-14]. The F1 subunit contains the fusion peptide (FP) on the N-terminus, two heptad repeats (HRA and HRB), and the transmembrane (TM) domain and cytoplasmic tail (CT) on the C-terminus. The F2 subunit contributes to fusogenicity, and contains a single heptad repeat (HRC) [7,15-18]. F2/F1 are covalently linked by 2-disulfide bonds to form a protomer. Three F2/F1 protomers associate through weak non-covalent interprotomeric bonds to form the functional F-trimer. Trimers are highly flexible and transiently open and close, dissociate, and monomerize within the lipid bilayer exposing a range of conformations recognized by B-cell receptors [19,20].

Prefusion F undergoes significant rearrangements to a stable postfusion F during attachment and fusion. Events triggering F-protein rearrangement are not understood, although elevated temperatures promote spontaneous rearrangement of the heptad repeats (HRA, HRB and HRC) to form the fusogenic six-helix bundle (6HB) that releases the FP from the hydrophobic cavity. FP insertion into the host membrane is essential for alignment of the virus and host cell membranes, fusion, pore formation, and release of the virus genome into the host cell [21]. RSV F variants with amino acid substitutions in the hydrophobic cavity stabilize F in the prefusion conformation. Disulfide-bond double mutant DS (S155C and S290C), Cav1 with cavity filling hydrophobic amino acids at positions S150F and V207L, and DS-Cav1 with four substitutions have been extensively characterized and shown to bind neutralizing monoclonal antibodies (mAbs) [22,23]. It has been proposed that prefusion F is a superior vaccine candidate since antibodies bind antigenic sites zero (Ø) and V with high affinity, although there is no clinical evidence supporting protective efficacy [24].

Monoclonal antibodies (mAbs) from convalescent patients have provided insight into the plasticity of the humoral immune response to subtle conformational changes in the F structure. For instance, human neutralizing mAbs D25, AM22, and RSD5 all bind prefusion-specific antigenic site Ø with high affinity [22,25]. Crystal structures of prefusion DS (PDB: 4MMQ) and DS-Cav1 (PDB: 4MMS), in complex with D25, AM22, and RSD5 Fabs (fragment antigen binding), show these Fabs bind site Ø at 3 distinct angles [26,27]. Gillman et al. [19] have recently compared binding of human mAbs CR9501 and hRSV90 to antigenic site V. Crystal structures of CR9501 Fab in complex with antigenic site V shows the Fab is rotated by 60° compared to hRSV90 and CDR makes contact with F2. Significantly, CR9501 binding

destabilizes F-trimers and promotes disassembly [19]. Similarly, mAbs palivizumab and motavizumab bind sites II at distinct angles and with different affinity [9,28,30]. We recently reported that murine mAb R4.C6 binds a quaternary epitope consisting of antigenic site II on one protomer and site IV on a neighboring protomer [28]. Collectively, these observations demonstrate that B-cells recognize a wide range of intermediate trimeric and monomeric F structures. These observations and others suggest trimeric vaccines candidates that mimic the breathable native F structure, allowing B-cell accessibility to sequestered epitopes, not only block receptor binding but, more importantly, may destabilize the trimeric structure.

We have described the development of a RSV prefusogenic F constructed from the near full-length RSV/A2 F protein in which furin cleavage site II was mutated to retain the native p27 with a truncated FP (Δ FP) on the N-terminus of F1. F1/F2 protomers assemble as trimers and 3-6 F-trimers spontaneously associate to form 35-40 nm nanoparticles. Prefusogenic F nanoparticles are immunogenic and protective in the cotton rats [29,31], elicit maternal antibodies that protect newborn infant baboons against pulmonary RSV challenge [32], and is safe and immunogenic in women of childbearing age (NCT02624947) [33], and older adults [34]. Here, we used prefusogenic F as a backbone to assess the effects of introducing prefusion stabilizing mutations (DS, Cav1, DS-Cav1) upon the immunogenicity of prefusogenic F. Overall we find that prefusogenic F was significantly more immunogenic than ridged DS or DS-Cav1 variants. Importantly, immunization with prefusogenic F elicited a broad range of antibodies that bound prefusion-specific antigenic sites \emptyset and V as well as serotype conserved neutralizing sites II and IV. Prefusion DS and DS-Cav1 variants were less immunogenic and elicited antibodies primarily to sites \emptyset and V with little or no antibodies to other neutralizing F epitopes. These findings suggest breathable F vaccine candidates produce broad range of antibodies to neutralizing sites that may collectively contribute to protection against RSV. These results have implications for vaccine strategies targeting other type 1 integral membrane proteins.

2. Materials and Methods

2.1. Cell Lines, Viruses, Synthetic peptides, and Monoclonal Antibodies (mAbs)

HEp-2 (ATCC, CCL-23, Manassas, VA, USA) cells were maintained in MEM with Earle's salts and L-glutamine (Gibco Laboratories, Gaithersburg, MD, USA), 10% fetal bovine serum (FBS; HyClone, Logan, UT, USA), and antibiotics (Life Technologies, Grand Island, NY, USA). RSV/A Long challenge stock (lot KJ060319, 2.21×10^7 pfu mL^{-1}) was propagated in HEp-2 cells and produced by Sigmovir Inc. (Gaithersburg, MD, USA). Synthetic p27 peptide (RARRELPRFMNYTLNPNKKTNTVLSKPKRRF) was produced by Thermo Fischer, Inc. Palivizumab (site II) was obtained from MedImmune, Inc. (Rockville, MD, USA). hRSV90 (site VIII also referred to as site V) [35] was provided Dr. James Crowe (Vanderbilt Vaccine Center, Vanderbilt University, Nashville, TN). D25 (site \emptyset) [8,22,23] was purchased from Creative BioLabs (Shirley, NY, USA). Mouse mAbs R1.42 (site IV) and R7.10 (p27) were produced by Novavax, Inc. [28,31]. World Health Organization (WHO) human RSV Reference Standard was purchased from the National Institute of Biological Standards and Control (NIBSC code 16/284, 2000 IU mL^{-1} , Potters Bar, Hertfordshire, UK).

Other reagents. RSV F DS-Cav1-293 reagent was produced by transient transfection of HEK293 T cells with pcDNA 3580 encoding the prefusogenic F gene with the DS (S55C and S290C) and the Cav1 (S190F and V207L) mutations. HEK293 T cells were transfected with lipofectamine 3000 following the manufacturer's procedure (Invitrogen, Carlsbad, CA, USA). Prefusion F (BV2129) reagent was

constructed from the prefusogenic F (BV1184) backbone with stabilizing triple mutations at positions N67L, S215P and E487Q [17] and produced in *Spodoptera frugiperda* (Sf9) insects cells.

2.2. RSV F Glycoprotein Constructs

Synthetic RSV F genes were cloned into the pFasBac1 (Invitrogen) downstream of the *AcMNPV* polyhedron promoter (GeneArt, Regensburg, DE, USA). RSV F transgenes were cloned into recombinant baculovirus (BV) using the Bac-to-Bac BV system [29]. Sf9 cells (Invitrogen, Grand Island, NY, USA) were cultured in serum-free medium. RSV F proteins were constructed from the near full-length RSV/A2 F gene sequence (Genbank Accession No. U63644) encoding 574 amino acids. The constructs were codon optimized for expression in insect cells. To produce the prefusogenic F construct (BV1184), furin cleavage site II was mutated (KKRKRR → KKQKQQ) to be protease resistant and the adjacent 10 amino acids deleted (F137-V146) from the fusion peptide (Δ FP) on the N-terminus of the F1 subunit and the native transmembrane (TM) and cytoplasmic tail (CT) retained on the C-terminus of the F1 subunit (GenBank Accession No. MN125707) [29,31,36]. Three prefusion F constructs were generated using the prefusogenic F (BV1184) backbone: (1) RSV F DS (BV2267) was produced by introducing disulfide bond double mutations (DS) at positions S155C and S290C within the F1 subunit; (2) RSV F Cav1 (BV2279) construct was generated by introducing hydrophobic amino acid substitutions at positions S190F and V207L; and (3) RSV F DS-Cav1 (BV2280) was constructed with the DS and Cav1 substitutions [22]. Postfusion F (BV2128) was constructed with the native furin cleavage sites I and II and the truncated fusion peptide (Δ FP). N-terminal TM and CT domains were deleted and replaced with a 6-histidine tag (6-H) [30,31].

2.3. Purification of RSV F Proteins

Prefusogenic F (BV1184), postfusion F (BV2128), prefusion F DS (BV2267), Cav1 (BV2279), and DS-Cav1 (BV2280) were produced by infection of Sf9 cells (MOI = 0.5 pfu/cell) with recombinant BV for 67 hours at 27°C. Cells were harvested by centrifugation and RSV F proteins extracted from detergent lysed cell membranes with 25 mM Tris-HCL (pH 8.0), 50 mM NaCl, 0.5% (v/v) Tergitol™ NP-9, and 2.0 μ g mL⁻¹ leupeptin (Sigma-Aldrich, St. Louis, MO, USA). RSV F proteins were purified by a combination of anion exchange, lentil lectin affinity, and cation ion exchange chromatography [29]. For soluble postfusion F (BV2128), Sf9 cells were infected with recombinant BV for 67 hours and supernatants were clarified by centrifugation. RSV F (BV2128) was purified with an immobilized metal affinity column (IMAC) and ion exchange chromatography [31]. Purified RSV F proteins (0.33-1.5 mg mL⁻¹) were formulated in 25 mM phosphate buffer (pH 6.8), 150 mM NaCl, 0.032% (v/v) polysorbate-80 (PS80), and 1% (w/v) histidine.

2.4. Antigenic Site-specific mAb Binding to RSV F Proteins Determined by ELISA

Site-specific mAbs binding to recombinant RSV F proteins was determined by ELISA. Purified prefusogenic F (BV1184) and prefusion F proteins DS (BV2267), Cav1 (BV2279), DS-Cav1 (BV2280), and DS-Cav1-293 were coated (2.0 μ g mL⁻¹) in 96 well micro titer plates overnight at 4°C. Following coating, the plates were blocked with 2% milk blocking buffer (Quality Biologicals, Gaithersburg, MD, USA). Primary mAbs D25 (site Ø), hRSV90 (site V), palivizumab (site II), R1.42 (site IV), R7.10 (p27) were serially diluted (1.0 μ g mL⁻¹ to 0.01 ng mL⁻¹). WHO human RSV reference antiserum (RSV Ref Std) was serially diluted 10⁻² to 10⁻⁸. Plates were incubated for 2 hours at room temperature and washed with phosphate buffered saline containing 0.05% Tween (PBS-T). Horseradish peroxidase (HRP)-conjugated goat anti-human IgG (2040-05, Southern Biotech) or HRP-conjugated goat anti-mouse IgG (1030-05, Southern Biotech) second antibodies were added to the wells. After 1 hour, the wells were washed with PBS-T and 3,3',5,5'-tetramethylbenzidine peroxidase substrate (TMB, T0440-IL, Sigma, St Louis, MO,

USA) added to the wells. Reactions were stopped with TMB stop solution (TSB999, Scytek Lab, Logan, UT, USA). Plates were read at an optical density (OD) 450 nm with a SpectraMax plus plate reader (Molecular Devices, Sunnyvale, CA, USA). EC50 values were calculated by 4-parameter fitting using Prism 7.05 software.

2.5. *Design of Animal Studies*

Animal ethics statement. Noble Life Sciences (Sykeville, MD, USA) performed the murine immunogenicity study. Noble Life Sciences is accredited by the Association for Assessment and Accreditation of Laboratory Animal Care (AAALAC International). Sigmovir Biosystems, Inc. (Rockville, MD, USA) performed the cotton rat immunogenicity and RSV challenge study. Sigmovir is licensed by the United States Department of Agriculture (USDA permit 51-R-0091) and the Office of Laboratory Animal Welfare (OLAW permit A4642-01). The cotton rat study was conducted in accordance with appropriate sections of the National Research Council Guide for the Care and Use of Laboratory Animals and the Animal Welfare Act Regulations. Both studies were performed in accordance with Institutional Animal Care and Use Committee (IACUC) approved protocols.

Murine study design. Female BALB/c mice (7-9 weeks old, 17-22 grams, N = 10 per group) were immunized by intramuscular (IM) injection with 1.0 µg of RSV F prefusion F, prefusion F variants (DS, Cav1, DS-Cav1) or postfusion F adjuvanted with 30 µg aluminum phosphate (Adju-Phos®, Brenntag Biosector, Frederikssund DK) administered in 2-doses spaced 21-days apart (day 0, 21). A separate group (N = 10) received a single intranasal (IN) challenge with 20 µL of 10⁶ pfu RSV/A Long (10 µL each nare) on study day 0. Serum was collected for analysis from individual animals on study day 35 (14-days after the second immunization).

Cotton rat study design. Female cotton rats (4-6 weeks old, 50-65 grams) were randomly assigned to groups (N = 6 per groups) and immunized by IM injection with 1.0 µg of the indicated RSV F protein with 30 µg aluminum phosphate (lot 9496, Brenntag Biosector) administered in 2-doses spaced 21-days apart (day 0, 21). A control group (N = 6) received placebo (non-immunized) and was used as a positive RSV challenge control. A separate group (N = 6) received a single IN challenge with 100 µL of 10⁶ pfu RSV/A Long divided between each nare on study day 0. Serum was collected via the orbital plexus 21-days after the second immunization (day 42) and stored at -20°C until analyzed. Animals were challenged with 100 µL of 10⁵ pfu RSV/A Long (50 µL each nare) 21-days after the second immunization (day 42). Nose and lung homogenates were prepared 4-days post-challenge (day 46) and virus load determined by a plaque assay.

2.6. *Anti-RSV F Glycoproteins IgG ELISA*

An ELISA was used to determine anti-RSV F IgG titers. Briefly, 96 well microtiter plates (Thermo Fischer Scientific, Rochester, NY, USA) were coated with 2.0 µg mL⁻¹ of prefusion F (BV1184), prefusion F (BV2129), DS-Cav1 (BV2280), or postfusion F (BV2128). Plates were washed with PBS-T and blocked with blocking buffer (Quality Biologicals, Gaithersburg, MD, USA). Serum samples were serially diluted (10⁻² to 10⁻⁸) and the plates incubated at room temperature for 2 hours. Following incubation, plates were washed with PBS-T and HRP-conjugated goat anti-mouse IgG or anti-rat IgG (Southern Biotech, Birmingham, AL, USA) added for 1 hour. Plates were washed with PBS-T and TMB peroxidase substrate (T0440-IL, Sigma, St Louis, MO, USA) added. Reactions were stopped with TMB stop solution (ScyTek Laboratories, Inc. Logan, UT). Plates were read at OD 450 nm with a SpectraMax plus plate reader (Molecular Devices, Sunnyvale, CA, USA) and data analyzed with SoftMax software. EC50 values

were calculated by 4-parameter fitting using GraphPad Prism 7.05 software. Individual animal anti-RSV F protein IgG titers and group geometric mean titers (GMT) and 95% confidence interval (\pm 95% CI) were plotted.

2.7. *RSV Neutralizing Assay*

Mouse and cotton rat serum samples were serially diluted in 96 well microtiter plates. Palivizumab and human RSV reference standard (NIBSC 16/284, Potters Bar, Hertfordshire, UK) were used to generate standard curves. RSV/A Long or RSV/B 18537 (TCID₅₀ = 200-350) was added to the wells and incubated for 2 hours at 37°C in 5% CO₂. Following adsorption, wells were seeded with 2.5x10⁴ HEp-2 cells and cultured for 4-days at 37°C in 5% CO₂. After incubation, the cultures were washed and air dried. To detect RSV infection, the cultures were treated with mouse monoclonal (RSV5H5) anti-RSV M2-1 (ab94805, Abcam, Cambridge, MA, USA). HRP-conjugated goat anti-mouse IgG second antibody (catalog number 1030-05, Southern Biotech, Birmingham, AL, USA) was added. TMB substrate was added to the wells and the plates read at OD 450 nm with a plate reader. Neutralizing titers were calculated relative to the human RSV reference standard (2000 IU mL⁻¹) or palivizumab standard (4600 IU mL⁻¹) using a 4-parameter fit using GraphPad Prism 7.05 software. Individual animal titers were plotted with the group GMT \pm 95% CI indicated.

2.8. *Prefusion and Postfusion F Specificity of Antibodies Determined by RSV F Competed ELISA*

Levels of serum antibodies specific for prefusion and postfusion F structures was determined with a competitive ELISA. Briefly, 96 well plates were coated with 2.0 μ g mL⁻¹ prefusion F (BV2129) overnight at 4°C. Immune cotton rat serum or RSV reference antiserum was 3-fold serially diluted and mixed (1:1 ratio) with 20 μ g mL⁻¹ of prefusion F (BV2129) or postfusion F (BV2128) for 1 hour at room temperature. Following incubation, 100 μ L of serum/protein mixture was transferred to the prefusion F coated microtiter wells. The mixture was incubated for 2 hours at room temperature and the wells washed with PBS-T. HRP conjugated goat anti-rat IgG or anti-human IgG was added followed by addition of TMB substrate. Plates were read at OD 450 nm with plate reader and the EC₅₀ values calculated by 4-parameter curve fitting with GraphPad Prism software.

2.9. *Prefusion and Postfusion F Specificity of Neutralizing Antibodies Determined by Competitive ELISA*

Levels of neutralizing antibodies bound by prefusion and postfusion F structures was determined with a competitive ELISA. Cotton rat serum was serially diluted in 96 well microtiter plates and 5 μ g mL⁻¹ prefusion F (BV2129) or postfusion F (BV2128) added for 1 hour at 37°C followed by addition of RSV/A (TCID₅₀ = 200-300) for 2 hours at 37°C. Following incubation, 2.5 x 10⁴ Hep2 cells were added to the wells and the plates incubated at 37°C and 5% CO₂ for 4 days. After incubation, the cultures were washed and air dried. Virus replication was detected with mouse mAb RSV5H5 anti-RSV M2-1 (ab94805, Abcam, Cambridge, MA, USA) followed by addition of HRP-conjugated goat anti-mouse IgG second antibody (catalog number 1030-05, Southern Biotech, Birmingham, AL, USA). TMB substrate was added and the plates read at OD 450 nm with a plate reader. Percent neutralization was calculated compared to the virus control cultures (no serum).

2.10. *Antigenic Site-Specific Antibodies Determined Antibody ELISA*

A competitive antibody binding assay was used to determine the levels of antibodies in mouse and cotton rat serum that competed the binding of RSV F site-specific mAbs [29]. Briefly, monoclonal antibodies D25 (site \emptyset), palivizumab (site II), R1.42 (site IV), and R7.10 (p27) were biotinylated using the

manufacturers protocol (Pierce, Rockford, IL, USA). For sites II, IV and p27 determinations, microtiter plates were coated with 2.0 $\mu\text{g mL}^{-1}$ prefusogenic RSV F protein (BV1184). For site \emptyset determinations, plates were coated with 2.0 $\mu\text{g mL}^{-1}$ prefusion RSV F protein (BV2129). After overnight incubation, the plates were washed with PBS-T and blocked with 2% milk blocking buffer for 1 hour at room temperature. Immune serum was 2-fold serially diluted and spiked with 10 ng mL^{-1} biotinylated D25, palivizumab, R1.42, or 50 ng mL^{-1} R7.10. Biotinylated antibodies without immune serum was used as a no competition control. After 2 hours at room temperature, the plates were washed with PBS-T and 100 μL diluted HRP conjugated Streptavidin (e-Bioscience, San Diego, CA, USA) was added for 1 hour and plates washed with PBS-T. TMB substrate was added, color developed, and reaction stopped with TMB stopping buffer. Plates were read at OD 450 nm with plate reader. Data was analyzed with Softmax pro software (Molecular Devices). Competing antibody concentration was determined from the RSV reference standard curve by interpolating the concentration of the reference standard that corresponds to 50% competition. Antigenic site-specific antibody levels ($\mu\text{g mL}^{-1}$) were plotted for individual animals and the group GMT \pm 95% CI indicated.

2.11. *Plaque Assay*

Nose and lung tissue homogenates were prepared from cotton rats 4-days following IN challenge with RSV/A Long. Homogenates were clarified by centrifugation. Confluent HEp-2 monolayers were infected in duplicate with 50 μL per well of serially diluted homogenate samples in 24-well plates. After 1 hour at 37°C in 5% CO_2 , 0.75% methylcellulose medium overlay added and the cells cultured for 6-days. Cells were fixed with 10% neutral formalin and stained with 0.01% crystal violet solution (1.5 mL/well). Plaques were counted and titers expressed as pfu g^{-1} of nasal or lung homogenate. Virus titers were plotted for individual animals and the group GMT \pm 95% CI indicated.

2.12. *Statistical Analysis*

Statistical comparisons between paired groups was performed with a student t-test and one-way ANOVA using Graph Pad Prism 7.05 software.

3. Results

3.1. *RSV F Glycoproteins Proteins*

We have described a prefusogenic RSV F (BV1184) vaccine candidate constructed from the near full-length RSV/A2 F-glycoprotein that form nanoparticles consisting of 3-5 F-trimers [29,31]. Using the prefusogenic F as a backbone, prefusion constructs were produced by introducing mutations within the hydrophobic cavity to stabilized the prefusion F structure as described by McLellan et al. [22,23]. RSV F DS (BV2267) was constructed with the disulfide bond double mutation (DS) at positions S155C and S290C; RSV F Cav1 (BV2279) with amino acid substitutions at positions S190F and V207L; and RSV F DS-Cav1 (BV2280) with the DS and Cav1 mutations (**Figure 1A and 1B**).

Figure 1. RSV F constructs used in this study. **(A)** Linear diagram of the RSV F glycoproteins constructed from the wild-type RSV/A2 fusion glycoprotein gene sequence (GenBank accession number MN125707). RSV F has two furin cleavage sites (arrows) that flank the intervening 27 amino acid (p27 fragment, green). To produce the prefusogenic RSV F (BV1184), furin cleavage site II was mutated (KKRKRR → KKQKQQ) to be protease resistant and is retained on the N-terminus of the F1 subunit. The first 10 amino acids of the adjacent fusion peptide were deleted (Δ FP, yellow). The native C-terminal native transmembrane (TM, black) and cytoplasmic tail (CT, white) were retained on the N-terminus of F1. The resulting F2 and F1 ectodomain is covalently coupled by two disulfide bonds (brackets). N-linked glycosylation sites are indicated by “Y”; palmitoleic acid (PAM) is indicated by “P”; heptad repeats A, B, and C are indicated by HRA (red), HRB (brown) and HRC (blue). The prefusogenic F structure (BV1184) backbone was used to produce three prefusion F variants. RSV F double mutant (DS) was generated by introducing a stabilizing disulfide bond at positions S155C and S290C within the F1 subunit (designated BV2267). RSV F Cav1 (BV2279) construct was generated by introducing two hydrophobic amino acids at positions S190F and V207L. The RSV F DS-Cav1 construct (BV2280) was generated with S155C/S290C disulfide bond and the S190F/V207L substitutions. Postfusion F (BV2128) was generated with intact furin cleavage sites I and II and truncation of the fusion peptide (Δ FP, yellow). The TM and CT domains were deleted and replaced with a 6-histidine tag (6-H, purple). **(B)** Surface structures of the RSV F prefusogenic and prefusion trimers indicating key neutralizing antigenic sites antigenic sites \emptyset (red), V (pink), II (orange), and IV (blue), and the p27 fragment (green). The native transmembrane (TM, black) and cytoplasmic tail (CT, grey) are indicated. The RSV F postfusion 6-heptarepeat bundle (6HR) is indicated.

3.2. Antigenic Site-specific Binding of mAbs to RSV F Proteins by ELISA

A panel of human and murine mAbs targeting RSV F neutralizing sites was used to characterize the antigenic profile of the RSV F constructs. Using an ELISA method, prefusion-stable RSV F DS (BV2267), Cav1 (BV2279), DS-Cav1 (BV2280), and DS-Cav1-293 were bound by prefusion-specific mAbs D25 (site \emptyset) and hRSV90 (site V) at low antibody concentrations ($ED_{50} = 1.07\text{-}1.66 \text{ ng mL}^{-1}$), indicating neutralizing sites \emptyset and V were intact on these prefusion constructs. Prefusogenic F nanoparticles (BV1184) also by bound by mAbs D25 and hRSV90, although at higher antibody concentrations ($ED_{50} = 47.40 \text{ ng mL}^{-1}$ and 26.80 ng mL^{-1} , respectively) suggesting these prefusion sites are present on prefusogenic F, although at lower levels (**Figure 2A** and **2B**). Humanized palivizumab [9] and murine R1.42 mAbs [28] were used to detect antigenic sites II and IV, respectively. Prefusogenic F and prefusion F variants (DS, Cav1, DS-Cav1 and DS-Cav1-293) bound palivizumab ($ED_{50} = 1.0\text{-}3.0 \text{ ng mL}^{-1}$) and R1.42 ($ED_{50} = 2.4\text{-}6.1 \text{ ng mL}^{-1}$) at low antibody concentrations indicating sites II and IV were present on prefusogenic F and the prefusion F variants (**Figure 2C** and **2D**). Prefusogenic F and prefusion DS, Cav1, and DS-Cav1 variants also bound mAb R7.10 at low concentration ($ED_{50} = 2.7\text{-} 3.17 \text{ } \mu\text{g mL}^{-1}$) consistent with the p27 being present on these F-trimers. DS-Cav1-293 bound R7.10 at a higher concentration ($ED_{50} = 22.1 \text{ } \mu\text{g mL}^{-1}$) compared to the other constructs, suggesting p27 may be less available to antibody binding (**Figure 2E**). All RSV F structures bound WHO RSV reference antibodies equally ($EC_{50} = 1.12\text{E-}04$ to $3.97\text{E-}04$) (**Figure 2F**). These results indicate that antigenic sites \emptyset , V, II, IV, and p27 were present on prefusogenic F and the DS, Cav1, DS-Cav1 and DS-Cav1-293 variants.

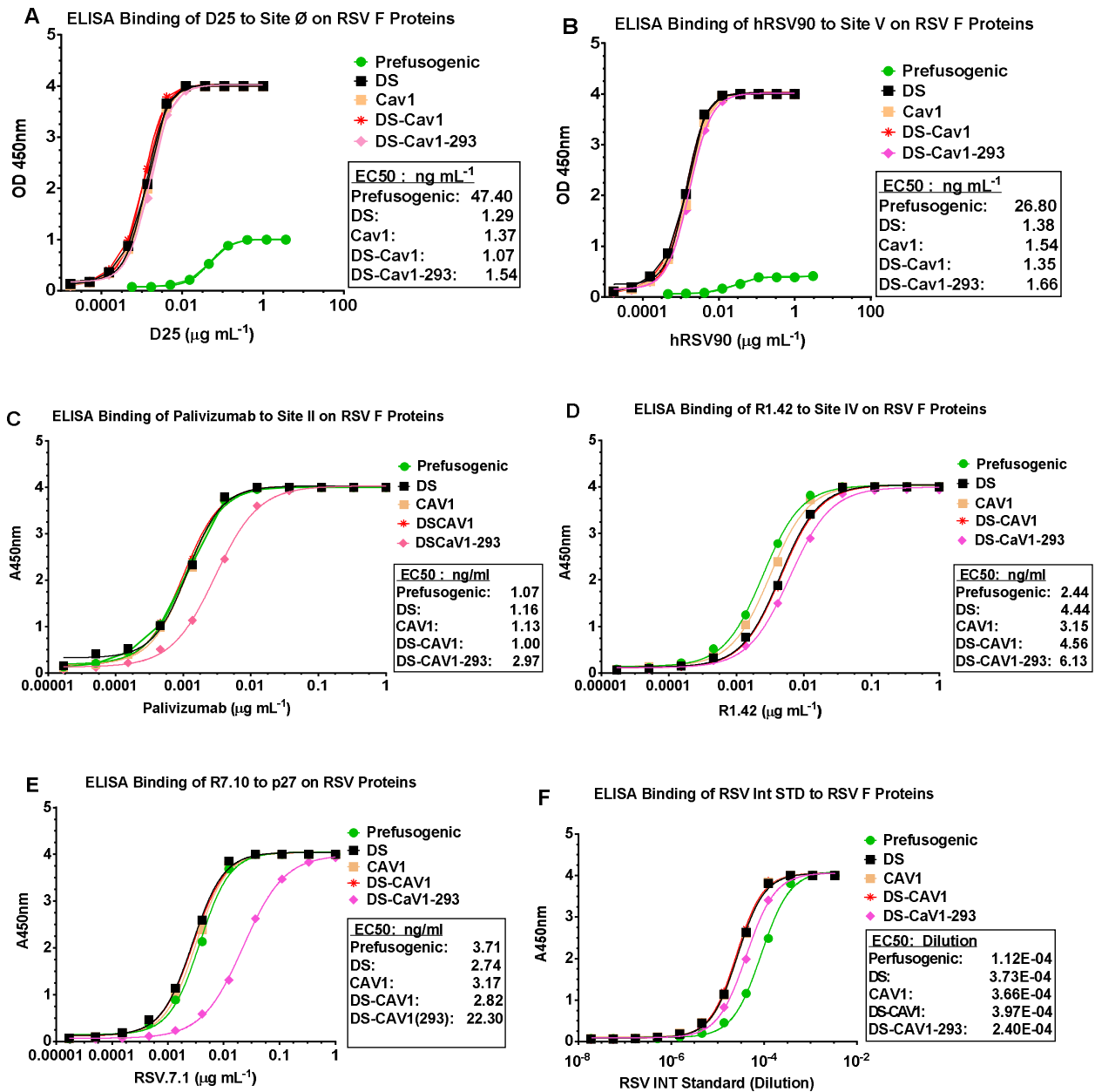


Figure 2. Antigenic site-specific monoclonal antibody binding to prefusogenic F and prefusion F variants. ELISA binding curves (A) D25 (site Ø), (B) hRSV90 (site V), (C) palivizumab (site II), (D) R1.42 (site IV), (E) R7.10 (p27), and (F) RSV international standard (RSV Int Std).

3.3. Immunogenicity of RSV F Proteins in Mice

The immunogenicity of prefusogenic F was compared to prefusion DS, Cav1, and DS-Cav1 F variants in mice. Animals immunized with prefusogenic F (BV1184) had significantly higher ($p < 0.0001$) antibody titers to prefusogenic F BV1184 (GMT = 384,547) compared to immunization with DS (GMT = 2736) or DS-Cav1 (GMT = 1038). Animals immunized with prefusogenic F also had significant higher ($p \leq 0.0007$)

antibody titers to prefusion F BV2129 (GMT = 223,670) compared to animals immunized with prefusion F variants DS (GMT = 20,527) or DS-Cav1 (GMT = 19,195) (**Figure 3A** and **3B**). Immunized with Cav1 (BV2279) also elicited high titer antibodies to prefusion F BV1184 (GMT = 161,953) and prefusion F BV2129 (GMT = 109,422) that were not significantly different from prefusion F (**Figure 3A** and **3B**).

Serum antibodies targeting neutralizing epitopes was determined by competitive ELISA using site-specific mAbs. Mice immunized with Cav1 had the highest levels of antibodies (GMT = 7.8 $\mu\text{g mL}^{-1}$) competitive with D25 (site \emptyset), while serum from mice immunized with prefusion F, DS, or DS-Cav1 variants had significantly lower ($p \leq 0.034$) levels of antibodies that competed D25 binding (GMT = 0.7-1.4 $\mu\text{g mL}^{-1}$). These results suggest that antigenic site \emptyset was poorly immunogenic in mouse regardless of the immunogen (**Figure 3C**).

Mice immunized with prefusion F and Cav1 had significantly higher ($p < 0.0001$) levels of antibodies that competed the binding of palivizumab to site II (GMT = 620.0 and 224.5 $\mu\text{g mL}^{-1}$, respectively), while mice immunized with prefusion F DS or DS-Cav1 had little or no antibodies targeting site II (GMT = 4.0 and 2.5 $\mu\text{g mL}^{-1}$, respectively) (**Figure 3D**). Similarly, animals immunized with prefusion F or Cav1 had significantly ($p < 0.0001$) elevated levels of antibodies that competed binding of R1.42 to site IV (GMT = 676.9 and 2044.56.0 $\mu\text{g mL}^{-1}$, respectively), while animals immunized with DS or DS-Cav1 had no detectable antibodies that competed binding of R1.42 to site IV (**Figure 3E**). Animals immunized with postfusion F also had low levels of antibodies competitive with palivizumab for site II (GMT = 33.8 $\mu\text{g mL}^{-1}$) and variable levels of antibodies competitive with R1.42 for site IV (GMT = 46.7 $\mu\text{g mL}^{-1}$) (**Figure 3D** and **3E**).

Antibody to p27 were also determined. Mice immunized with prefusion F or Cav1 had significantly higher ($p < 0.0001$) levels of antibodies that competed with R7.10 to the p27 peptide (GMT = 112.2 and 80.9 $\mu\text{g mL}^{-1}$, respectively) (**Figure 3F**). In contrast, immunization with DS or DS-Cav1 variants elicited no measurable antibodies to p27 (GMT $< 0.35 \mu\text{g mL}^{-1}$) suggesting that p27 was not accessible to B-cells in DS and DS-Cav1 stabilized F (**Figure 3F**). Animals immunized with the prefusion F, DS, or DS-Cav1 variants had similar levels of RSV/A Long neutralizing antibodies (GMT = 86.2 to 225.4 IU mL^{-1}) (**Figure 3G**). Immunization with postfusion F (BV2128) produced significantly lower ($p = 0.011$) levels of neutralizing antibodies. Taken together, these results demonstrate that wild-type prefusion F was significantly more immunogenic than prefusion F DS or DS-Cav1. Antigenic site \emptyset was poorly immunogenic and elicited little or no antibodies competitive with D25 in mice regardless of the immunogen. Prefusion F and Cav1 were equally immunogenic and elicited antibodies targeting sites II and IV while immunization with prefusion stable DS and DS-Cav1 had little or no antibodies to antigenic sites II or IV or p27, suggesting these sites were not accessible to humoral immune response on the prefusion-stable F structures.

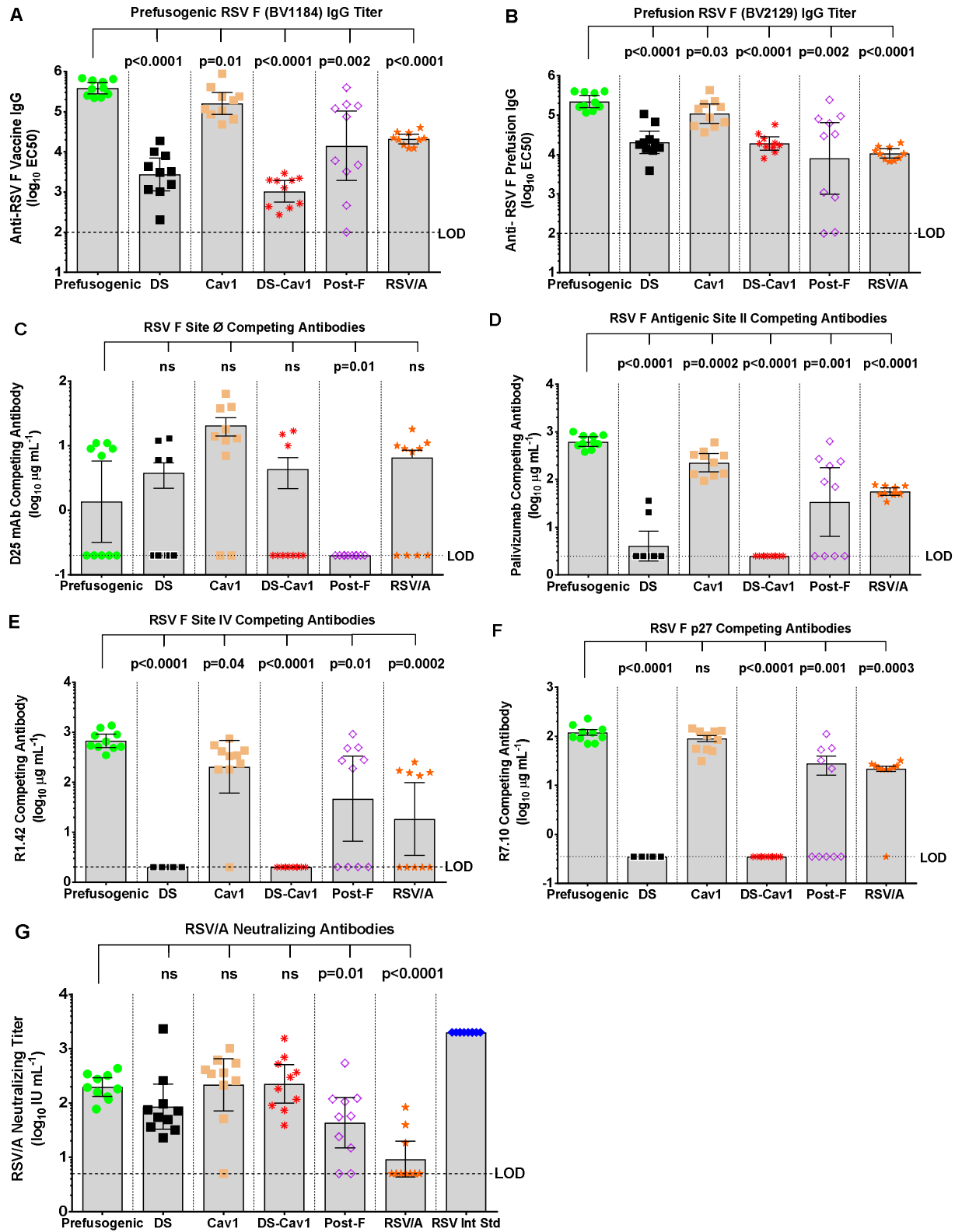


Figure 3. Immunogenicity of prefusogenic F and F variants in mice. Groups of mice were immunized by intramuscular injection with prefusogenic F vaccine (BV1184), DS (BV2267), Cav1 (BV2279), DS-Cav1 (BV2280) or postfusion F (BV2128) adjuvanted on days 0 and 21. A separate group was challenged intranasal with 10^6 pfu RSV/A Long. Serum was collected for analysis from all animals on study day 35 (14-days after the second immunization). **(A)** Anti-prefusogenic F vaccine IgG (BV1184) and **(B)** anti-postfusion F IgG (BV2128) determined by ELISA. A competitive antibody ELISA was used to determine serum levels of antibodies that competed the binding of site-specific mAbs **(C)** D25 (site Ø), **(D)** palivizumab (site II), **(E)** R1.42 (site IV), and **(F)** R7.10 (p27). **(G)** RSV neutralizing antibody titers were determined by inhibition of HEp-2 cell infection with RSV/A Long. The horizontal bars indicate the group (N = 10 per group) geometric mean titer (GMT) and the error bars indicate the 95% confidence interval ($\pm 95\%$ CI). Significance between the RSV F prefusogenic vaccine (BV1184) group and paired RSV F protein immunized groups is indicated. Limit of detection (LOD). Not significant (ns).

3.4. Immunogenicity of RSV F Proteins in Cotton Rats

The immunogenicity of prefusogenic and postfusion F variants was also compared in cotton rats. Animals immunized with prefusogenic F had significantly higher ($p < 0.0001$) titer antibodies to prefusogenic F BV1184 (GMT = 285,691) compared to animals immunized with DS (GMT = 18,361) or DS-Cav1 (GMT = 4590) (**Figure 4A**). Animals immunized with prefusogenic F also had significantly higher ($p < 0.0001$) antibody titers to postfusion F BV2128 (GMT = 164,082) compared to DS or DS-Cav1 (GMT = 3424 and 2177, respectively) (**Figure 4B**). Interestingly, animals immunized with prefusogenic F and DS, Cav1, or DS-Cav1 had equivalent antibody titers to postfusion F (BV2128) and DS-Cav1 (BV2280) (**Figure 4C** and **4D**). These results demonstrate that prefusogenic F elicited high titer antibodies to postfusion and postfusion F structures while cotton rats immunized with DS and DS-Cav1 produced antibodies primarily targeting postfusion structures.

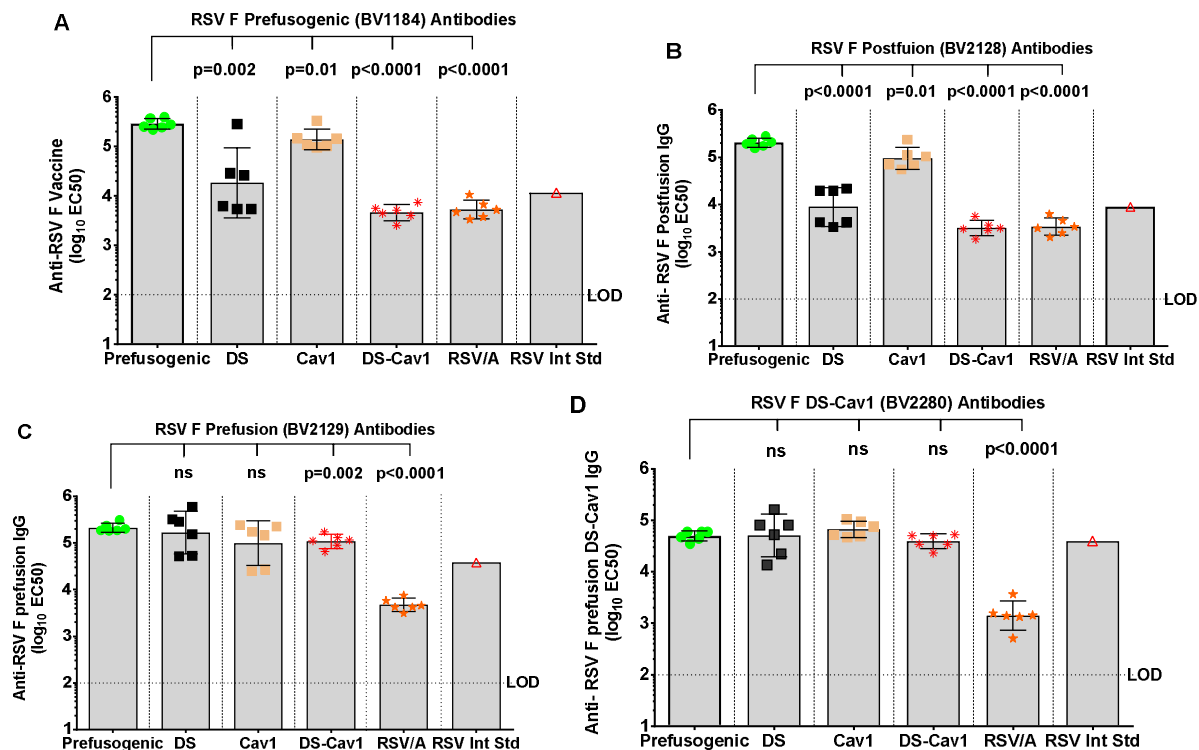


Figure 4. Immunogenicity of prefusion F and F variants in cotton rats. Cotton rats were immunized by intramuscular injection with 1.0 μg of prefusion F vaccine (BV1184), DS (BV2267), Cav1 (BV2279), DS-Cav1 (BV2280) or postfusion F (BV2128) with 30 μg aluminum phosphate on days 0 and 21. A comparative control group was intranasal challenged with 10^6 pfu RSV/A Long. Serum was collected for analysis 21-days after the second immunization (day 42). (A) Anti-prefusion F vaccine IgG (BV1184). (B) Anti-postfusion F IgG (BV2128). (C) Anti-prefusion F IgG (BV2129). (D) Anti-DS F IgG (BV2280). Not significant (ns). Horizontal bars indicated the GMT and error bars indicate the $\pm 95\%$ CI. Statistical significance between the prefusion F (BV1184) compared to paired groups. Limit of detection (LOD). Not significant (ns).

3.5. Prefusion and Postfusion F Antibody Specificity

A competitive ELISA was used to determine the specificity of antibodies to prefusion and postfusion F conformers. Competition with prefusion F (BV2129) resulted in a 288- to 468-fold reduction in prefusion-specific antibodies in immune serum from animals immunized with DS or DS-Cav1 while competition with postfusion F (BV2128) resulted in only a <3-fold reduction in antibody levels, consistent with the majority of antibodies being directed to prefusion F structure (Figure 5A). In contrast, animals immunized with prefusion F or Cav1 produced antibodies that competed with both prefusion and postfusion F structures. Competition with prefusion resulted in 34-fold reduction in prefusion-specific F antibodies while competition with postfusion F resulted in a 13-fold reduction in antibodies, which indicates prefusion F and Cav1 elicited a mix of antibodies that bound both pre- and postfusion F structures (Figure 5A).

We next determined the conformation specificity of RSV/A neutralizing antibodies elicited by immunization with prefusion F compared to immunization with the prefusion F variants. Competition with prefusion F (BV2129) resulting in complete adsorption of neutralizing antibodies in the serum of cotton rats immunized with prefusion F, DS, and DS-Cav1 and near complete adsorption of neutralizing antibodies in serum of Cav1 immunized animals (Figure 5B). In contrast, adsorption with postfusion F (BV2128) resulted in partial or no loss of neutralizing antibodies. These results indicate that immunization with prefusion F, DS, Cav1, and DS-Cav1 elicited neutralizing antibodies that primarily recognize the prefusion F structure (Figure 5B).

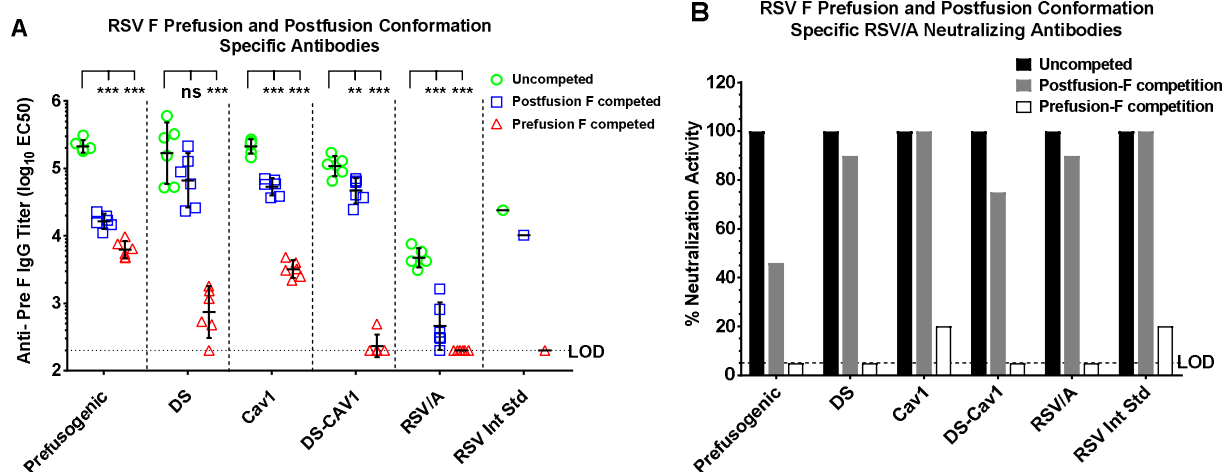


Figure 5. Prefusion and postfusion F specificity of antibodies elicited by prefusion F and prefusion F variants. Cotton rats were immunized and serum analyzed 21 days after the second immunization as

described in **Figure 4**. **(A)** A competitive ELISA was used to determine the level of antibodies targeting prefusion F (BV2129) and postfusion F (BV2128) in serum of immunized cotton rats. **(B)** Conformation-specific neutralizing antibodies in serum competed with prefusion (BV2129) or postfusion F (BV2128). The horizontal bars indicate the group (N = 6 per group) geometric mean titer (GMT) and the error bars indicate the 95%CI. Lower limit of detection (LOD). **p = 0.003, ***p < 0.0001, not significant (ns).

3.6. RSV F Antigenic Site-specific Antibodies

We next determined the specificity of the humoral response to key RSV F neutralizing epitopes with a panel of neutralizing mAbs in a competitive ELISA. Animals immunized with prefusion F DS, Cav1 or DS-Cav1 had significantly higher levels (p<0.001) of antibodies competitive with D25 (GMT = 40.8 – 76.2 $\mu\text{g mL}^{-1}$) compared to animals immunized with prefusogenic F (GMT = 9.0 $\mu\text{g mL}^{-1}$) (**Figure 6A**). Importantly, animals immunized with prefusogenic F or Cav1 had higher levels (p<0.0001) of antibodies competitive with palivizumab (site II) and R1.42 (site IV). In contrast, animals immunized with prefusion DS or DS-Cav1 had significantly lower (p<0.0001) levels of antibodies that were competitive with palivizumab (site II), R1.42 (site IV) or p27 (**Figure 6B**, **6C**, and **6D**). These results are consistent with the mouse immunogenicity result and support the hypothesis that the flexible native prefusogenic F elicits antibodies that recognize neutralizing epitopes on prefusion and postfusion F conformations, while prefusion variants DS and DS-Cav1 produced antibodies primarily targeting prefusion site \emptyset but little or no antibodies to other neutralizing sites (II and IV).

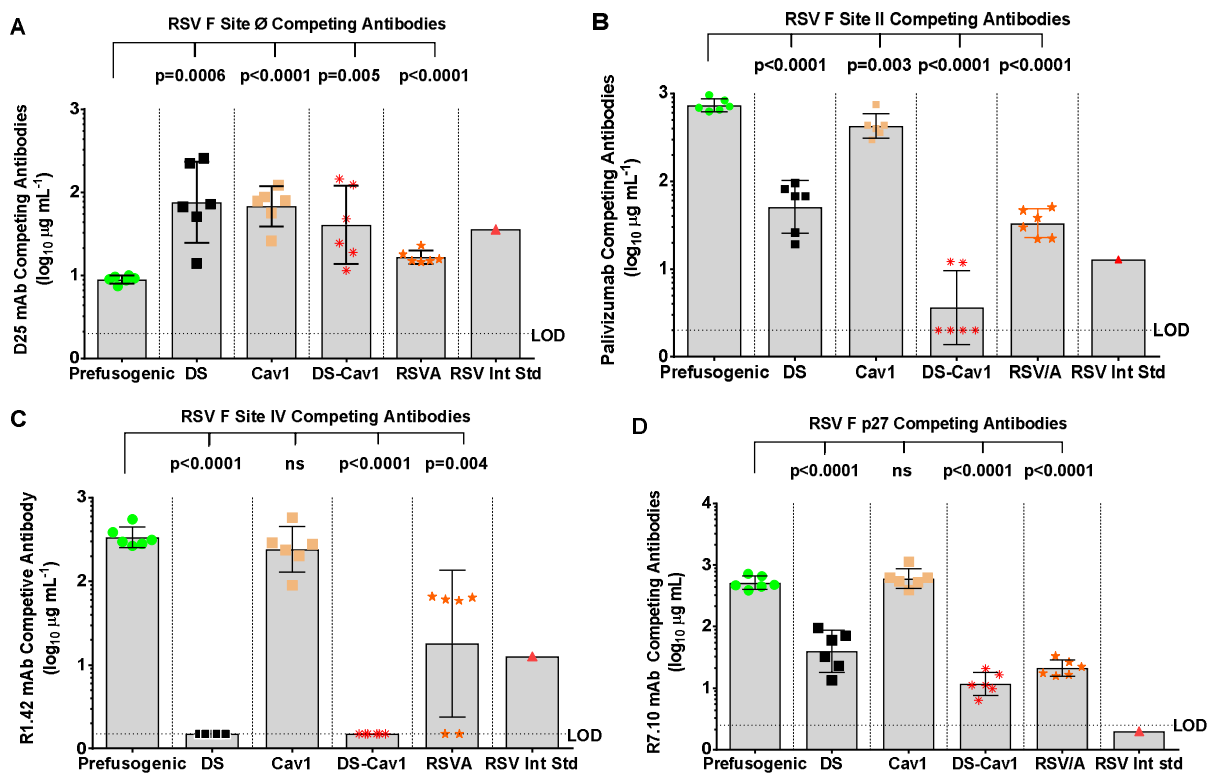


Figure 6. Specificity of antibodies elicited by prefusogenic F compared to prefusion F variants in cotton rats. Female cotton rats were immunized with 1.0 μg RSV F proteins with 30 μg aluminum phosphate and a control group was intranasal challenged with RSV/A. Cotton rats were immunized and serum analyzed 21 days after the second immunization as described in **Figure 4**. The specificity of the humoral response was determined by the competitive antibody binding ELISA using site-specific mAbs: **(A)** D25

(site Ø), (B) palivizumab (site II), (C) R1.42 (site IV), and (D) R7.10 (p27). The horizontal bars indicate the group (N = 6 per group) GMT and the error bars indicate the $\pm 95\%$ IC. Statistical significance between the RSV prefusogenic F group (BV1184) and the paired groups is indicated. WHO RSV International Standard (RSV Int Std). Limit of detection (LOD). Not significant (ns).

3.7. Neutralizing Antibodies and Protection against RSV/A Challenge in Cotton Rats

RSV A and B subtype cross neutralizing antibody titers were determined in cotton rats immunized with prefusogenic F and prefusion F variants. RSV/A neutralizing antibody titers were significantly higher ($p \leq 0.04$) in animals immunized with prefusogenic F or Cav1 (GMT = 2,346 and 1,594 IU mL⁻¹, respectively) compared to animals immunized with DS or DS-Cav1 (GMT = 532 and 282 IU mL⁻¹, respectively) (**Figure 7A**). Significantly, animals immunized with prefusogenic F and Cav1 had 13- to 20-fold higher levels of neutralizing antibodies that cross-neutralized RSV/B (18537) subtype (GMT = 1559 and 1264 IU mL⁻¹, respectively) while animals immunized with DS or DS-Cav1 had significantly lower ($p \leq 0.001$) levels of RSV/B cross neutralizing antibody titers (GMT = 116 IU mL⁻¹ and GMT = 79 IU mL⁻¹, respectively) (**Figure 7B**).

To assess the protective effects of the F proteins, immunized cotton rats were challenged 3 weeks after 2 immunization by intranasal challenge with 10⁵ pfu RSV/A Long. Virus load was significantly lower ($p \leq 0.005$) nose homogenates of all vaccinated animals (GMT = 1,077 – 14,024 pfu g⁻¹) compared to non-immunized animals (GMT = 558,308 pfu g⁻¹). Furthermore, animals immunized with prefusogenic F had significantly lower ($p < 0.01$) virus load in nasal homogenates compared to animals immunized with the DS-Cav1 (GMT = 2587 pfu g⁻¹ vs. GMT = 14,024 pfu g⁻¹, respectively). Nasal virus loads were not significantly in animals immunized with prefusogenic F, DS or Cav1 GMT = 1077 – 9290 pfu g⁻¹) (**Figure 7C**). Virus load in lung homogenates were significantly lower ($p < 0.0001$) in lungs of animals immunized with RSV F proteins were below the limit of detection (LOD <100 pfu g⁻¹) compared to placebo treated animals (GMT = 449,018 pfu g⁻¹) (**Figure 7D**). Animals immunized with RSV F proteins had significantly reduced upper and lower respiratory tract virus load compared to non-immunized cotton rats.

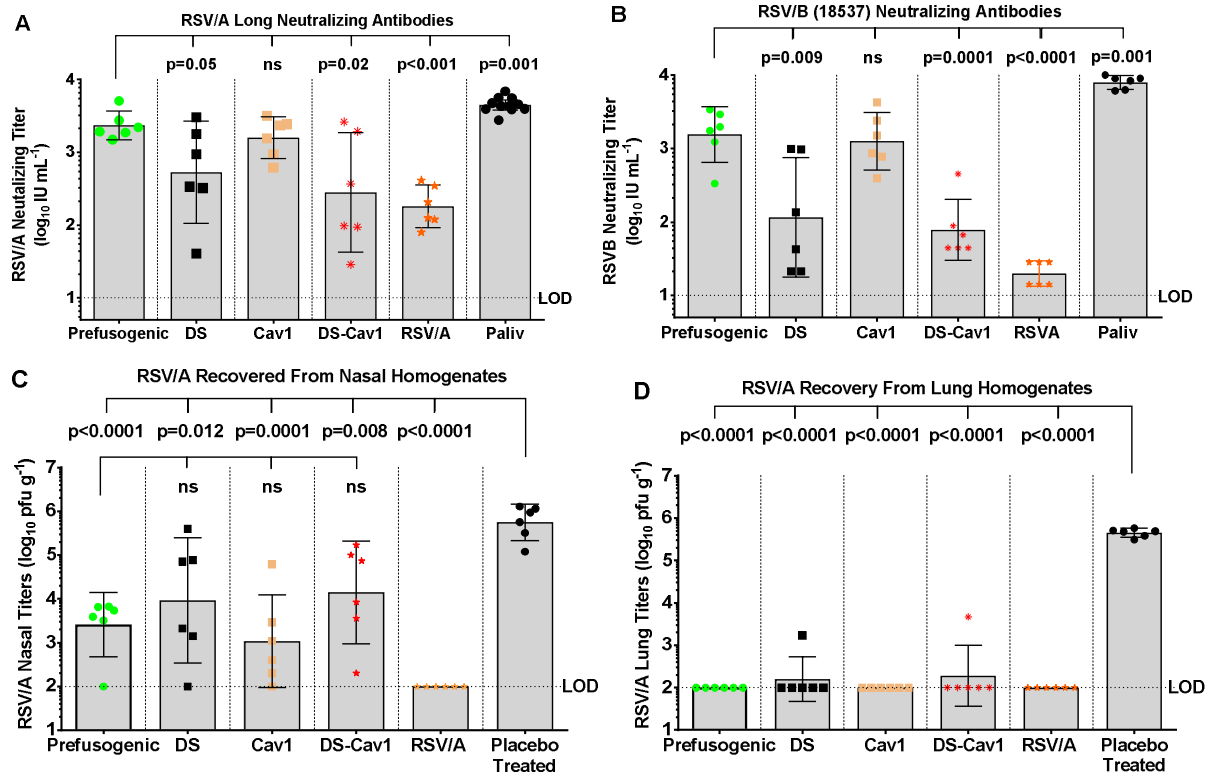


Figure 7. RSV A and B subtype neutralizing antibodies and protection against RSV challenge in cotton rats. (A, B) RSV A and B subtype cross-neutralizing antibody titers in animals immunized with prefusogenic F and prefusion F DS, Cav1 and DS-Cav1. (C, D) Nose and lung tissues were collected 4-days post-challenge (day 46) and infectious virus determined with a plaque assay. Individual animal antibody titers and virus load in nasal and lung homogenates are plotted. The horizontal bars indicate the group GMT and error bars indicate the $\pm 95\%$ CI. Statistical significance between paired groups is indicated by brackets. Limit of detection (LOD). Not significant (ns).

4. Discussion

Prefusion stabilized F DS-Cav1 has been proposed to be a potentially superior vaccine candidate since high affinity neutralizing antibodies bind to conformation-dependent sites \emptyset and V [22,25,35], although there is no clinical evidence to support the assumption that prefusion-specific immunity alone is protective against RSV A and B subtypes [24]. There is however considerable clinical and preclinical evidence that multiple F-specific neutralizing epitopes contribute to protection. For instance, passive immunization of high-risk infants with mAbs palivizumab or motavizumab targeting antigenic site II is protective against developing RSV-LRTI in high-risk infants [3,4,37]. Infants born with maternal anti-F antibodies to site II develop less severe RSV-related disease compared to infants without maternal antibodies [38-40]. Further, we have also shown that infant baboons born to mothers' immunization with prefusogenic F are substantially protected against developing RSV-LRTI compared to infants born to non-immunized mothers [32]. We have shown that women of childbearing age and older adults immunized with near full-length prefusogenic F nanoparticles produce antibodies that bind site II [33,34,41].

Here, we compared the humoral immune response elicited by native prefusogenic F to prefusion F variants. Using flexible prefusogenic F backbone, prefusion stable variants DS, Cav1 and DS-Cav1 were generated. Mice and cotton rats immunized with DS or DS-Cav1 had significant levels of antibodies targeting prefusion-specific sites Ø and V (**Figure 3C, Figure 6A** and **Supplementary Figure S1**) but had little or no antibodies to key neutralizing antigenic sites II or IV (**Figure 3D, 3E** and **Figure 6B, 6C**). Immunization with prefusogenic F also elicited elevated levels of antibodies competitive with D25, although at lower levels compared to the prefusion variants (**Figure 6A**). In contrast, immunization with the flexible prefusogenic F elicited antibodies that recognized prefusion and postfusion F structures (**Figure 5A**). Importantly, immunization with prefusogenic F produced antibodies that broadly cross-neutralized RSV A and B subtypes while immunization with prefusion DS or DS-Cav1 elicited antibodies that were RSV/A neutralizing and little RSV/B cross-neutralizing antibodies (**Figure 7A and 7B**). Collectively, these results show that prefusogenic F was significantly more immunogenic and the flexible structure allowed B-cell access to important neutralizing epitopes that were not available more ridged F structures.

The concept of type 1 trimer breathing, dissociation, and monomerization on the lipid bilayer has been described for the HIV envelope (Env) glycoprotein gp120 and gp41. Lee et al [42-44] have shown that antibody binding to flexible Env gp promotes the dissociation of the Env gp120/gp40, which promotes trimer instability and may be an unrecognized mechanism of protection. These observations support the concept that Env immunogens that closely mimic the flexible Env gp structure may be critical considerations in vaccine design and allows accessibility of B-cells to Env epitopes that promote trimer instability and decay. This same mechanism may also be important consideration for design of RSV F trimeric vaccines. Gillman et al. [19] recently reported that CR9501 mAb binding to RSV F at a specific angle promotes trimer disassembly within the lipid bi-layer. These observations suggest that vaccine strategies targeting type 1 trimers should consider flexible trimeric structures and the plasticity of humoral immunity. Such vaccine strategies may be superior to ridged structures.

Conclusions

In this report, we hypothesized trimeric vaccine strategies that are based on breathable F structures, and provide accessibility of B-cells to epitopes on intermediate conformations, may collectively contribute to broad protective immunity, while less flexible F structures restrict B-cell access to important protective epitopes. We compared the humoral immune responses to prefusogenic F to prefusion stabilized variants. Native prefusogenic F was significantly more immunogenic and elicited antibodies to conformation-dependent and conformation-independent epitopes that broadly cross neutralized RSV A and B subtypes while prefusion variants were less immunogenic and elicited antibodies primarily targeting prefusion F structure. These results have implications for vaccine strategies targeting other type 1 integral membrane proteins.

Supplementary Materials: Appendix A Figure S1. Competitive binding of RSV F site-specific monoclonal antibodies with immune serum determined by bio-layer interferometry (BLI) in cotton rats.

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