Refining the Balance of Attenuation and Immunogenicity of Respiratory Syncytial Virus by Targeted Codon Deoptimization of Virulence Genes

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ABSTRACT Respiratory syncytial virus (RSV) is the most important pathogen for lower respiratory tract illness in children for which there is no licensed vaccine. Live-attenuated RSV vaccines are the most clinically advanced in children, but achieving an optimal balance of attenuation and immunogenicity is challenging. One way to potentially retain or enhance immunogenicity of attenuated virus is to mutate virulence genes that suppress host immune responses. The NS1 and NS2 virulence genes of the RSV A2 strain were codon deoptimized according to either human or virus codon usage bias, and the resulting recombinant viruses (dNSh and dNSv, respectively) were rescued by reverse genetics. RSV dNSh exhibited the desired phenotype of reduced NS1 and NS2 expression. RSV dNSh was attenuated in BEAS-2B and primary differentiated airway epithelial cells but not in HEp-2 or Vero cells. In BALB/c mice, RSV dNSh exhibited a lower viral load than did A2, and yet it induced slightly higher levels of RSV-neutralizing antibodies than did A2. RSV A2 and RSV dNSh induced equivalent protection against challenge strains A/1997/12-35 and A2-line19F. RSV dNSh caused less STAT2 degradation and less NF-kB activation than did A2 in vitro. Serial passage of RSV dNSh in BEAS-2B cells did not result in mutations in the deoptimized sequences. Taken together, RSV dNSh was moderately attenuated, more immunogenic, and equally protective compared to wild-type RSV and genetically stable.

IMPORTANCE Respiratory syncytial virus (RSV) is the leading cause of infant viral death in the United States and worldwide, and no vaccine is available. Live-attenuated RSV vaccines are the most studied in children but have suffered from genetic instability and low immunogenicity. In order to address both obstacles, we selectively changed the codon usage of the RSV nonstructural (NS) virulence genes NS1 and NS2 to the least-used codons in the human genome (deoptimization). Compared to parental RSV, the codon-deoptimized NS1/NS2 RSV was attenuated in vitro and in mice but induced higher levels of neutralizing antibodies and equivalent protection against challenge. We identified a new attenuating module that retains immunogenicity and is genetically stable, achieved through specific targeting of nonessential virulence genes by codon usage deoptimization.

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RESPIRATORY SYNCYTIAL VIRUS (RSV) is the leading cause of lower respiratory tract illness (LRTI) in young children, manifested as bronchiolitis and pneumonia. In the United States, there are 132,000 to 172,000 estimated annual RSV-associated hospitalizations in children less than 5 years of age, with the highest hospitalization rates seen in very young infants (1). RSV-associated LRTI results in an annual 66,000 to 199,000 deaths in children younger than 5 years old globally (2). Prophylaxis currently available to prevent RSV-associated disease is a humanized monoclonal antibody (palivizumab) targeting the RSV fusion (F) protein, but it is prescribed only to infants with certain risk factors (pneumonia). RSV is a member of the Paramyxoviridae family, which contains important human pathogens. RSV carries 10 genes from which 11 proteins are produced. Two promoter-proximal nonstructural (NS1 and NS2) proteins inhibit interferon (IFN) pathways, including type I and type III IFN and potentially type II IFN (6–14). NS1 and NS2 exert their immune-suppressive functions on human dendritic cells (DC) as well as CD4+ and CD8+ T cells (15–17). NS1 and NS2 have also been shown to inhibit apoptosis in infected cells to facilitate viral growth (18). Deletion of either NS1 or NS2 results in virus attenuation, while simultaneously deleting both NS1 and NS2 overattenuates the virus for vaccine purposes (19–22). Combined with other attenuating cold-passage (cp) and/or temperature-sensitive (ts) point mutations, viruses with ΔNS1 or ΔNS2 were evaluated as potential live-attenuated vaccine candidates, and ΔNS1 was highly attenuated, whereas ΔNS2 was underattenuated (19, 20, 22–24). Deletion of nonessential virulence genes provides a limited range of attenuation. Another challenge associated with setting the attenuation level of live-attenuated vaccines containing cp or ts point mutations is reversion or compensatory mutations. This is especially the case for RNA viruses (23, 25, 26), highlighting the need to further stabilize...
vaccine candidates. Attenuating mutations can also be associated with loss of immunogenicity due to reduced replicative fitness, as seen with RSV rA2ΔM2-2 (19, 27).

The codon usage deoptimization strategy was first used to address the problem of genetic instability of live-attenuated poliovirus vaccines (28, 29). Codon deoptimization of the poliovirus capsid gene by incorporation of the rarest codons in the human genome reduced translation of capsid protein, resulting in virus attenuation (28, 29). Another attenuation strategy, codon pair deoptimization, has been used to recode viral genes using rare codon pairs, which does not necessarily alter codon usage (30). In this study, we applied codon usage deoptimization combined with selective targeting of viral immune-suppressive genes to a human pathogen and characterized the genetic stability, replicative fitness, immunogenicity, and protective efficacy of the recoded virus. To our knowledge, this is the first example of virus attenuation by codon deoptimization specifically of nonessential virulence genes. Our results demonstrate that targeting RSV NS1 and NS2 by codon deoptimization can be an effective strategy for developing live-attenuated vaccines with controllable attenuation, wild-type replication in Vero cells, genetic stability, and improved immunogenicity.

RESULTS
Generation of codon-deoptimized NS1 and NS2 RSV. We compared codon usage in the NS1 and NS2 genes of several RSV strains to the codon usage bias of the human genome (31). Of the 18 amino acids used in the RSV NS1 and NS2 genes, 6 (33%) share the same least-used codons as those of human genes. Therefore, because we could not rule out the possibility that RSV utilizes a unique codon usage bias, we designed two mutant viruses with codon-deoptimized NS1 and NS2 genes, namely, dNSv (wherein every codon in NS1 and NS2 is the least used for that amino acid in humans) and dNSv (all NS1 and NS2 codons are the least used by RSV). The dNSv design included 84 silent mutations in NS1 and 82 in NS2, the dNSv design included 145 silent mutations for NS1 and 103 mutations for NS2, and these nucleotide changes were distributed across the coding regions for both genes (Fig. 1). Wild-type NS1 and NS2 genes were replaced by deoptimized NS1 and NS2 genes using MscI and EcoRV sites (Fig. 2). The kRSV-dNSv and kRSV-dNSv mutants (k designates inclusion of the far-red fluorescent protein mKate2 in the first gene position, as described previously [32]) were rescued by reverse genetics, and the sequences of NS1 and NS2 genes were confirmed for all viral stocks. To test the genetic stability of the human codon-deoptimized virus, we serially passaged the virus in three separate lines in BEAS-2B cells at 37°C for 10 passages and sequenced the final passage stocks (P10). All three P10 lines maintained the original deoptimized sequences for both NS1 and NS2 genes.

Reduced NS1 and NS2 protein expression by kRSV-dNSh in vitro. In order to examine the effect of codon deoptimization on NS1 and NS2 expression, HEp-2, BEAS-2B, and Vero cells were infected at a multiplicity of infection (MOI) of 5 with either parental kRSV-A2 or deoptimized virus kRSV-dNSh or kRSV-dNSv. Twenty-four hours postinfection (p.i.), total cell lysates were harvested and analyzed by Western blotting (Fig. 3). Relative steady-state NS1 and NS2 levels were determined by densitometry. Compared to kRSV-A2, human-codon-bias-deoptimized virus (kRSV-dNSh) expressed 75 to 90% less NS1 protein and 70 to 90% less NS2 protein in these cell lines (Fig. 3). In contrast, RSV-codon-bias-deoptimized virus (kRSV-dNSv) expressed higher levels of NS1 and NS2 than did the parental virus, especially in HEp-2 and Vero cell lines (Fig. 3A and C). As the kRSV-dNSh virus exhibited the desired phenotype of reduced NS1 and NS2 levels, we chose this mutant for further studies.

In vitro replication of human-deoptimized NS1/NS2 RSV. Multistep growth curve analyses were done in several cell lines as well as primary, normal human bronchial epithelial (NHBE) cells differentiated at the air-liquid interface (ALI). kRSV-dNSh grew to similar levels as kRSV-A2 in HEp-2 and Vero cell lines (Fig. 4A and B). In BEAS-2B cells, the two viruses replicated to similar levels at earlier time points p.i., but growth of kRSV-dNSh was attenuated at 72 and 96 h p.i. (Fig. 4C). Primary differentiated airway epithelial cells provide a more accurate model than immortalized continuous cell lines for rank ordering RSV attenuation levels (33). We therefore compared the growth kinetics of kRSV-dNSh and kRSV-A2 in differentiated NHBE/ALI cultures. At MOIs of 0.2 and 2.0, kRSV-dNSh virus exhibited a more restricted growth phenotype in these cell cultures, unlike kRSV-A2, which maintained its replication throughout the experiments (Fig. 4D to F). Although the two viruses started with similar levels of infection (Fig. 4F, day 1), only kRSV-A2 infectious yield increased (Fig. 4F). Similarly to previously published data, the RSV-infected NHBE cells exhibited no obvious cytopathic effect over the course of infection (Fig. 4F) (34).

Attenuation, protection, and immunogenicity in BALB/c mice. BALB/c mice were infected with either kRSV-A2 or kRSV-dNSh virus, and lung viral loads were measured at indicated days postinfection. Both viruses peaked between days 4 and 6, and kRSV-dNSh exhibited approximately 1-log10-lower titer than kRSV-A2 on both days (Fig. 5A). As complete protection against RSV challenge is commonly achieved in the BALB/c mouse model with experimental RSV vaccines, we increased the stringency of efficacy determination in this model by using a dose titration of vaccines to evaluate breakthrough of protection. When given as a single vaccination of 10^3 fluorescent focus units (FFU) intranasally (i.n.), both kRSV-A2 and kRSV-dNSh elicited complete protection against heterologous subgroup A RSV strain A/1997/12-35 (12-35 [35]) challenge at 100 days postvaccination (Fig. 5B). Vaccination using kRSV-A2 or kRSV-dNSh with 10^4 or 10^5 FFU resulted in equivalent levels of protection and breakthrough. This protection correlated with induction of RSV-neutralizing antibodies (nAb) over time, and kRSV-dNSh induced slightly but statistically significantly higher levels of nAb than did kRSV-A2 (Fig. 5C). Mice vaccinated with a dose range of either kRSV-A2 or kRSV-dNSh also showed similar levels of protection against challenge with RSV A2-line19F at day 28 postvaccination (Fig. 5D). Taken together, kRSV-dNSh was significantly attenuated in mice but was equally protective and slightly more immunogenic than the parental kRSV-A2 strain.

STAT2 degradation and NF-κB activation. We characterized the effect of kRSV-dNSh infection on STAT2, a known target for NS2 and potentially NS1 (6, 7, 9, 10, 13). 293T cells were mock infected or infected with either wild-type kRSV-A2 or kRSV-dNSh. As expected, kRSV-A2 infection caused 50% STAT2 degradation compared to mock infection. In contrast, we found that kRSV-dNSh infection had no effect on STAT2 levels compared to mock infection (Fig. 6A), suggesting that reduced NS1 and NS2 protein levels may augment host immune responses due to less STAT2 degradation compared to wild-type virus infection.
Early during RSV infection, NS1 and NS2 proteins activate host cell prosurvival signals to promote viral growth, including NF-κB, so that NF-κB activation is a measure of NS1/NS2 function (12, 18). Activation of NF-κB leads to expression of proinflammatory cytokines. Pediatric live-attenuated vaccine strains should preferably be less proinflammatory than wild-type strains for safety concerns. HEK-Blue-Null 1 cells, which contain an NF-κB reporter gene, were mock infected or infected with kRSV-A2 or kRSV-dNSh. Tumor necrosis factor alpha (TNF-α), as a positive control for NF-κB activation, induced a high level of reporter activity, followed by kRSV-A2 virus infection (Fig. 6B). Mock infection and kRSV-dNSh infection resulted in equivalent low levels of NF-κB activation, indicating a reduced NF-κB activation and inflammatory potential of kRSV-dNSh compared to the wild-type virus.

DISCUSSION
Here, we adapted the codon deoptimization strategy to specifically target RSV nonessential virulence genes NS1 and NS2, which function in immune suppression. Deoptimization was based on human codon usage bias, which resulted in reduced target protein expression. Unlike previous RSV attenuation strategies focusing on virus replication per se, diminishing expression of NS1 and NS2 led to attenuation without loss of immunogenicity or infectious yield in Vero cells. Codon deoptimization of nonessential target virulence genes can simultaneously fine-tune attenuation and immunogenicity. The stability of the recoded sequences was also confirmed by in vitro passaging the virus stocks in a restrictive cell line (BEAS-2B) without new mutations.

NS1 and NS2 proteins of RSV suppress type I and type III IFN responses in human epithelial cells and macrophages and have similar effects on mouse cells (11, 13, 15). Homologous genes in related viruses, pneumonia virus of mice (PVM) and bovine respiratory syncytial virus (BRSV), also antagonize type I and type III IFN responses (36–39). Targets of NS1 and NS2 genes include various members of the type I IFN pathways, including STAT2 (6–10, 13, 14). In agreement with these studies, we found that

FIG 1  (A and B) Nucleotide sequence alignment of RSV A2 strain NS1 open reading frame (ORF) with human-codon-deoptimized NS1 (dNS1h) (A) or virus-codon-deoptimized NS1 (dNS1v) (B) ORF. (C and D) Nucleotide sequence alignment of RSV A2 strain NS2 ORF with human-codon-deoptimized NS2 (dNS2h) (C) or virus-codon-deoptimized NS2 (dNS2v) (D) ORF. Nucleotide changes compared to A2 are highlighted in blue in the consensus sequences.
STAT2 protein levels were reduced in kRSV-dNSh infection. There is functional overlap between NS1 and NS2 proteins (6, 9, 11, 12, 15, 17). For example, NS1 and NS2 can cooperatively suppress the maturation of dendritic cells (DC), as ΔNS1/ΔNS2 RSV treatment resulted in higher DC maturation than did ΔNS1 or ΔNS2 treatment (17). Both NS1 and NS2 inhibit IRF3 activation in human epithelial cells (12). Both genes inhibit induction of human alpha interferon (IFN-α), IFN-β, and IFN-λ (11). The overlapping functions of NS1 and NS2 could be explained by their potential interaction and formation of multimeric complexes in the infected cell (7, 13, 40). Collectively, these studies show that double deletion of NS1 and NS2 attenuates the virus more than the single deletions and induces the highest level of antiviral immune responses, supporting our strategy to target both nonstructural genes of RSV.

Deletion of either RSV NS gene or both results in virus attenuation both in vitro and in vivo (11, 19–22, 41). ΔNS1 and ΔNS2 single deletion mutants and the ΔNS1/ΔNS2 double deletion mutant replicate at lower levels than wild-type virus, with ΔNS2 only slightly attenuated and ΔNS1/2 about 2 log10 attenuated in BALB/c mice (15). We found that kRSV-dNSh virus had a milder phenotype than ΔNS1/2 because kRSV-dNSh was 1 log10 attenuated in mice. This is promising because ΔNS1/2 was overattenuated as a vaccine. Also, kRSV-dNSh virus was not attenuated in Vero cells, the presumed vaccine strain producer cells, unlike ΔNS1/2, which exhibited a 20-fold-lower titer in Vero cells (11).

Transcription factor NF-κB activates many host antiapoptotic and proinflammatory genes and is activated by NS1 and NS2 genes rapidly during RSV infection to delay apoptosis (12, 18). Blocking apoptosis by activation of NF-κB to promote viral replication has been documented for other viruses, such as HIV-1, influenza virus, hepatitis B virus (HBV), and HCV (42). We speculate that reduced NF-κB activation by kRSV-dNSh, compared to kRSV-A2, may contribute to the attenuated phenotype of this mutant due to accelerated cell death (Fig. 6B).

Virus attenuation through manipulation of codon usage or codon pair usage in conjunction with synthetic biology has been achieved for several viruses, including poliovirus and influenza virus (28–30, 43, 44). Rather than recoding the protein sequence by using the rarest codons for each amino acid (codon deoptimization), codon pair deoptimization recodes the protein sequence to maximize the occurrence of the rarest adjacent codon pairs. The mechanisms behind virus attenuation using either method are not completely defined. Decreased translational efficiency of codon-deoptimized or codon pair-deoptimized genes is considered the main principle, although mRNA stability was not examined in each case (28, 30, 43, 44). According to the “mutation-selection-drift balance” model, codon bias may be under weak selection for translation efficiency or accuracy, although other mechanisms are proposed, such as mutational bias (45). Studies of tRNA concentration either by gene copy number or by direct measure of cellular tRNA pools have provided a consistent correlation between tRNA abundance and corresponding codon usage frequency, lending support to translational selection on codon bias (46). Codon usage appears to be an important cellular strategy to control protein expression level, activity, function, and ultimately physiology (47–50). Other than selection on translation efficiency, codon usage pattern has also been implicated in regulating the protein-folding process (50, 51). Thus, the production of a functional protein from mRNA is tightly regulated by its codon usage pattern in every step of the process.

We propose codon deoptimization of nonessential virus virulence genes as a general strategy in generating live-attenuated vaccine candidates with retained immunogenicity in vivo, genetic stability, and replication in producer cell lines in vitro. This may be important for RSV vaccines because the wild-type virus is not potently immunogenic and does not grow to high titers in vitro. Additionally, reversion to wild-type virulence in the context of codon deoptimization is minimized due to the additive contribution of silent mutations across the coding region (30), and our in vitro serial passaging experiment provided support for this. Because the recoding method does not change protein sequence, it will maximally preserve immune epitopes. Although more studies are needed to understand the effect of recoding protein sequences...
using either rare codons or codon pairs on protein translation, targeting virulence genes using this method should be widely applicable to many viruses for live-attenuated vaccine development.

MATERIALS AND METHODS

Cell lines. Vero (ATCC CCL-81) and HEp-2 (ATCC CCL-23) cells were maintained in minimal essential medium (MEM) with Earle’s salts and L-glutamine (Gibco) supplemented with 10% fetal bovine serum (FBS) (HyClone) and 1 μg/ml penicillin, streptomycin sulfate, and amphotericin B solution (PSA) (Invitrogen). BEAS-2B cells were maintained in RPMI 1640 (Cellgro) with 10% FBS, as described previously (35). HEK-Blue-Null 1 cells, which express secreted embryonic alkaline phosphatase (SEAP) under the control of the IFN-β minimal promoter fused to NF-κB and AP-1 binding sites, were maintained in Dulbecco’s modified Eagle medium supplemented with 10% FBS, L-glutamine, 4.5 g/liter D-glucose, and 1 μg/ml PSA, as recommended by the provider (InvivoGen, San Diego, CA). BSR-T7/5 cells were a gift from Ursula Buchholz (National Institutes of Health, Bethesda, MD) and were cultured in Glasgow’s minimal essential medium (GMEM) containing 10% FBS and 1 μg/ml PSA. BSR-T7/5 cells were selected with 1 mg/ml Geneticin every other passage.

RSV strains. We performed RSV reverse genetics by cotransfection of five plasmids, an RSV antigenomic cDNA cloned in a bacterial artificial chromosome (BAC) and four codon-optimized helper plasmids that express RSV N, P, M2-1, or L protein, into BSR-T7/5 cells as we described previously (32). The pSynkRSV-line19F BAC produces A2-line19F RSV with the far-red fluorescent protein monomeric Katushka-2 (mKate2) in the first position (32) to mark infected cells. We modified pSynkRSV-line19F by replacing the line 19 strain fusion (F) gene, flanked by SacII-to-SalI sites in the BAC, with a synthetic cDNA (GeneArt) containing the A2 strain F open reading frame (A2 from Barney Graham, Vanderbilt University; GenBank accession number FJ614814) flanked by noncoding regions identical to those in pSynkRSV-line19F and corresponding SacII-to-SalI sites (32). The resulting BAC (pSynk-A2) was used as the genetic background for insertion of codon-deoptimized RSV nonstructural (NS) genes. The PmeI-AvrII fragment from pSynk-A2 BAC (Fig. 2) was subcloned. The MscI-to-EcoRV fragment of pSynk-A2 was replaced with a corresponding synthetic fragment (GeneArt, Life Technologies, Gaithersburg, MD) in which only the NS1 and NS2 open reading frames were codon deoptimized based on either human or viral codon usage to generate kRSV-dNSv or kRSV-dNSh, respectively. Recombinant RSV was...
FIG 4  (A to E) Growth kinetics of kRSV-A2 (open circles) and kRSV-dNSh (closed circles) in HEp-2 (A), Vero (B), and BEAS-2B (C) cells at 37°C infected at an MOI of 0.01, as well as in differentiated NHBE/ALI cells infected at an MOI of 0.2 (D) and 2.0 (E). (F) Time course images for NHBE cells infected at an MOI of 0.2, showing mKate2 fluorescence produced by recombinant viruses. For growth curves in panels A, B, and C, each graph is compiled from two independent experiments performed in duplicate wells. For panels D and E, each single experiment was performed in triplicate wells. *, \( P < 0.05 \) between kRSV-dNSh- and kRSV-A2-infected groups. NHBE, normal human bronchial epithelial cells; ALI, air-liquid interface; LOD, limit of detection.
FIG 5  Attenuation, efficacy, and immunogenicity. (A) BALB/c mice (5 per group) were infected i.n. with either kRSV-A2 (open circles) or kRSV-dNSh (closed circles), and lung viral loads on indicated days are shown. *, $P < 0.05$ comparing kRSV-A2 to kRSV-dNSh at days 4 and 6. (B) BALB/c mice (5 per group) were vaccinated i.n. with various doses of either kRSV-A2 (open circles) or kRSV-dNSh (closed circles) or mock infected (gray circles) and challenged with the RSV 12-35 strain 100 days later. Lung peak viral loads after challenge are shown. Each symbol represents a single mouse. (C) Serum nAb titers measured at indicated days after vaccination with $10^5$ FFU. There were 5 mice per group. *, $P < 0.05$ comparing the bracketed kRSV-dNSh- and kRSV-A2-infected groups. (D) Lung peak viral loads after rA2-line19F challenge 28 days postvaccination. Each symbol represents a single mouse. All data represent one of two replicate experiments with similar results. Dotted lines indicate limit of detection for plaque assay. i.n., intranasal; FFU, fluorescent focus unit; PFU, plaque-forming unit; n.s., not significant.
recovered by transfection as described previously, except that virus stocks were propagated in Vero cells (32). All the virus stocks used in this study were sequenced for the NS1 and NS2 genes and confirmed to be mycoplasma negative using the Venor GeM Mycoplasma detection kit (Sigma-Aldrich, St. Louis, MO). Challenge virus strains A2-19F and RSV 12-35 were generated in HEp-2 cells as described previously and titrated by plaque assay on HEp-2 cells (35).

Normal human bronchial epithelial cells at air-liquid interface (NHBE/ALI). NHBE cells (Lonza, Allendale, NJ) were cultured according to the recommended protocols. Cells were seeded onto collagen-coated (BD Bioscience, Bedford, MA) 24-well transwell supports (Corning Costar, NY) for differentiation. Briefly, 5 × 10^4 cells (100 µl) in B-ALI growth medium (Lonza) were seeded per insert with 500 µl B-ALI growth medium added to the basal chamber. Growth medium in both apical and basal chambers was changed the next day. Air-lift was performed on day 3 by removing B-ALI growth medium from both chambers followed by adding 500 µl B-ALI differentiation medium (Lonza) supplemented with inducer only to the basal chamber. Differentiation medium in the basal chamber was replaced with fresh medium every other day for 21 days before experiments.

Western blotting. Cells (HEp-2, Vero, or BEAS-2B) at 70% confluence were infected with kRSV-A2, kRSV-dNSh, or kRSV-dNSv at an MOI of 5, and cell lysates were harvested 20 h p.i. in RIPA buffer (Sigma-Aldrich, St. Louis, MO) containing 1× protease inhibitor cocktail (Thermo Scientific, Rockford, IL). Proteins were separated by SDS-PAGE and transferred onto polyvinylidene difluoride membranes. Blots were blocked with 5% nonfat milk in Tris-buffered saline (TBS) plus 0.1% Tween 20. Polyclonal rabbit antisera against NS1 and NS2 proteins (gifts from Michael Teng, USF Health) were used to quantify protein levels. Blots were stripped and reprobed with a mouse monoclonal antibody against RSV N protein (clone D14, a gift from Edward Walsh) as a loading control. For STAT2 protein detection, 293T cells at 70% confluence were mock infected or infected with either kRSV-A2 or kRSV-dNSh virus at an MOI of 3. Cell lysates were harvested 12 h later. Blots were probed with rabbit anti-STAT2 polyclonal antibody (C20; Santa Cruz Biotechnology, Santa Cruz, CA). Blots were stripped and reprobed with mouse anti-glyceraldehyde-3-phosphate dehydrogenase (anti-GAPDH) antibody (6C5; GeneTex, Irvine, CA) as a loading control.

**FIG 6** STAT2 degradation and NF-κB activation. (A) 293T cells infected with either kRSV-A2 or kRSV-dNSh or mock infected at an MOI of 3. Twenty hours p.i., total STAT2 protein level was analyzed by Western blotting and densitometry (from three independent experiments). (B) HEK-Blue-Null 1 cells treated with either kRSV-A2, kRSV-dNSh, mock treatment, TNF-α (1 ng/ml), or double-distilled water (ddH2O) for 72 h. Supernatants were measured for reporter activity using a colorimetric assay (from three independent experiments). *, P < 0.05 between the groups indicated by the open brackets.

**Virus growth kinetics.** Seventy percent confluent cells in 6-well plates were infected at an MOI of 0.01 in a volume of 500 µl. After 1-h incubation at room temperature, cells were washed once with 2 ml of phosphate-buffered saline (PBS), and 2 ml MEM with 10% FBS (for the Vero cell line) or RPMI 1640 with 10% FBS (for the BEAS-2B cell line) was added. At
time points p.i., cells were scraped in medium and resuspended, and aliquots were frozen until use. Differentiated NBHE cells were infected at an MOI of 0.2 or 2. Virus inoculum (100 µl) was applied apically after PBS wash (100 µl) followed by a 2-h incubation at 37°C. Inoculum was removed by three apical PBS washes. To collect virus for time points, differentiated medium without inducer (150 µl) was added to the apical chamber and cells were incubated for 10 min at 37°C. This step was repeated twice for each well (300-µl total). This apical supernatant was snap-frozen in liquid nitrogen and stored until use.

**Viral load and protection and vaccine efficacy.** All animal studies were approved by the Emory University Institutional Animal Care and Use Committee (protocol number 20015335) and carried out in accordance with recommendations in the Guide for Care and Use of Laboratory Animals of the National Institutes of Health, as well as local, state, and federal laws. Six- to eight-week-old female BALB/c mice (The Jackson Laboratory, Bar Harbor, ME) maintained under specific-pathogen-free conditions were intranasally (i.n.) infected with 1.6 × 10^6 FFU of virus per mouse. The left lung from each mouse was harvested for viral titer by FFU assay (described above) at days 1, 2, 4, 6, and 8 postinfection. For vaccine protection and efficacy assays, mice were vaccinated with various doses of virus inoculum (100 µl) by intranasal or intramuscular (i.m.) route. Vaccination was performed twice, once at day 0 and again at day 14 postinfection. At day 28 after vaccination, mice were challenged with 100 FFU of virus per mouse. The left lung from each mouse was harvested for viral titer by FFU assay (described above) at days 1, 2, 4, 6, and 8 postchallenge. For all mouse experiments, titers below the limit of detection were scored as 0.5. Titers below the limitation of detection were assigned half the value of the limit of detection.

**Microneutralization assay:** HEp-2 cells were seeded in 96-well plates to attain 70% confluence in 24 h. Heat-inactivated (56°C, 30 min) serum samples were 2-fold serially diluted in MEM and added to 50 to 100 FFU kRSV-A2 in an equal volume. The virus and serum mixture was incubated at 37°C for 1 h. Then, half of the serum-virus mixture was transferred onto HEp-2 cell monolayers in 96-well plates in duplicate, and plates were incubated at 2,000 × g for 30 min at 4°C. Fluorescent foci were counted 36 h p.i. The 50% effective concentration (EC_{50}) was calculated using a nonlinear regression analysis with four-parameter fitting in GraphPad Prism version 6.0.

**NF-κB activation assay:** NF-κB activation was assayed according to the manufacturer’s protocol (InviVoGen, San Diego, CA). Briefly, HEK-Blue-Null 1 cells were seeded at 5 × 10^3 cells/well in a 96-well plate and infected with either kRSV-A2 or kRSV-dNSh or mock infected for 72 h. Supernatants were incubated with Quant-i-Blue (InviVoGen, San Diego, CA) substrate prepared according to instructions for 1 to 3 h before reading optical density at 630 nm on a microplate reader (Bio-Tek, Winooski, VT).

**Statistical analysis:** Statistical analysis was performed using GraphPad Prism software version 6.0 (San Diego, CA). Data are represented as means with standard deviation (SDs). One-way and two-way analyses of variance (ANOVA) with Tukey’s post hoc test with a P value of 0.05 were used, as indicated. Student’s t test (unpaired, two-tailed) was used for Fig. 6A.

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M.L.M. and Emory University are entitled to licensing fees derived from various agreements Emory has entered into related to products used in the research described in this paper. This study could affect his personal financial status. The terms of this agreement have been reviewed and approved by Emory University in accordance with its conflict of interest policies.

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