A Glycoprotein Subunit Vaccine Elicits a Strong Rift Valley Fever Virus Neutralizing Antibody Response in Sheep

Bonto Faburay,1 Maxim Lebedev,2 D. Scott McVey,3 William Wilson,3 Igor Morozov,1 Alan Young,2 and Juergen A. Richt1

Abstract

Rift Valley fever virus (RVFV), a member of the Bunyaviridae family, is a mosquito-borne zoonotic pathogen that causes serious morbidity and mortality in livestock and humans. The recent spread of the virus beyond its traditional endemic boundaries in Africa to the Arabian Peninsula coupled with the presence of susceptible vectors in nonendemic countries has created increased interest in RVF vaccines. Subunit vaccines composed of specific virus proteins expressed in eukaryotic or prokaryotic expression systems are shown to elicit neutralizing antibodies in susceptible hosts. RVFV structural proteins, amino-terminus glycoprotein (Gn), and carboxyl-terminus glycoprotein (Gc), were expressed using a recombinant baculovirus expression system. The recombinant proteins were reconstituted as a GnGc subunit vaccine formulation and evaluated for immunogenicity in a target species, sheep. Six sheep were each immunized with a primary dose of 50\(\mu\)g of each vaccine immunogen with the adjuvant montanide ISA25; at day 21, postvaccination, each animal received a second dose of the same vaccine. The vaccine induced a strong antibody response in all animals as determined by indirect enzyme-linked immunosorbent assay (ELISA). A plaque reduction neutralization test (PRNT\(_{80}\)) showed the primary dose of the vaccine was sufficient to elicit potentially protective virus neutralizing antibody titers ranging from 40 to 160, and the second vaccine dose boosted the titer to more than 1280. Furthermore, all animals tested positive for neutralizing antibodies at day 328 postvaccination. ELISA analysis using the recombinant nucleocapsid protein as a negative marker antigen indicated that the vaccine candidate is DIVA (differentiating infected from vaccinated animals) compatible and represents a promising vaccine platform for RVFV infection in susceptible species.

Key Words: Rift Valley fever virus—Glycoproteins—Subunit vaccine—Neutralizing antibodies—Sheep.

Introduction

Rift Valley fever virus (RVFV) is a mosquito-borne zoonotic pathogen that causes high morbidity and mortality in both livestock and humans. The virus has caused outbreaks in ruminants and humans in Africa and the Arabian Peninsula (Flick and Bouloy 2005) and represents a threat to the well-being and livelihood of resource-poor farmers in much of Africa (Smith et al. 2013). It is classified as a select agent and risk group-3 pathogen by the Centers for Disease Control and Prevention (CDC) and the United States Department of Agriculture (USDA). In ruminant livestock, RVF is characterized by high mortality in young animals, notably in lambs, fetal malformations, and widespread abortion storms; sheep are the most susceptible, with neonatal mortalities approaching 100% (Swanepoel and Coetzer 1994). Human infections are often characterized by benign fever, but in a small proportion of individuals RVF could lead to more serious complications such as retinitis, encephalitis, neurological disorders, hepatic necrosis, or fatal hemorrhagic fever (Bird et al. 2009). Although human fatal hemorrhagic cases have been historically estimated at 2% in infected
individuals, case fatalities of up to 20% have been reported (Bird et al. 2009), including the recent outbreak in Mauritania (Heald 2012). The reasons or factors contributing to this increase in fatalities are unknown.

RVFV belongs to the genus *Phlebovirus* within the family Bunyaviridae, which includes over 350 named isolates (Walter and Barr 2011). It has a tripartite single-stranded RNA genome of negative polarity consisting of small (S), medium (M), and large (L) RNA segments. The M segment encodes two structural glycoproteins, amino-terminus glycoprotein (Gn) and carboxyl-terminus glycoprotein (Gc), the 78-kDa protein, and the nonstructural protein, NSm, whereas the S segment encodes the nucleocapsid protein (N) protein and the nonstructural protein NSs. The L segment encodes an RNA-dependent RNA polymerase (Elliott 1996). The N and L proteins are required for viral RNA synthesis. The NSs protein, the major virulence factor, has been shown to inhibit host transcriptional immune response (Bouloy et al. 2001) through generalized transcription downregulation, including repression of interferon-β (IFN-β) (Billecocq et al. 2004, Le May et al. 2004, Le May et al. 2008) and degradation of protein kinase R (Habjan et al. 2009, Ikegami et al. 2009). The NSm protein functions to suppress virus-induced apoptosis (Won et al. 2007) and there is evidence that it plays a functional role in the vector competence of mosquitoes for RVFV at the level of the midgut barrier (Kading et al. 2014). The glycoproteins Gn and Gc are surface proteins that play a role in virus attachment to initiate infection and have been shown to carry epitopes that elicit the production of neutralizing antibodies, a correlate of protective immunity (Besselaar et al. 1991, Besselaar and Blackburn 1992, Besselaar and Blackburn 1994).

There are currently no RVFV vaccines fully approved for commercial use outside its endemic area in Africa and the Arabian Peninsula. Given the potential for viral spread elsewhere, including the mainland United States, there is an urgent need for a safe and efficacious vaccine. Attributes essential for a vaccine for use in nonendemic areas include safety and the ability to generate a rapid (with primary vaccination) protective immune response in a susceptible host. In endemic regions, RVFV in livestock has been controlled traditionally by using the live-attenuated Smithburn strain or inactivated whole virus (Grobbelaar et al. 2011). The Smithburn vaccine is highly immunogenic, but it is teratogenic in pregnant sheep and cattle (Coetzer and Barnard 1977, Botros et al. 2006). The formalin-inactivated whole-virus vaccines are safe but less immunogenic (Lubroth et al. 2007). Clone 13, another live attenuated natural mutant, is now in commercial use in South Africa for the control of the disease in livestock (Muller et al. 1995, Dungu et al. 2010). MP12, a chemically attenuated virus derived from ZH548, an Egyptian wild-type isolate (Caplen et al. 1985, Vialat et al. 1997), is being evaluated as a potential vaccine for human and veterinary use. The immunogenicity and pathogenicity of these latter two candidate vaccines have been evaluated in various animal species (Muller et al. 1995, Morrill et al. 1997), and, although both vaccine candidates showed promising results, MP12 was reported to induce fetal malformations during the first trimester (Hunter et al. 2002); however, a recent study reported the absence of fetal malformation in pregnant ewes inoculated with the virus (Morrill et al. 2013).

Strategies to develop RVFV vaccines include subunit (Schmaljohn et al. 1989, Naslund et al. 2009, Mandell et al. 2010a), DNA (Spik et al. 2006, Lagerqvist et al. 2009), virus-like particles (VLPs) (Naslund et al. 2009, de Boer et al. 2010, Kortekaas et al. 2012), virus replicon particles (Kortekaas et al. 2011, Dodd et al. 2012, Oreshkova et al. 2013), virus-vectored (Wallace et al. 2006, Heise et al. 2009) modified live vaccines, developed from recombinant viruses engineered using reverse genetics (Ikegami et al. 2006, Bird et al. 2008, Billecocq et al. 2008, Habjan et al. 2008, Bird et al. 2011), live attenuated (Smithburn 1949, Caplen et al. 1985, Muller et al. 1995, Dungu et al. 2010, Pittman 2012, Morrill et al. 2013), and inactivated whole virus vaccines (Pittman et al. 2000). Although subunit vaccines for RVFV are generally considered safe, and recently some progress has been made in their development, evaluation of immunogenicity and/or efficacy in a target species, sheep, has been performed for a few candidates (Kortekaas et al. 2012, Oreshkova et al. 2013). On the other hand, production of live vaccines requires high level of biosafety, and their use is associated with potential risk to personnel and reversion to virulence in vaccinated hosts. Therefore, the general availability of a safe, efficacious vaccine with DIVA (differentiating infected from vaccinated animals) compatibility will be extremely valuable to endemic and nonendemic countries outside Africa.

In this study, we describe the expression of the RVFV structural glycoproteins Gn and Gc and assess their immunogenicity in a natural target species, sheep. The results demonstrate the induction of a strong virus neutralizing antibody response with potentially protective titers induced within 2 weeks after the primary vaccination without any noticeable adverse reactions. Taken together, the findings suggest that a baculovirus-expressed RVFV glycoprotein subunit vaccine could be a potential viable candidate for development of an effective vaccine against RVFV.

**Materials and Methods**

**Cell cultures**

The African green monkey cells, Vero E6 (American Type Culture Collection [ATCC], Manassas, VA), were maintained in Eagle’s minimal essential medium (Corning Cellgro, Manassas, VA) supplemented with 10% fetal bovine serum (FBS), L-glutamine, and penicillin-streptomycin (Invitrogen–Life Technologies, Carlsbad, CA). The cultures were maintained at 37°C in humidified 5% CO₂ atmosphere. The *Spodoptera frugiperda* (Sf9) cells (Invitrogen–Life Technologies) were maintained in SFM 900 II medium supplemented with 10% FBS and penicillin-streptomycin (Invitrogen–Life Technologies). These cells were maintained at 27°C.

**Construction of recombinant bacmid**

The cloning and construction of the recombinant bacmid was carried out as described previously (Faburay et al. 2013). Briefly, the coding sequences of the RVFV structural proteins Gn and Gc were retrieved from the RVFV ZH548 isolate sequences (GenBank accession no. DQ380151), and the sequences were synthesized (GENEWIZ, Inc., San Diego, CA) with molecular modifications. For Gn, the transmembrane domain and cytoplasmic tail were deleted from the coding.
sequence and designated as the Gn ectodomain (Gne) (Fig. 1A). To ensure translocation of both Gne and Gc into the endoplasmic reticulum (ER) and processing through the cellular glycosylation pathway, a specific 54-nucleotide signal peptide (SP) sequence was identified in the M segment of the Rift Valley fever virus (RVFV) genome, as described previously (Faburay et al. 2013) (sequences are available on request). This SP sequence was fused to the 5′ terminus of Gne and Gc. (γ) Putative glycosylation sites (Gerrard and Nichol 2007). (C) Coomassie Blue stain of purified recombinant Gne, Gc, and empty baculovirus-infected, mock-purified cell lysate (EBM). M, molecular weight marker.

FIG. 1. Creation of amino-terminus glycoprotein ectodomain (Gne) and carboxyl-terminus glycoprotein (Gc) recombinant constructs and recombinant protein expression. (A) Diagrammatic representation of deletion of the transmembrane (TM) domain (black boxes) and the cytoplasmic tail of Gn. (B) Diagrammatic representation of signal peptide (SP) with a signal peptidase cleavage site (scissors) identified in the M segment of the Rift Valley fever virus (RVFV) genome is fused to the 5′ terminus of Gne and Gc. (γ) Putative glycosylation sites (Gerrard and Nichol 2007). (C) Coomassie Blue stain of purified recombinant Gne, Gc, and empty baculovirus-infected, mock-purified cell lysate (EBM). M, molecular weight marker.

The nucleoprotein (N) nucleotide sequence was amplified by PCR from the pET30 Ek/LIC recombinant plasmid (a gift from Dr. Friedeman Weber, University of Marburg, Germany), which contained the entire coding region of the S segment of the RVFV strain ZH548. The RVFV-specific sequences were cloned into a pFastBac/CT-TOPO vector (Invitrogen–Life Technologies) in-frame with a carboxy-terminal polyhistidine-tagged protein, resulting in the donor plasmids pRF-Gne, pRF-Gc, and pRF-N. The individual donor plasmids were transformed into MAX Efficiency DH10Bac-competent E. coli to generate recombinant bacmids. Recombinant bacmids were purified and used for transfection to express the respective proteins.

Recombinant baculovirus expression and purification of RVFV proteins

Recombinant baculovirus expression of the respective RVFV proteins was carried out as described previously (Faburay et al. 2013). Briefly, purified recombinant bacmids
carrying the respective coding sequences (Gne, Gc, and N) of RVFV were transfected, using Cellfectin II Reagent (Invitrogen–Life Technologies), into SF9 cells (Invitrogen–Life Technologies) grown in SF-900 II SFM medium (Invitrogen–Life Technologies) supplemented with 10% FBS and 100 U/mL 100 µg/mL penicillin-streptomycin. Protein expression was carried out using P2 or higher-passage recombinant baculovirus stock (>10^7 plaque-forming units (pfu)/mL). Western blot analysis of the lysate was performed to confirm specific protein expression as described below. The proteins were expressed with a carboxy-terminal 6xHistag, and purification using Ni-NTA Superflow resin (Novagen, Rockland, MA) was performed according to the protocol described previously (Faburay et al. 2013). The purified proteins were stained with Coomassie Blue, and protein concentrations were determined using the bicinchoninic acid (BCA) assay (Thermo Scientific, Rockford, IL) at an absorbance of 562 nm, using bovine serum albumin (BSA; Sigma-Aldrich, St. Louis, MO) as the protein standard. Aliquots were stored at −80°C until used.

Expression and purification of RVFV N protein

Recombinant Escherichia coli expression of RVFV N protein was carried out using an expression construct kindly provided by Dr. Friedeman Weber (University of Marburg, Germany). The E. coli cells B21(DE3)pRARE harboring the expression vector pCDNA3.1 were grown on Luria–Bertani (LB) agar plates containing 100 µg/mL ampicillin and 34 µg/mL chloramphenicol. A single clone was picked and inoculated into 10 mL of LB medium for overnight growth. Ten milliliters of the overnight culture was then inoculated into 1 liter of LB medium, and expression was induced with the addition of 0.4 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) when optical density at 600 nm (OD600) reached 0.6. The culture was grown overnight at 15°C. Cells were harvested by centrifugation at 4500 × g for 15 min at 4°C, resuspended in 1 mL of lysis buffer (50 mM Tris-HCl, pH 8.0, 150 mM NaCl), and lysed by three freeze–thaw cycles and then sonicated three times for 5 sec each (Sonic Dismembrator, Model 100, Fischer Scientific, Inc.). The lysate was fractionated by centrifugation for 30 min at 10,000 × g. The supernatant was clarified through a 0.45-µm filter and then loaded onto a 5-mL HisTrap HP column (GE HealthCare) in conjunction with an AKTA Xpress purification system (GE HealthCare) at a flow rate of 1 mL/min. The columns were washed with Buffer A (50 mM Tris, 300 mM NaCl, pH 8.0) over 10 column volumes. Nonspecifically bound proteins were removed by washing with five column volumes of 10% Buffer B (50 mM Tris, 300 mM NaCl, pH 8.0, 500 mM imidazole). Protein was eluted from the column with a gradient of 10–100% Buffer B over 10 column volumes. The elution fractions, which contained the purified N protein, were pooled and stored at −80°C after addition of an equal volume of 100% glycerol.

Detection of recombinant protein expression and analysis of immunoreactivity

The procedure for detection of recombinant protein expression by western blot has been described previously (Faburay et al. 2013). Expression of the recombinant proteins N and Gne was further confirmed using primary antibodies, mouse anti-N (R3-ID8), and the mouse anti-Gn monoclonal antibody 4D4 (a gift from Dr. Connie Schmaljohn, United States Army Medical Research Institute for Infectious Diseases), respectively, at a dilution of 1:2000. To probe immunoreactivity of antiserum obtained from sheep vaccinated with recombinant RVFV Gne and Gc glycoprotein subunit vaccine, the blots were incubated with individual sheep sera at 1:100 dilution for 1 h at room temperature. After washing, the membrane was incubated for 1 h at room temperature with Protein G-HRP (Abcam, Cambridge, MA) diluted 1: 25,000. After the final washing steps, specific reactivity was detected using 3-amin-9-ethyl-carbazole (AEC) peroxidase substrate (Sigma-Aldrich, St. Louis, MO) or an enhanced chemiluminescent (ECL) detection system.

Preparation of the vaccine, animal immunization, and MP12 virus antiserum

The purified glycoproteins were formulated in montanide ISA25 water-in-oil adjuvant (Seppic, France) to obtain a concentration of 50 µg of each immunogen per vaccine dose according to the manufacturer’s instruction. Six adult Rambouillet breed sheep (#9, #36, #163, #169, #170, #179) were each immunized subcutaneously with a primary dose of 50 µg of purified Gne and 50 µg of purified Gc. At day 21 after primary vaccination, hereafter referred to as post-vaccination (pv), each sheep was given a booster with the same amount of vaccine (a second dose). At different time points after vaccination, the injection sites were inspected for any possible adverse reaction. Blood samples were collected from the jugular vein of each sheep for the separation of sera. Prior to vaccination (day 0) to establish a baseline pre-vaccination immune response status. Thereafter, serum samples were collected from each sheep weekly on days 7, 14, 21, 28, 35, 42, and 49 pv. All sera were stored at −80°C until used. The RVFV MP12 control antiserum was obtained from a sheep at day 28 postinfection in a previous experiment conducted at the Kansas State University Biosecurity Research Institute (BRJ (Faburay et al. 2013). Animal experiments complied with institutional guidelines of South Dakota State University and were approved (Committee approval no. 12-037A) by South Dakota State University Institutional Animal Care and Use Committee (IACUC).

Immunogen-specific antibody response

Specific antibody response in serum was measured by enzyme-linked immunosorbent assay (ELISA) using nickel column–purified baculovirus-expressed RVFV Gne, Gc, and E. coli-expressed RVFV N proteins. The N protein was expressed in E. coli using an expression construct kindly provided by Friedeman Weber (University of Marburg, Germany) and was used as a negative diagnostic marker antigen to assess the DIVA compatibility of the recombinant subunit vaccine. To perform an ELISA, each well in a 96-well plate (Nunc, Maxisorp) format was coated overnight at 4°C with approximately 100 ng of each purified recombinant protein in 100 µL of Dulbecco’s coating buffer (pH 7.4) (Invitrogen–Life Technologies). The ELISA assays were performed as described previously (Faburay et al. 2013). The cutoff point was determined for each sheep in each specific ELISA by the addition of two standard deviations to the corresponding mean OD value of the prevaccination serum. Mean OD values equal to or greater than the cutoff value were considered positive.
Anti-RVF virus plaque-reduction neutralization 80% titers

The stock of MP12 RVFV was diluted to 50 plaque-forming units (pfu) in 250 μL of 1 × minimum essential medium (MEM) containing 4% BSA (Sigma-Aldrich, St. Louis, MO). Separately, aliquots of serum from each vaccinated sheep were diluted as follows: 1:10, 1:20, 1:40, 1:160, 1:320, 1:640, and 1:1280 in 1 × MEM containing 2% BSA and 1% penicillin-streptomycin. Diluted serum (250 μL) was mixed with an equal volume of diluted MP12 virus and incubated at 37°C for 1 hr. Thereafter, each mixture of serum plus RVFV was used to infect confluent monolayers of Vero E6 cells in 12-well plates.

After 1 hr adsorption at 37°C and 5% CO₂, the mixture was removed, and 1.5 mL of nutrient agarose overlay (1 × MEM, 4% BSA, and 0.9% SeaPlaque agarose) was added to the monolayers. After 5 days of incubation, the cells were fixed with 10% neutral buffered formalin for 3 hr prior to removal of the agarose overlay. The monolayer was stained with 0.5% Crystal Violet in phosphate-buffered saline (PBS), and plaques were enumerated. The calculated plaque reduction neutralization 80% titers (PRNT₈₀) corresponded to the reciprocal titer of the highest serum dilution, which reduced the number of plaques by 80% or more relative to the virus control. As positive neutralizing serum control, a 1:40 dilution of day-28 serum obtained from a sheep previously immunized with RVFV GneGc VLP was used. The serum showed a PRNT₈₀ titer of 1280 (data not shown). Studies in humans and nonhuman primates show that a PRNT₈₀ titer of ≥ 40 is protective against virulent RVFV challenge (Pittman et al. 2000, Papin et al. 2011); thus, in this study a PRNT₈₀ titer of ≥ 40 was considered potentially protective.

Electron microscopy

Co-expression of the RVFV glycoproteins Gne and Gc in cell culture has been shown to result in the formation of VLPs (de Boer et al. 2010, Mandell et al. 2010b). Thus, to rule out the possibility of assembly of recombinant Gne and Gc upon mixing of the proteins into vaccine formulation, an examination by transmission electron microscopy (TEM) was performed. Briefly, equal amounts of purified Gne and Gc were mixed together in a single tube and incubated for 30 min at room temperature. As controls, aliquots of purified Gne and Gc were also made in separate tubes and incubated for 30 min as described above. Subsequently, the proteins were nebulized on copper Formar carbon-coated grids (Ted Pella Inc., Redding, CA), dried at room temperature for 30 min, and stained with phosphotungstic acid (PTA). Images were recorded at a calibrated magnification of 30,000 × or 60,000 × using an electron microscope (FEI Tecnai G2 Spirit BioTwin, Hillsboro, OR).

Statistical analysis

We used a t-test of independent samples with equal or unequal variances for statistical analysis. To determine the significance of observed differences in the OD values of prevaccination and postvaccination sera, a serum reactivity index (SRI) for each sheep, defined as the ratio of postvaccination OD value to the prevaccination OD value, was determined.
antibodies in vaccinated sheep. Using sera from vaccinated sheep, an increase in immunoreactivity with Gne antigen was observed from day 7 pv to day 28 pv, which later plateaued until the end of the experiment (Fig. 3A). In contrast, immunoreactivity with N antigen was barely detectable, remaining at baseline levels throughout the experiment (Fig. 3A). To confirm the specific immunoreactivity of the N protein as a marker antigen in ELISA, a day-28 MP12 post-infection antiserum (positive control) was tested alongside sera obtained from sheep vaccinated with the glycoprotein-based vaccine (Fig. 3B). Reactivity with these sera remained at baseline negative levels at all the time points, whereas the MP12 control serum showed strong reactivity as indicated by a high OD value ($p < 0.05$).

Recombinant RVFV glycoproteins elicit a neutralizing antibody response

To examine a vaccine-induced neutralizing antibody response, a PRNT assay was performed using an attenuated RVFV virus strain, MP12. A representative sample of a negative and positive PRNT80 assay using prebled and day-28 pv sera (sheep #170), respectively, is shown (Fig. 4A, B). In this study, a serum neutralizing antibody titer of 40 or greater is considered potentially protective (Pittman et al. 2000, Papin et al. 2011). Five of the six vaccinated sheep showed protective neutralizing titers at day 14 pv in response to the primary vaccination, with antibody titers ranging from 40 to 160; one sheep, #9, showed a protective neutralizing titer of 40 as early as day 7 pv (Table 1). Protective levels of virus neutralizing titers were maintained in all the sheep until day 21 pv, with three of the five sheep (#170, #179, #36) showing a titer increase. A second vaccine dose administered day 21 pv significantly boosted response in all six sheep above the 1280 titer at day 28 pv (Fig. 4C) (Table 1). Neutralizing antibody titers remained high in all sheep until day 49 pv (Fig. 4C), and all animals were positive for RVFV neutralizing antibodies when tested at day 328 pv with titers ranging from 80 to 160 (Table 1).

Electron microscopy

The purpose of TEM was to rule out the formation of VLPs following mixing of the recombinant glycoproteins Gne and Gc into the vaccine formulation. Images revealed by TEM showed clumps of protein aggregates that showed no resemblance to VLPs (Fig. 5).
Discussion

The impact of RVF outbreaks in Africa and the Arabian Peninsula, and the potential for viral spread to nonendemic areas, makes the development of safe and efficacious vaccines urgent. RVFV is a suitable candidate for a one health–focused approach to prevent both livestock and human disease through animal vaccinations (Oreshkova et al. 2013). However, there are currently no fully licensed or commercial vaccines for human or livestock use outside endemic areas,

![Diagram 3](image3.png)

**FIG. 3.** Differentiating infected from vaccinated animals (DIVA) analysis of the subunit shows indirect immunoglobulin G (IgG) enzyme-linked immunosorbent assay (ELISA) of Rift Valley fever virus (RVFV) anti-amino-terminus glycoprotein (Gn) and anti-nucleoprotein (N) antibody response in the vaccinated sheep. (A) Reactivity of sera with amino-terminus glycoprotein ectodomain (Gne) antigen indicates a time-dependent increase in antibody response, whereas in N-ELISA, reactivity remains at baseline prevaccination levels at all time points for all sera obtained from the vaccinated sheep. (B) Comparison of the reactivity of sera obtained from sheep vaccinated with the glycoprotein-based subunit vaccine to sera obtained from RVFV MP12-infected sheep, the positive control (PC) serum. The N antigen was positively reactive with only the PC serum indicated by high mean optical density (OD) value; day 0 to day 49 sera were obtained from sheep #169. The cutoff value for individual sheep in Gne-ELISA: #163 = 0.354; #169 = 0.167; #170 = 0.507; #179 = 0.365; #36 = 0.252; #9 = 0.668. The cutoff value in N-ELISA for individual sheep tested: #169N = 0.288; #163N = 0.237; #170N = 0.212; #9N = 0.407; #179N = 0.188; #36N = 0.239. N denotes recombinant N antigen used in ELISA. A cutoff value was determined for each sheep as described in Materials and Methods.

![Diagram 4](image4.png)

**FIG. 4.** An illustration of a plaque reduction neutralization test (PRNT<sub>80</sub>) shows negative (A) and positive (B) Rift Valley fever virus (RVFV) neutralization test results. Protective levels of RVFV neutralizing antibody titers (≥40) are detectable in the animals within 2 weeks postvaccination. A marked increase in RVFV neutralizing antibody titer is detected in all animals following administration of the second vaccine dose (C). PC, positive control serum (1:40 dilution); VC, virus control (50 pfu).
despite numerous potential vaccine candidates. Essential attributes for a RVFV vaccine include safety and high immunogenicity, DIVA compatibility, and the ability to induce a rapid onset of protective response with single vaccination, at most within 2 weeks of administration in susceptible host species.

Herein, the immunogenicity of a recombinant baculovirus-expressed RVFV Gne and Gc glycoprotein-based vaccine candidate in a natural host species, the sheep, is reported. Gne and Gc are presented as glycosylated proteins on the surface of RVF virions and have been shown to carry epitopes that elicit neutralizing antibodies, the only established correlate of protective immunity against virus infection (Besselaar, et al. 1991, Besselaar and Blackburn 1992). They are also used by the virus for attachment to target cells (Schmaljohn and Nichol 2006). Thus, the surface glycoproteins represent ideal targets for vaccine development, and we hypothesized that antibodies targeting epitopes on both structural glycoproteins in a subunit vaccine would generate a strong virus neutralizing effect. To produce vaccine immunogens, Gne and Gc expression constructs were designed to include a signal peptide at their amino terminus containing a signal peptidase cleavage site that ensures processing through translocation into the ER and the cellular glycosylation pathway and as well enhancing protein expression (Faburay et al. 2013).

Subsequently, sheep were immunized with the purified baculovirus-expressed Gne and Gc proteins with the adjuvant montanide ISA25 resulting in induction of virus neutralizing antibody response, followed by a strong response after the booster dose, in all vaccinated animals. However, due to lack of approval at the time of this study to perform challenge experiments with wild-type RVFV, it was not possible to conduct further studies to test the efficacy of the vaccine. The vaccine induced potentially protective (i.e., 1:40) virus neutralizing titers with single vaccination in five of the six animals within 2 weeks pv (Table 1). These results compared favorably with the outcome of recently reported vaccinations using vaccines based on RVFV glycoproteins, such as GnGc VLPs and Gne (Naslund et al. 2009, de Boer et al. 2010, Mandell et al. 2010a, Kortekaas et al. 2012, Oreshkova et al. 2013), as well as a Newcastle disease virus-vectored vaccine (NDFL-GnGc) (Kortekaas et al. 2010a, Kortekaas et al. 2010b) and virus replicon particles (Dodd et al. 2012, Oreshkova et al. 2013), some of which have also been reported to elicit neutralizing antibodies with single vaccination in sheep (Kortekaas et al. 2010a, Kortekaas, et al. 2012, Oreshkova, et al. 2013).

It is significant that the GnGc-based recombinant protein vaccine candidate tested in this study induced potentially protective neutralizing antibody titers with single vaccination.

### Table 1. Reciprocal PRNT80 Titers in Sheep in Response to Vaccination with RVFV Recombinant GnGc Glycoprotein Subunit Vaccine

<table>
<thead>
<tr>
<th>Sheep no.</th>
<th>Days: 0 prevac</th>
<th>7 pv</th>
<th>14 pv</th>
<th>21 pv</th>
<th>28 pv</th>
<th>35 pv</th>
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<th>328 pv</th>
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<td>77</td>
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RVFV, Rift Valley fever virus; Gne, amino-terminus glycoprotein; GC, carboxyl-terminus glycoprotein; PRNT80, plaque reduction neutralization 80% titers; prevac, prevaccination; pv, postvaccination; nd, not determinable.

FIG. 5. Electromicrograph of purified recombinant Rift Valley fever virus (RVFV) proteins used in the vaccine formulation. Recombinant amino-terminus glycoprotein ectodomain (Gne), carboxyl-terminus glycoprotein (Gc), and mixed GnGc show clumps of protein aggregates that are structurally distinct from RVF VLPs, shown as control (de Boer et al. 2010). Gne and Gc are mixed in equal amounts and incubated at room temperature to allow possible assembly of proteins into VLPs. This figure illustrates the nonformation of VLPs.
in 80% (5/6) of sheep within 2 weeks of vaccination and 100% (6/6) sheep at three weeks post-vaccination (pv) (Table 1). The robust neutralizing antibody response elicited by the RVFV vaccine candidate could be attributed to the concurrent use of Gne and Gc proteins as vaccine immunogens. Gc is known to contain virus-neutralizing epitopes, whereas inclusion of Gc in the vaccine is suggested to provide an additional target for neutralizing antibodies (Besselaar et al. 1991, Besselaar and Blackburn 1992). Neutralizing antibody titers increased sharply in all animals following the booster, and this high anamnestic response was maintained in all animals for more than 3 weeks, i.e., up to day 49 pv. Remarkably, neutralizing antibody titers, ranging from 80 to 160, above the potentially protective threshold, were detected in all sheep at day 328 pv (Table 1). This is against the backdrop that in the United States, the withdrawal time for the adjuvant used in the current vaccine is 90 days. An early-onset vaccine-induced immunoglobulin G (IgG) antibody response to Gne occurred in half of the sheep within 7 days pv followed by seroconversion in 100% of the animals at 2 weeks pv for both Gne and Gc (Fig. 2A–D). It is noteworthy that high ELISA background was detected in some sheep. Although this is speculated to be due to a host of unknown factors, including the physiological status of individual animals, it nonetheless indicates the need for further development of the assay. Taken together, these results support the conclusion that the RVFV recombinant GneGc glycoprotein-based vaccine candidate is highly immunogenic, eliciting a strong immune response in sheep, the natural and most susceptible species to RVFV.

Differentiating infected from vaccinated animals during RVF disease outbreaks is of fundamental epidemiological importance. Therefore, DIVA compatibility of a vaccine with accompanying diagnostic tests represents an important factor for consideration when designing vaccines, especially for use in countries or regions nonendemic for RVFV. Using the RVFV glycoproteins and the nucleocapsid protein as diagnostic antigens, it was possible to distinguish vaccine-induced antibody responses (Fig. 3A) from RVFV MP12 infection in sheep (Fig. 3B). The increase in international trade in livestock coupled with the potential for RVFV outbreaks in nonendemic areas provides strong incentives for the development of DIVA vaccines. The absence of the nucleocapsid protein in the vaccine offers the possibility of developing a DIVA vaccine with a companion diagnostic assay using the recombinant N and Gne/Gc ELISA.

The N protein represents a suitable diagnostic antigen because it is the most abundant viral protein and is highly immunogenic, inducing antibodies within the first days after infection (Swanepoel et al. 1986, Paweska et al. 2008, Faburay et al. 2013). Furthermore, the recombinant GmGc glycoprotein subunit vaccine candidate elicited strong neutralizing and IgG antibody responses in the natural host, which can be easily detected by ELISA assays. Analysis of the structural morphology of the vaccine immunogens by electron microscopy confirmed that the proteins upon reconstitution formed into clumps or aggregates that were very much distinct from VLPs (Fig. 5). To date, RVFV VLP assembly has been reported to occur only by simultaneous production of Gn and Gc in both mammalian (Mandell et al. 2010a,b) and insect cells (Liu et al. 2008, de Boer et al. 2010) and in all cases involved the co-expression of nonhistidine-tagged proteins. These results are anticipated, for unlike with naked viruses, where VLPs are reassembled proteins, formation of an enveloped virus such as RVFV requires live cells.

RVFV represents a threat to human and animal health, and there is no commercially available vaccine or effective antiviral therapeutic agent for human use. Efforts to produce live attenuated vaccines (Bouloy and Flick 2009) are tempered by safety concerns; however, such safety concerns are comparatively lower for recombinant protein–based vaccines. However, unlike live-attenuated vaccines, recombinant protein–based vaccines generally require boosters to elicit long-term protective immune response. Therefore, future research will be directed at further improving the duration and onset of protective neutralizing antibody responses. Additionally, the fact that RVFV has low genetic diversity and consists of a single serotype suggests that the recombinant Gne and Gc glycoprotein vaccine would likely confer protection against all strains of the virus. Further evaluation of the efficacy of the vaccine in livestock and nonhuman primates is the next step toward developing a safe and efficacious vaccine for livestock and human use.

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Author Disclosure Statement

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The Bunyaviridae


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Address correspondence to:
Bonto Faburay
Department of Diagnostic Medicine/Pathobiology
College of Veterinary Medicine
Kansas State University
1800 Denison Avenue
Mosier Hall K218
Manhattan, KS 66506
E-mail: bfaburay@vet.k-state.edu

and

Juergen A. Richt
Department of Diagnostic Medicine/Pathobiology
College of Veterinary Medicine
Kansas State University
1800 Denison Avenue
Mosier Hall K224
Manhattan, KS 66506
E-mail: jricht@vet.k-state.edu