Chimeric rabies glycoprotein with a transmembrane domain and cytoplasmic tail from Newcastle disease virus fusion protein incorporates into the Newcastle disease virion at reduced levels

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Rabies remains an important worldwide health problem. Newcastle disease virus (NDV) was developed as a vaccine vector in animals by using a reverse genetics approach. Previously, our group generated a recombinant NDV (LaSota strain) expressing the complete rabies virus G protein (RVG), named rL-RVG. In this study, we constructed the variant rL-RVGTM, which expresses a chimeric rabies virus G protein (RVGTM) containing the ectodomain of RVG and the transmembrane domain (TM) and a cytoplasmic tail (CT) from the NDV fusion glycoprotein to study the function of RVG’s TM and CT. The RVGTM did not detectably incorporate into NDV virions, though it was abundantly expressed at the surface of infected BHK-21 cells. Both rL-RVG and rL-RVGTM induced similar levels of NDV virus-neutralizing antibody (VNA) after initial and secondary vaccination in mice, whereas rabies VNA induction by rL-RVGTM was markedly lower than that induced by rL-RVG. Though rL-RVG could spread from cell to cell like that in rabies virus, rL-RVGTM lost this ability and spread in a manner similar to the parental NDV. Our data suggest that the TM and CT of RVG are essential for its incorporation into NDV virions and for spreading of the recombinant virus from the initially infected cells to surrounding cells.

Keywords: Newcastle disease virus, antibody response, chimeric rabies glycoprotein, rabies virus, viral vector

Introduction

Rabies is a zoonotic disease that causes severe nerve damage in humans and mammals [8]. Rabies virus (RV) is an enveloped virus from the genus *Lyssavirus* of the *Rhabdoviridae* family. The RV virion is composed of five structural proteins [9,11,15,16,24,25], and its G protein binds to receptors at the cell surface and undergoes a pH-dependent conformational change to mediate membrane fusion and endocytosis, allowing RV to enter cells [1]. Rabies virus G protein (RVG) is also the main protein responsible for inducing and reacting with virus-neutralizing antibody (VNA), which is useful as a marker of productive immunization [35].

Newcastle disease virus (NDV) causes severe disease symptoms in all species of birds [7]. Some NDV strains were developed as live-virus vectors by using a reverse genetics approach [3,17,18,20,27,29,32]. In order to find an effective, safe, and affordable RV vaccine, our group has generated rL-RVG, which is an NDV vector (avirulent NDV LaSota strain) expressing RVG [19]. Our results showed that RVG is efficiently incorporated into the NDV virion, and rL-RVG can spread on BHK-21 cells in a manner similar to that of RV but different from NDV’s manner of spreading. Safe in cats and dogs, L-RVG can induce a substantial and protective rabies virus neutralization antibody in intramuscular-vaccinated animals [19]. To determine the function of the transmembrane domain (TM) and cytoplasmic tail (CT) of RVG, we generated a recombinant chimeric RVG-NDV variant, in which the ectodomain of RVG has been fused with the TM and CT of the NDV F protein. In the present study, we evaluated the influence of the RVG TM and CT on immune responses and incorporation into NDV.
Materials and Methods

Cells and viruses

BHK-21 cells were grown in Dulbecco’s modified Eagle medium containing 5% fetal bovine serum. Our group generated the parental NDV vector of the LaSota vaccine strain (rL) and rL-RVG, which is a recombinant NDV strain expressing RVG [19]. A modified vaccinia virus expressing T7 RNA polymerase [36] was grown and titered in primary chicken embryo fibroblasts. Recombinant RV and NDV expressing enhanced green fluorescent protein were generated by our laboratory group.

Construction of recombinant NDV expressing chimeric rabies virus G protein (RVGTM)

A RVGTM gene containing sequences encoding the TM domain and CT of the NDV F protein was constructed by gene synthesis. We introduced the modified G gene into the vector pLa through a unique Pmel site and rescued the virus as described previously [17]. The rescued recombinant rL-RVGTM was confirmed by assessing hemagglutination activity, and the nucleotide sequences of the incorporated RVGTM gene in the rescued virus were confirmed by performing reverse transcription-polymerase chain reaction (RT-PCR) and nucleotide sequencing.

Western blotting

BHK-21 cells were incubated with rL, rL-RVG, or rL-RVGTM at an multiplicity of infection (MOI) of 5.0, and cell sediments were prepared for analysis at 60 h post-infection. To compare the incorporation of RVG and RVGTM into NDV virions, allantoic fluid was harvested from specific pathogen-free (SPF) chicken eggs at 72 h post-infection, and virus particles were purified as described previously [19]. We determined the amounts of cell-associated protein and purified virion protein by assaying the thickness of β-actin and NDV bands on western blots. Equal amounts of protein were analyzed by western blotting with SDS-12% PAGE. After being incubated with chicken serum against NDV or rabbit serum against RV, horseradish peroxidase-conjugated rabbit anti-chicken IgG (A9046; Sigma) or goat anti-rabbit IgG (ZB-2301; ZSGB-BIO) was used to detect chicken or rabbit serum binding.

Pathogenicity of recombinant virus

To assess the pathogenicity of rL-RVGTM, the mean death time (MDT), intracerebral pathogenicity index (ICPI), and intravenous pathogenicity index (IVPI) of White Leghorn chickens were determined [17]. Values for rL-RVG and rL virus were determined simultaneously for comparison purposes. Two groups of 12 mice (6-week-old mice) were inoculated intramuscularly (i.m.) in the gastrocnemius muscle with 5 × 10⁷ times the 50% egg infective dose (EID₅₀) of rL-RVG or rL-RVGTM. Mouse weight change was determined to assess viral pathogenicity in mammals.

Care of animals and animal experimentation in our laboratory were performed according to approved animal ethics guidelines and protocols. All animal studies were approved by the Animal Ethics Committee of Harbin Veterinary Research Institute of the Chinese Academy of Agricultural Sciences (SYXK (H) 2006-032).

Growth kinetics of rL-RVGTM in cells or eggs

To determine whether RVGTM expression alters the growth properties of rL, 10⁴.⁵ EID₅₀ of rL, rL-RVG, or rL-RVGTM were inoculated into SPF chicken eggs. Allantoic fluid from five eggs was harvested every 12 h (12–108 h) post-infection, and the TCID₅₀ of each virus was determined by assessing immunofluorescence.

The three viruses at MOIs of 0.1 were inoculated into BHK cells on 24-well plates, and the supernatant harvested at 24, 36, 48, 60, 72, and 84 h post-infection. The viruses were titrated by determining both their immunofluorescence and EID₅₀. Cells were evaluated by examining immunofluorescence in order to investigate the manner of spread of rL-RVGTM in cells after the supernatant was collected. Part of the collected supernatant was used to determine, via immunofluorescence, whether the incorporation of RVGTM alters the trypsin-dependent infectivity of the NDV vector in mammalian cells.

ELISA to detect the incorporation of RVGTM

Purified viruses were assessed by enzyme-linked immunosorbet assay (ELISA) with dog serum against RV or chicken antibody against NDV along with horseradish peroxidase-conjugated rabbit anti-dog IgG (A6792; Sigma) or rabbit anti-chicken IgG (A9046; Sigma). We determined the protein concentrations to standardize the amount of loading (10 µg, 5 µg, 2.5 µg, 1.25 µg, 0.625 µg, 0.3125 µg, 0.15625 µg, and 0.078125 µg) of duplicate samples of the purified virions on the ELISA plate. Data was collected by using an enzyme standard instrument (Model 650; Bio-Rad Laboratories, USA).

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**Immunization comparison**

Two groups of 12 mice were inoculated i.m. with rL-RVG or rL-RVGTM as described in section “Pathogenicity of recombinant virus”. After 4 weeks, the mice received second doses. For serological assessment, blood was obtained from the retro-orbital sinus under isoflurane inhalation anesthesia at 3 and 4 weeks after the first dose and at additional times (1, 2, 3, 5, 7, 9, 13, 17, and 21 weeks) after the second dose. With a World Health Organization standard as a reference, titers of RV VNA were expressed in international units per milliliter of serum (IU/mL) [19].

**Results**

**Generation of recombinant NDVs expressing the RVGTM gene**

To determine the effects of replacing the TM and CT regions of RVG with corresponding domains of the NDV F protein, we prepared a synthetic RVGTM and inserted it into the unique Pmel site of pLa to produce the resulting plasmid pRL-RVGTM (Fig. 1). The rescued recombinant rL-RVGTM was confirmed by its hemagglutination activity results, and the nucleotide sequences of the incorporated RVGTM gene in the rescued virus were confirmed by the RT-PCR and nucleotide sequencing.

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**Fig. 1.** Construction of a viral vector for expressing the chimeric NDV/RV protein RVGTM. (A) NDV has two transmembrane glycoproteins; HN protein and F protein. The NDV F protein and rabies virus G protein (RVG) are type I transmembrane proteins. (B) Schematic of the generation of recombinant NDV rL bearing the RVGTM gene. RVGTM is a chimera of the RVG ectodomain fused with the sequence for the transmembrane domain and cytoplasmic tail of the NDV protein. The RVGTM sequence was inserted into the vector PBRN-rL at the Pmel site to produce prL-RVGTM. NDV, Newcastle disease virus; RV, rabies virus; RVGTM, chimeric rabies virus G protein.

**Fig. 2.** The rL-RVGTM expresses RV and NDV components. (A) Detection of RVGTM expression by immunofluorescence. BHK-21 cells were infected with rL, rL-RVG, or rL-RVGTM at a multiplicity of infection of 0.1. At 24 h post-infection, the cells were fixed and stained with chicken anti-NDV and dog anti-RV sera followed by fluorescein isothiocyanate (FITC)-conjugated rabbit anti-dog and tetramethyl rhodamine isocyanate (TRITC)-labeled goat anti-chicken antibodies. The cells were permeabilized with 0.02% Triton X-100 and cell nuclei were stained with DAPI. (B) Western blot analysis of recombinant NDV expressing RVGTM. Lysates of BHK-21 cells infected with rL, rL-RVG, or rL-RVGTM were separated by 12% SDS-PAGE and incubated with dog serum against RV, chicken serum against NDV, or anti-β-actin monoclonal antibody as a loading control. After incubation with peroxidase-conjugated secondary antibody, the proteins were visualized with 3,3-diaminobenzidine reagent. The locations of marker proteins are indicated on the left, and the antiserum or antibody used is indicated on the right. RVGTM, chimeric rabies virus G protein; RV, rabies virus; NDV, Newcastle disease virus; RVG, rabies virus G protein; MK, protein molecular marker.
Verification of rL-RVGTM expression

To verify the expression of RVGM, we performed laser confocal microscopy. The rL-RVG and rL-RVGTM were stained by both mouse monoclonal IgG1 against NDV HN antibody and dog serum against RV, while the rL was only stained by mouse monoclonal IgG1 against NDV HN (panel A in Fig. 2). Western blot analysis confirmed the expression levels of RVG and RVGM in BHK-21 cells were similar (panel B in Fig. 2).

Viral growth kinetics

To further assess the ability of RVGM to produce virus, we measured viral titers after infection of SPF chicken eggs. The rL, rL-RVG, and rL-RVGTM grew to similar levels in the eggs (panel A in Fig. 3). The rL-RVGTM reached a peak titer of 8.375 log TCID₅₀/mL at 72 h post-infection, which is similar to the titer level of rL and slightly higher than that of rL-RVG. These results confirm that rL-RVGTM produces virus at levels equivalent to or greater than those produced by rL and rL-RVG.

Comparison of the virulence of different recombinant NDVs

Determination of MDT, ICPI, and IVPI (panel B in Fig. 3) levels was performed to obtain measures of virulence. The MDT of rL-RVGTM was more than 120 h while its ICPI and IVPI values were 0, indicating that expression of the RVGM gene does not alter the pathogenicity of the NDV vector in poultry.

A previous report indicated no differences between the body weight of rL and rL-RVG infected mice. To determine the effects of RVGM on pathogenicity in mice, we inoculated mice with rL-RVG and rL-RVGTM; all mice survived and body weight changes between two groups were similar (panel C in Fig. 3). Therefore, RVGM expression does not alter the pathogenicity of the recombinant NDV vector in mice.

RVGM incorporates into the NDV virion at reduced levels

To determine whether RVGM incorporates into the NDV virion, we collected viral particles 72 h post-infection of SPF chicken eggs and assessed the contents by western blot analysis. RVG was incorporated more efficiently than RVGM into the NDV virions, and RVGM was undetectable by western blotting (panel A in Fig. 4). These results indicate that RVGM incorporates into the NDV virion at reduced levels.

To verify these results, we performed ELISA testing with anti-RV and anti-NDV antibodies. ELISA performed with dog anti-RV demonstrated that rL-RVG levels were notably higher than rL-RVGTM levels at low dilutions, but the two levels tended to be similar at high dilutions. In contrast, ELISA with chicken anti-NDV yielded similar results at every dilution for both rL-RVG and rL-RVGTM, suggesting that the rL-RVG and RVGM incorporate into the NDV virion at reduced levels.
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Fig. 4. RVGTM incorporates poorly into the NDV virion. (A) To determine whether the RVGTM incorporates into the NDV virion, the recombinant rL-RVG and rL-RVGTM viruses were purified through sucrose gradients. Equivalent amounts of rL, rL-RVG, and rL-RVGTM virions were analyzed by SDS-PAGE. (B and C) Top: ELISA analysis of the incorporation of RVGTM and RVG into the NDV virion was performed by testing for G protein. Bottom: ELISA analysis of NDV demonstrates that the amount of the two recombinant viruses is equivalent despite differences in G protein incorporation. RVGTM, chimeric rabies virus G protein; NDV, Newcastle disease virus; RVG, rabies virus G protein; MK, protein molecular marker; NP, nucleo protein; M, matrix protein; GP, Glycoprotein.

rL-RVGTM virion amounts were similar (panel C in Fig. 4). These results confirm that native G protein in rL-RVG incorporates more effectively than the G protein in rL-RVGTM.

Chimeric rL-RVGTM virus has altered ability to spread

To assess the ability of the rL-RVGTM virus to spread, we performed immunofluorescence assays. The percentages of cells infected by rL and rL-RVGTM were similar and did not change significantly during the 24 to 96 h post-infection time course in the absence of tosylsulfonyl phenylalanlyl chloromethyl ketone (TPCK) trypsin. However, with elapsing
Fig. 6. rL-RVGTM elicits reduced levels of RV VNA. (A) Induction of serum antibodies specific to NDV and RV were assessed following intramuscularly (i.m.) immunization of mice with equivalent levels of rL-RVG and rL-RVGTM. Twelve mice from each group were given the first immunization at week 0 and a second dose 4 weeks later. Serum samples were collected at the third and fourth weeks after the first immunization and at appropriate times after second dose. Neutralization test of NDV VNA. Values represent the dilution ratio of NDV specific antibodies that can protect half of the cells from being infected by NDV. (B) Neutralization test of RV VNA. Values are presented as IU/mL. *p < 0.05, **p < 0.01. RVGTM, chimeric rabies virus G protein; RV, rabies virus; VN A, virus-neutralizing antibody; NDA, Newcastle disease virus; RVG, rabies virus G protein.

time, rL-RVG fluorescence spread notably in the absence of TPCK trypsin (panel A in Fig. 5). These results are in contrast to the results obtained in the presence of TPCK trypsin, in which both RVG and RVGTM spread over time, which suggests that the reduced infection level of rL-RVGTM is due to a reduced ability to spread rather than due to effects on the number of viral particles. These findings were verified by analyzing the EID₅₀ of supernatant collected from the three progeny viruses; the three viruses grew to similar levels in eggs, but the titer of rL-RVG was a little higher than those of rL and rL-RVGTM (panel B in Fig. 5).

**Induction of anti-NDV and anti-RVG serum antibody responses in mice immunized with recombinant NDVs**

To determine whether rL-RVG and rL-RVGTM differ in their immunogenic properties, we immunized mice twice (at weeks 0 and 4) and performed neutralization tests over a 21-week period using serum collected from the mice. The NDV VNA levels of the rL-RVG and rL-RVGTM group both increased sharply and then gradually declined, with similar levels in both viruses after both the first and second immunization and at all times tested (panel A in Fig. 6). In contrast, after the first immunization, the mean titer of RV VNA for rL-RVG was 0.77, while the mean titer of RV VNA for rL-RVGTM was only 0.21 (p = 0.01). After the second immunization, the RV VNA levels were induced to a much greater extent and remained high for a longer period for rL-RVG than for rL-RVGTM (panel B in Fig. 6). Additionally, more than 83.3% of the mice in the rL-RVG group were protected, while only 33.3% of the mice in the rL-RVGTM group were protected after a single dose (data not shown). These results suggest that substitution of the TM and CT regions of RVG greatly diminishes its ability to induce a VNA response.

**Discussion**

Previously, our group generated a recombinant NDV expressing RVG and evaluated its potential as a novel vectored vaccine against rabies in animals [19]. In this study, we replaced the TM domain and CT of RVG with that of the NDV F protein by performing gene synthesis. We generated a recombinant NDV expressing a chimeric RV glycoprotein and tested its characteristics. The expression of RVGTM was verified by undertaking laser confocal microscopy and western blotting, and the results showed that the expression levels of native RVG and RVGTM were similar in BHK-21 cells. Furthermore, MDT, IVPI, and ICPI measurements from mouse experiments indicated that the expression of RVGTM did not increase the virulence of the NDV vector; rL-RVGTM was shown still to comprise a lentogenic strain that is safe for poultry and mice.

Foreign glycoproteins can be incorporated into recombinant NDV virions during their envelopment from the host cell plasma membrane [2,12,13,26]. Western blotting and ELISA tests with equal amounts of purified rL, rL-RVG, and rL-RVGTM viral particles indicated that the incorporation of RVGTM into NDV virions decreased compared to the incorporation of RVG. Mouse experiments showed that the NDV VNA levels of the two groups were similar after the first and second immunization, whereas the level of RV VNA induced by rL-RVGTM was obviously lower than that of rL-RVG. This result is consistent with the result of a study of recombinant NDVs expressing native gD protein and a
chimeric gD from bovine herpes virus-1 [22], but is contrary to
the result of a study of recombinant NDV expressing native HA
protein and a chimeric HA protein of avian H7 influenza [3,29].
Therefore, whether chimeric proteins are incorporated into
virions may depend on specific structural criteria of different
viral glycoproteins.

RVG is a typical type I glycoprotein that is anchored in the
evelope, forming spikes as a trimer, which is the functional
unit of its assembly into virions, and mediating virus entry. The
basis for the highly efficient incorporation of the RVG into the
NDV virion is still not fully described, and the reason for the
reduced incorporation when the TM and CT of RVG are
replaced is unknown. According to a previous study [22], one
possibility is that some amino acid sequence characteristics of
the TM domain or CT of the native G protein contribute to their
inclusion in particles. TM and CT replacement of G proteins
results in slower transportation from the endoplasmic reticulum
to the Golgi complex, and the rate at which G protein is
transported from the endoplasmic reticulum to the Golgi
complex influences the rate of accumulation on the plasma
membrane [14,34]. Thus, it is possible that native RVG might
accumulate in higher molar amounts at the cell surface leading
to efficient incorporation.

Progeny of RV can spread from the initial infected cells to
contiguous or noncontiguous cells surrounding the interstitial
space to form large plaques [10,19]. However, rL is a
low-pathogenicity NDV strain that can infect individual cells
but cannot spread to adjacent BHK-21 cells without TPCK
trypsin (1 mg/mL). The rL-RVG acquired the ability to spread
from cell to cell in BHK-21 cells [19], whereas rL-RVGTM lost
this ability, as assessed by immunofluorescence. This change
may be explained by several considerations. The vesicular
stomatitis virus (VSV) G protein lost its fusion properties when
its TM segment was replaced with a glycoprophatidylinositol
anchor [28], and a similar phenomenon could explain the lost
spreading ability of rL-RVGTM. Mutations in the glycine
residues of the TM segment of VSV G protein block fusion at
the hemifusion stage [6]. These results suggest that the TM
segment of the G protein has a major role in the fusion process
of Rhabdoviridae. Furthermore, the 13 membrane-proximal
amino acids close to the TM are reported to influence cell-to-cell
spreading [21,23]. Therefore, the replacement of the TM and
CT may influence the interaction between the 13 membrane-proximal amino acids of RVG and the NDV F
protein. Refolding of the C-terminal portion of the molecule
also facilitates the switch between the pre- and post-conformations
of the G protein [33], a switch that is essential for the fusion
process, and the loss of the native C-terminal portion may
prevent the structural transition of RVGTM. An additional
consideration is that the F protein is a class I fusion protein,
while RV G protein is class III fusion protein, and their
C-terminal segments have different roles in the pre- and
post-fusion states [31]. Thus the TM and CT from the F protein
may not be able to compensate for the roles of G protein’s TM
and CT. The HA protein of avian H7 influenza and the NDV F
protein are both class I fusion proteins. Although the trimer
structure of NDV F glycoprotein is different from that of HA, in
which the orientation of the heptad repeat A is opposite to that
of the central, coiled trimer observed in HA [4,5], they undergo
a similar conformational change during the fusion process.
Amino acid alignment has revealed that all rhabdovirus
glycoproteins can be said to share the same fold as VSV; on that
basis, the RV G is a class III fusion protein [1]. Although the
recognition of G protein domains is similar to that of class II
fusion proteins, G protein’s refolding process from pre-
to post-fusion is similar to that of class I proteins, which form
6-helix bundles by first forming the trimeric central core of the
post-fusion conformation. However, in the pre-fusion state, G
proteins show a unique property that differs from those of both
class I and class II fusion proteins. Their fusion loops are not
buried in an oligomeric interface, but instead they point toward
the viral membrane [31]. These differences between F, HA, and
G proteins may lead to different results. Finally, there are
estimated to be at least 15 G protein spikes in the fusion
complex that is involved in achievement of the fusion reaction
[30,31], and the reduced incorporation of RVGTM may
contribute the reduced fusion of RVGTM. We speculate that the
chimeric TM and CT results in the loss of the fusion properties
of RVGTM.

In summary, we have demonstrated that RVG incorporates
more efficiently into recombinant virions than RVGTM does.
The replacement of the TM and CT of RVG with that of the
NDV F protein may change the conformation of RVGTM and
influence the effective antigen epitopes of the G protein. The
decreased incorporation of RVGTM into NDV virions and the
loss of fusion function of RVGTM further contribute to the
reduction of the RV VNA levels induced by rL-RVGTM
directly. Our results demonstrate that the C-terminal portion
of RVG has an important role in the spreading of the virus from the
initially infected cells to surrounding cells.

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Conflict of Interest

The authors declare no conflicts of interest.
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