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Citation for published version:

Digital Object Identifier (DOI):
10.1016/j.vaccine.2009.09.024

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
Vaccine

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A robust lentiviral pseudotype neutralisation assay for in-field serosurveillance of rabies and lyssaviruses in Africa

Edward Wrighta,†, Suzanne McNabbb,c, Trudy Goddardd, Daniel L. Hortone,d,e, Tiziana Lemboe,g, Louis H. Nel1, Robin A. Weissa, Sarah Cleavelandc,e,g, Anthony R. Fooksd,∗∗

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1. Introduction

Rabies is spreading at an alarming rate in some regions of the developing world [1–3]. Collaborative efforts among human and animal health care professionals are required to monitor this situation and allow a timely and proportioned therapeutic response, limiting unnecessary use of valuable vaccines. The combination of rabies awareness campaigns, improved vaccine coverage and disease surveillance has already resulted in the successful elimination of rabies awareness campaigns, improved vaccine coverage and limiting unnecessary use of valuable vaccines. The combination of canine rabies in North America [4]. However, vaccines need to be efficacious enough to elicit a response that will confer protection and diagnostic techniques are needed to ensure that adequate levels of virus-neutralising antibodies (VNAbs) have been achieved in response to vaccination. The problem with current tests is that they are not easily used in endemic areas within developing countries because they need to be set up in containment laboratories since highly pathogenic zoonotic viruses are required to perform the assay, or such assays are prohibitively expensive. Those that are e.g., ELISA-based assays, do not have the sensitivity or specificity of the fluorescent antibody virus neutralisation (FAVN) assay that is widely used within Office international des épizooties (OIE) rabies reference laboratories. Therefore, in addition to the need for further implementation and development of improved anti-rabies biologicals, improved techniques to assess seroconversion are required before canine rabies is success-
fully eliminated in developing countries as it has been in North America.

Within the Lyssavirus genus, classical rabies viruses (genotype 1) are not the only pathogen to cause morbidity and mortality in mammalian populations. Clearly, infection by lyssaviruses of the other genotypes (2–7) can result in a clinical manifestation that is indistinguishable from rabies. The other genotypes are distributed geographically predominantly within African, European and Australian bat populations [5]. Recently, additional variants that are more divergent than genotypes 1–7 have been isolated suggesting further genotypes may yet exist [6–10]. Isolates representing genotypes 1, 2 and 4–7 have been identified in insectivorous, fruit bat species [5]. Mokola virus (MOKV, genotype 3) along with genotypes 1, 2 and 4–7 have been identified in insectivorous, fruit bat species [5].

Lagos bat virus (LBV, genotype 2) and Duvenhage virus (DUVV, genotype 4) comprises the African lyssaviruses. Interestingly, only a few clinical isolates representing these genotypes have been identified to date [11]. The handful of cases that have been reported has given us a limited understanding of the epidemiology and zoonotic threat that these genotypes pose in their respective hosts. Recently, surveillance programs and greater access to serosurveillance techniques have resulted in the discovery of a high seroprevalence against LBV in East and West African megachiropterans [12,13] and a common presence of LBV in South African bats collected for routine surveillance [11].

These reports and others [2,14] emphasise that the potential for increased incidence levels of rabies and related lyssavirus infections is a concern in Africa, mainly because of the lack of awareness of these infections in the population. However, there is also poor accessibility to vaccines and post-exposure treatments for those exposed to these viruses and there are difficulties with undertaking serosurveillance measures in many countries within Africa [15]. While the most important factor in reducing rabies prevalence is the implementation of vaccination campaigns, it was highlighted at the recent Southern and Eastern African Rabies Group meeting that poor infrastructure becomes a major barrier when attempting to control rabies in Africa [16]. These views are shared by the OIE and World Health Organization (WHO), that list the development of novel diagnostics as an urgent requirement [17,18].

Serological techniques that can be employed to study naturally occurring or vaccine-induced humoral responses to rabies virus infection include the FAVN assay [19], rapid fluorescent focus inhibition test (RFFIT) [20] and enzyme linked immunosorbant assay (ELISA) [21]. Variations of these assays have been described previously [22,23]. The routinely used FAVN assay and RFFIT are the current assays of choice with OIE/WHO reference laboratories but must be performed in BSL3/SAPO4 high containment facilities because live virus is handled as part of the assay. While a modified RFFIT that combines green fluorescent protein (GFP) with live recombinant virus can remove the need for expensive conjugates, work with recombinant virus still requires the use of high containment facilities [23]. With the recent addition to the above mentioned set of neutralisation assays of the ELISA-based method that uses plates coated with whole, inactivated virus, the need for live virus has been eliminated. Since both non-neutralising and neutralising antibodies are detected, the level of circulating, protective VNAb alone cannot be determined. There are also issues with low sensitivity when using the ELISA.

We recently described the use of surrogate viruses known as lentiviral pseudotypes as replacements for live or inactivated whole virus to accurately determine anti-rabies VNAb responses in vaccine recipients. The samples tested were taken from vaccinated humans, dogs and cats in the United Kingdom (UK) [24]. Here we report the results of the largest virus neutralisation study published to date using the surrogate lentiviral pseudotypes rather than the live native or recombinant rabies virus with field serum samples from Tanzanian dogs. We further increase the utility of our pseudotype neutralisation assay for laboratories undertaking vaccine trials and serosurveillance in resource-limited, rabies endemic countries by exploring the use of lacZ as a reporter gene and incorporating the glycoproteins of a further three lyssavirus
genotypes, in addition to genotype 1, which will allow improved serosurveillance for lyssaviruses other than classical rabies. This report describes a highly sensitive yet flexible platform that can be adapted to allow the evaluation of vaccine and antiviral drugs against highly pathogenic viruses without the need for high level containment facilities or expensive reagents and equipment.

2. Methods

2.1. Study area

Dogs enrolled in this study were selected from domestic animals living in four villages (Ngarawani, Runga’bure, Nyamburi and Bisarara) within 20 km of the Serengeti National Park perimeter in the Serengeti District, northwestern Tanzania (Fig. 1A). In each case the dog owner’s consent was sought before enrolling their dog in the study. This project was part of the annual vaccination campaign undertaken by the Viral Transmission Dynamics Project, which works to prevent the spread of diseases through animal populations in the Serengeti region.

2.2. Vaccination history

A detailed medical history of each dog enrolled in this study, which included any prior vaccinations, was taken at the first visit. Sixty-six and a half percent (n = 125) of the dogs had not been vaccinated against rabies prior to this study (referred to as “primary”), 27.2% (n = 52) had received at least one rabies vaccine previously (“booster”) and there was no vaccination history available or taken (“no record”; Table 1). The Nobivac® rabies virus vaccine (donated by Intervet Schering-Plough Animal Health) was administered to all dogs in a single dose inoculation given subcutaneously. It comprises an inactivated vaccine containing ≥2 IU rabies virus (Pasteur strain).

2.3. Serum samples

At the first visit, a blood sample was taken from each dog before the rabies virus vaccine was administered. The second visit was 20–21 days post-vaccination, the optimum time to detect an immune response stimulated by the vaccine. The overall study protocol is shown in Fig. 1B.

Blood was drawn from the cephalic or jugular vein, stored on ice, and processed at the end of each day (centrifugation at 3000 rpm for 10 min). Serum was prepared, inactivated at 57°C for 30 min and then frozen for transport. In total, 321 samples were taken from the enrolled dogs, blinded and sent to University College London (UCL) and Veterinary Laboratories Agency (VLA) for testing. The OIE standard reference dog serum diluted to 0.5 international units/ml (IU/ml) with PBS was used as a positive control.

Table 1

<table>
<thead>
<tr>
<th>Village</th>
<th>Mean age (years)</th>
<th>Gender (% female)</th>
<th>Number of sera collected</th>
<th>Vaccination history at 1st visit (% of total)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>First visit</td>
<td>Second visit</td>
</tr>
<tr>
<td>Ngarawani</td>
<td>2.1</td>
<td>47.9</td>
<td>48</td>
<td>37</td>
</tr>
<tr>
<td>Runga’bure</td>
<td>2.65</td>
<td>48</td>
<td>50</td>
<td>38</td>
</tr>
<tr>
<td>Nyamburi</td>
<td>1.4</td>
<td>54.2</td>
<td>48</td>
<td>26</td>
</tr>
<tr>
<td>Bisarara</td>
<td>1.56</td>
<td>55.6</td>
<td>45</td>
<td>29</td>
</tr>
<tr>
<td>TOTAL</td>
<td>1.94</td>
<td>51.3</td>
<td>191</td>
<td>130</td>
</tr>
</tbody>
</table>

“Primary” refers to dogs that had never received a rabies vaccination prior to this study, “booster” refers to dogs that had previously received ≥1 rabies vaccination and “no record” means there was no vaccination history available or taken.

a One sample vial was empty on arrival in the UK.

b One sample vial was empty on arrival in the UK.

Rabbit anti-sera raised against LBV isolate RV1 (n = 2) and DUVV isolate RV131 (n = 4) were used in neutralisation assays with African lyssavirus pseudotypes. Anti-sera to MOKV were not available but anti-LBV serum is known to cross-neutralise MOKV. One serum raised against uninfected tissue culture supernatant and another against Australian bat lyssavirus (RV634) were used in the same panel as the LBV and DUVV sera.

2.4. Infection and neutralisation assays

Human embryonic kidney 293T cells [25] were used for production of the lentiviral pseudotypes (lyssavirus surrogates). Neutralisation assays were undertaken on baby hamster kidney 21 cells clone 13 (BHK; [26]). The NP2 human glioma cell line expressing CD4 and CXCR4 was used as target cells for HIV-1 pseudotypes [27].

2.4.1. FAVN assays

Live virus experiments were undertaken using a restricted version of the FAVN that is identical to the existing FAVN assay [19] but with serum samples diluted 3-fold to a final titre of 1:81, roughly equivalent to 5.92 IU/ml.

2.4.2. Pseudotype assays

Standard plasmids and the transfection protocol for lentiviral pseudotype (lyssavirus surrogate) production have been described elsewhere [24]. Additionally, glycoprotein (G) gene sequences from LBV (LBV.SA2004; accession number EF547428), MOKV (MOKV.98/071 RA361; accession number GQ500108) and DUVV (DUVV.RSA2006; accession number EU623444) were amplified by PCR using specific primers (detailed in Supplementary Table S1). Pseudotypes containing the HIV-1 envelope gp160 gene were generated using pSVI1env [28]. G gene sequence analysis was undertaken using ClustalW [29] and Treeview [30].

Infection and neutralisation assays using lyssavirus pseudotypes were performed as previously described [24] with the following three modifications: (1) plates were centrifuged at 500 rpm for 10 s after the virus was added and once more following addition of BHK cells; (2) to ensure the OIE standard reference dog serum recorded an IC100 at a 1:40 dilution, approximately 30× TCI50 of pseudotype was used; (3) serum was diluted 5-fold to a final titre of 1:640.

2.5. Pseudotype reporter genes

2.5.1. β-Galactosidase

To enable pseudotype particle detection using the appropriate β-galactosidase (β-gal) substrates, we constructed pCSLZW, which contains the lacZ gene from pMFG-nls-lacZ (a kind gift from Dr. Yasuhiro Takeuchi; [31]) cloned into pCSGW (primers detailed in Supplementary Table S1). Where β-gal was the reporter protein, β-gal-pseudotype-infected cells were detected.
**Fig. 2.** Strong correlation between FAVN and pseudotype neutralising antibody titres using Tanzanian dog sera. (A) Neutralising titres achieved with the pseudotype assay increase concordantly with those detected in the FAVN assay. The distribution, according to VNAb titres determined by the (B) pseudotype and (C) FAVN assays, of sera collected at the first visit from primary (dogs that had not received a previous rabies vaccination—black columns), booster (dogs that had received ≥1 rabies vaccination previously—grey columns) or dogs with no vaccination record (white columns) is shown. Similar analyses for samples collected at the second visit are shown in (D) and (E) for the pseudotype and FAVN assays, respectively. The dotted line marks the level of VNAb that was achieved by the OIE positive control serum. Percentages of samples with an inadequate or adequate VNAb response are given left and right of the dotted line, respectively. Vaccination history was available for 169 dogs (first visit: primary \( n = 119 \), booster \( n = 50 \), no record \( n = 13 \); second visit: primary \( n = 84 \), booster \( n = 37 \), no record \( n = 1 \)). (F) Results from the pseudotype neutralisation and FAVN assays reveal a high degree of concordance (Sn: sensitivity and Sp: specificity) and a strong correlation (\( r \)) between titres. Sn and Sp are relative to the 0.5 IU/ml threshold and \( r \) values were calculated using Pearson’s product-moment correlation. Arrows indicate VNAb titres for which there were no serum samples containing that level of antibodies.

Using 5-bromo-4-chloro-3-indolyl-beta-D-galactopyranoside (X-gal; Sigma), chlorophenol red-beta-D-galactopyranoside (CPRG; Sigma) or o-nitrophenyl-beta-D-galactopyranoside (ONPG; Sigma). Detection using the X-gal substrate was accomplished by fixing the infected cell monolayer and then staining the cells with substrate buffer (10 mM deoxycholic acid, 5 mM potassium ferrocyanide, 5 mM potassium ferricyanide, 4.3 mM magnesium chloride and 0.02% (v/v) NP40) containing 1 mg/ml X-gal. For CPRG and ONPG substrates, nuclei were lysed and 50 \( \mu l \) of reaction buffer (120 mM Na2HPO4·2H2O, 80 mM NaH2PO4·H2O, 2 mM MgCl2, 100 mM \( \beta \)-mercaptoethanol) was added containing CPRG or ONPG at a final concentration of 29.2 and 1.3 mg/ml, respectively. This reaction was...
Table 2
Distribution of sera according to FAVN neutralising antibody titre.

<table>
<thead>
<tr>
<th>FAVN titre (IU/ml)</th>
<th>Number of samples (n / %)</th>
<th>Discordant results (n / %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.07</td>
<td>32 / 10.5</td>
<td>0 / 0</td>
</tr>
<tr>
<td>0.1</td>
<td>9 / 2.9</td>
<td>0 / 0</td>
</tr>
<tr>
<td>0.13</td>
<td>13 / 4.3</td>
<td>0 / 0</td>
</tr>
<tr>
<td>0.17</td>
<td>17 / 5.6</td>
<td>0 / 0</td>
</tr>
<tr>
<td>0.22</td>
<td>21 / 6.9</td>
<td>0 / 0</td>
</tr>
<tr>
<td>0.29</td>
<td>19 / 6.2</td>
<td>0 / 0</td>
</tr>
<tr>
<td>0.38</td>
<td>10 / 3.3</td>
<td>0 / 0</td>
</tr>
<tr>
<td>0.5</td>
<td>18 / 5.9</td>
<td>12 / 70.6</td>
</tr>
<tr>
<td>0.66</td>
<td>7 / 2.3</td>
<td>1 / 5.9</td>
</tr>
<tr>
<td>0.87</td>
<td>7 / 2.3</td>
<td>3 / 17.6</td>
</tr>
<tr>
<td>1.14</td>
<td>4 / 1.3</td>
<td>1 / 5.9</td>
</tr>
<tr>
<td>1.5</td>
<td>1 / 0.3</td>
<td>0 / 0</td>
</tr>
<tr>
<td>1.97</td>
<td>14 / 4.6</td>
<td>0 / 0</td>
</tr>
<tr>
<td>2.6</td>
<td>6 / 2.0</td>
<td>0 / 0</td>
</tr>
<tr>
<td>3.42</td>
<td>8 / 2.6</td>
<td>0 / 0</td>
</tr>
<tr>
<td>4.5</td>
<td>14 / 4.6</td>
<td>0 / 0</td>
</tr>
<tr>
<td>5.92</td>
<td>104 / 34.2</td>
<td>0 / 0</td>
</tr>
</tbody>
</table>

Percentages given are the proportion in that category out of the total number for the column.

*Samples that were positive by FAVN assay but negative by the pseudotype assay.

inhibited, after 1–2 h at 37 °C, by adding stop solution (1 M sodium carbonate).

2.5.2. GFP and luciferase

GFP and luciferase reporter genes were employed to detect GFP-positive cells, which were visualised using a fluorescent microscope and FACS Calibur (BD Biosciences) and luciferase expression, which was detected using the Bright-Glo reagent and GloMax 96 micropalte luminometer (Promega).

3. Results

3.1. Serum samples analysed

Of the 191 dogs enrolled in this study, the mean age was 1.94 years and 51.3% were female (Table 1). In total, 321 samples were taken during both visits (Table 1). However, in transport one sample disappeared (#27) and one leaked so no serum remained (#192). Of the remaining 319 samples, the volume in 12 was too small to test in the FAVN assay and a further three samples could not initially be tested in the pseudotype assays because of contamination and toxicity. The remaining 304 samples were tested in both assay types.

While the serum process times were kept as constant as possible there were varying degrees of haemolysis and volumes of sera produced varied with each collection (Fig. 1C). Haemolysis did not affect the results obtained with either the FAVN or the pseudotype assays.

3.2. Sensitivity and specificity of CVS-11 pseudotypes using Tanzanian canine sera

Sera that failed to achieve an antibody titre of 1:10 by the pseudotype assay, the lowest recordable result from the dilutions that were used, were given an arbitrary titre of 1:5 for this analysis. The OIE positive control serum routinely neutralised 100% of CVS-11 pseudotype particles at a dilution of 1:40 (in this study the OIE mean dilution = 37.5 and standard deviation = ±14.4) and 100% of live CVS-11 at a dilution of 15.59, normally equivalent to 0.5 IU/ml, for the FAVN assay (in this study the OIE mean dilution = 17.2 and standard deviation = ±4.0). The lowest titre a serum sample was given by the FAVN assay in this study is 0.07 IU/ml. Any dogs with VNAb levels that attained an IC100 at a dilution of greater than or equal to 1:40 (by pseudotype assay) or a titre of 0.5 IU/ml (by FAVN assay) were considered positive (also referred to as “adequate” in this study).

When results from the FAVN and pseudotype assays were compared, we observed a concurrent increase in VNAb titres reported by the pseudotype assay as the FAVN titre increased (Fig. 2A). There were 17 discordant samples, classified as negative by the pseudotype assay but positive with the FAVN assay (Table 2). As a result, the average titre of samples classified as borderline positive by the FAVN assay (0.5 IU/ml) is 1:25, a borderline negative result with the pseudotype assay. While these discordant samples reduced the sensitivity of the pseudotype neutralisation assay to 94.4%, the specificity of the assay was 100% (n = 304). The majority of the discrepancies clustered around 0.5 IU/ml, the titre achieved using the OIE positive control serum and currently used to classify VNAb responses as inadequate (levels below that achieved by the OIE positive control serum) or adequate (levels greater than or equal to that achieved by the OIE positive control serum). Twelve of the discordant results (71%) were in the lowest dilution classified as positive by the FAVN assay (0.5 IU/ml), one (5.9%) fell within the 0.66 IU/ml category, three (17.6%) in the 0.87 IU/ml category and one (5.9%) scored 1.14 IU/ml by the FAVN assay (Table 3).

There were no discrepant results achieved using the FAVN or pseudotype assays and samples from the second visit.

Prior to vaccination at the first visit, results using the pseudotype assay classified 89.9% of the dogs that had never been vaccinated

Table 3
Titres recorded for discordant sera using the FAVN and pseudotype assays.

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>FAVN titre (IU/ml)</th>
<th>Pseudotype titre (dilution)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>0.5</td>
<td>5</td>
</tr>
<tr>
<td>41</td>
<td>0.5</td>
<td>20</td>
</tr>
<tr>
<td>60</td>
<td>0.5</td>
<td>5</td>
</tr>
<tr>
<td>73</td>
<td>0.5</td>
<td>25</td>
</tr>
<tr>
<td>75</td>
<td>0.5</td>
<td>5</td>
</tr>
<tr>
<td>81</td>
<td>0.5</td>
<td>20</td>
</tr>
<tr>
<td>94</td>
<td>0.5</td>
<td>5</td>
</tr>
<tr>
<td>157</td>
<td>0.5</td>
<td>5</td>
</tr>
<tr>
<td>171</td>
<td>0.5</td>
<td>5</td>
</tr>
<tr>
<td>189</td>
<td>0.5</td>
<td>5</td>
</tr>
<tr>
<td>198</td>
<td>0.5</td>
<td>5</td>
</tr>
<tr>
<td>200</td>
<td>0.5</td>
<td>5</td>
</tr>
<tr>
<td>69</td>
<td>0.66</td>
<td>20</td>
</tr>
<tr>
<td>42</td>
<td>0.87</td>
<td>30</td>
</tr>
<tr>
<td>102</td>
<td>0.87</td>
<td>10</td>
</tr>
<tr>
<td>165</td>
<td>0.87</td>
<td>30</td>
</tr>
<tr>
<td>168</td>
<td>1.14</td>
<td>20</td>
</tr>
</tbody>
</table>

* Vaccination history was not available for this dog (“no record”).
* Serum samples taken from dogs with no rabies vaccination history prior to this study (“primary”).
Fig. 3. Radial tree and panels showing degree of glycoprotein identity and viral titre of lyssavirus pseudotypes. The envelope G genes from LBV, MOKV and DUVV were cloned into pI.18 and titres were compared with that achieved by pseudotypes bearing the CVS-11 G. Identity of the full-length G amino acid sequence was used to construct the tree. Viral titres were assessed using GFP carrying pseudotypes and are given in IFU/ml. The branch lengths and scale correspond to the number of amino acid substitutions per site.

Fig. 3. Radial tree and panels showing degree of glycoprotein identity and viral titre of lyssavirus pseudotypes. The envelope G genes from LBV, MOKV and DUVV were cloned into pI.18 and titres were compared with that achieved by pseudotypes bearing the CVS-11 G. Identity of the full-length G amino acid sequence was used to construct the tree. Viral titres were assessed using GFP carrying pseudotypes and are given in IFU/ml. The branch lengths and scale correspond to the number of amino acid substitutions per site.

3.3. African lyssavirus pseudotypes for surveillance

We cloned the G gene sequences of LBV, MOKV and DUVV into our expression plasmid and tested their ability to be incorporated into lentiviral pseudotype particles. Using GFP as a marker for infection of BHK cells we observed comparable titres for LBV and MOKV pseudotypes as we achieved using CVS-11 (3.3 × 10^5, 3.6 × 10^5 and 3.2 × 10^5 IFU/ml, respectively; Fig. 3). The titre recorded for pseudotypes expressing the DUVV G was 1.5-fold higher at 5.2 × 10^5 IFU/ml. To test the specificity of these African lyssavirus pseudotypes, neutralisation assays were run with sera raised against LBV (RV1; serum nos. 1 and 2) and DUVV (RV131; serum nos. 3, 4, 111 and 112) isolates. Neutralising titres with the pseudotypes correlated strongly with the titres observed using the FAVN assay (Table 4). Anti-LBV sera cross-neutralised MOKV in both the FAVN and pseudotype assays. However, the FAVN assay failed to detect anti-DUVV VNAb in serum 4 and 111 and only low levels in serum 3 (IC100 = 8) compared to the pseudotype assay, which recorded IC100 titres of 160, 8 and 28 for sera 3, 4 and 111, respectively (Table 4). The negative control and Australian bat lyssavirus (ABLV) sera did not achieve an IC100 against any genotype, in either assay, and no cross-neutralisation of DUVV by the LBV sera was observed (Table 4).

3.4. Utility of rabies (CVS-11) pseudotypes carrying the lacZ reporter gene

To broaden the lyssavirus pseudotype platform, we cloned the lacZ gene as a reporter. Using a panel of 33 serum samples selected to contain low, medium and high VNAb titres, we ran parallel assays and determined that using lacZ as the reporter gene, instead of luciferase, did not alter the high correlation (r = 0.918, p < 0.001 [Pearson’s product–moment correlation]; Fig. 4A) or sensitivity and specificity with respect to the titres achieved using the FAVN assay. While the flexibility to use different read-outs is a key requirement of any assay to be used in Africa, it must also be robust, able to withstand fluctuations in temperatures. Therefore, we tested the stability of the pseudotype stocks stored under different conditions. Freeze-thaw cycles of CVS-11 pseudotypes revealed an average decrease of 3.7% in viral titre per cycle, while pseudotypes bearing the VSV or HIV-1 envelope proteins lost 2.6% and 9.2%, respectively (Fig. 4B). Compared to the CVS-11 pseudotype stock stored at −80 °C, aliquots stored at room temperature (average of 23 °C) had a half-life of 1–2 weeks, which increased to 2–4 weeks for aliquots stored at +4 °C. Pseudotypes stored at −20 °C were relatively stable for over 6 months (Fig. 4C). This stability was similar to that observed with
Table 4

Levels of African lyssavirus-neutralising antibodies in sera from immunised rabbits as determined by FAVN and pseudotype assays.

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Immunogen</th>
<th>Lagos bat virus</th>
<th>Mokola virus</th>
<th>Duvenhage virus</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FAVN titre</td>
<td>Pseudotype titre</td>
<td>FAVN titre</td>
<td>Pseudotype titre</td>
</tr>
<tr>
<td>1</td>
<td>RV 1 (LBV)</td>
<td>128</td>
<td>508</td>
<td>128</td>
</tr>
<tr>
<td>2</td>
<td>RV 1 (LBV)</td>
<td>362</td>
<td>508</td>
<td>64</td>
</tr>
<tr>
<td>3</td>
<td>RV 131 (DUVV)</td>
<td>•</td>
<td>•</td>
<td>8</td>
</tr>
<tr>
<td>4</td>
<td>RV 131 (DUVV)</td>
<td>•</td>
<td>•</td>
<td>•</td>
</tr>
<tr>
<td>5</td>
<td>RV634 (ABLTV)</td>
<td>•</td>
<td>••</td>
<td>••</td>
</tr>
<tr>
<td>6</td>
<td>Negative control</td>
<td>•</td>
<td>••</td>
<td>••</td>
</tr>
<tr>
<td>111</td>
<td>RV 131 (DUVV)</td>
<td>•</td>
<td>••</td>
<td>•</td>
</tr>
<tr>
<td>112</td>
<td>RV 131 (DUVV)</td>
<td>•</td>
<td>••</td>
<td>••</td>
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(•): Failed to record a positive titre. (–): Experiment not undertaken. Titres are given as the geometric mean dilution that achieved an IC100.

Fig. 4. Validation of lacZ-based pseudotypes as a tool for in-field diagnostics to detect neutralising antibodies. (A) Selected samples were run using the lacZ-based CVS-11 pseudotypes. The results show an identical correlation \( r \) between the FAVN results and those achieved using luciferase-based pseudotypes. Sensitivity (Sn) and specificity (Sp) relative to the 0.5 IU/ml cut-off are given and \( r/p \) values were calculated using Pearson’s product–moment correlation. The stability of CVS-11 lacZ-pseudotypes was tested by (B) subjecting aliquots to round of freeze–thaw or (C) storing aliquots at different temperatures over time. Results for pseudotypes bearing the VSV envelope glycoprotein and was 7-fold greater than the half-life of pseudotypes bearing the HIV envelope glycoprotein stored at +4 °C or room temperature (data not shown).

The inclusion of lacZ as a reporter gene increased the applicability of our pseudotype system in resource-limited laboratories as it allows the determination of VNAb titres without the need for expensive regents or equipment. Cells were infected with lacZ-based pseudotypes and subsequently stained with one of three β-gal substrates. Punctate blue staining of infected cell nuclei was achieved in the presence of X-gal (Fig. 5, far left panels). We also adapted the assay to allow the β-gal colorimetric substrates CPRG and ONPG to be used (Fig. 5, second left and middle panels). These changes, and resulting VNAb titres, were recorded using a microplate reader, reading at 550 nm (CPRG) or 405 nm (ONPG), or simply by eye. The incorporation of lacZ is an additional option to the much used GFP and luciferase reporter genes, which allow a more high-throughput approach but at a far greater expense (Fig. 5, second right and far right panels, respectively).

4. Discussion

While mass culling of stray dogs has previously been shown to be ineffective in controlling the spread of rabies [33,34], this practice is once again being adopted by a number of countries. On the contrary, dog vaccination programs have been enormously successful in controlling canine rabies [35–37], and improved diagnostics to enable rapid serosurveillance in countries undertaking these programs are therefore important. The assay reported here offers a practical, effective and robust solution for rapid lyssavirus serosurveillance. Using lentiviral pseudotypes we can accurately measure the concentration of VNAb and, coupled with the use of lacZ as the reporter gene for pseudotype particle production, removes the need for high containment laboratories and expensive equipment or reagents. This allows the assay to be undertaken in laboratories previously unable to use the existing FAVN. Furthermore, the assay is not just restricted to lyssavirus serosurveillance as other highly pathogenic viruses that have been pseudotyped could be incorporated into this platform [24,38–41].

To our knowledge this is the largest study published to date using pseudotypes in a diagnostic format for serosurveillance. Of the 319 samples received, titres for three could not be determined using the pseudotype assay: one had low level contamination and two caused cellular cytotoxicity in the assay. These issues were overcome using higher concentrations of antibiotics and increasing the number of BHK cells used 2-fold, to \( 2 \times 10^4 \) per assay. In comparison, there were 12 samples that could not be titrated with the FAVN assay because of insufficient serum volume. While serum volume varied between samples, there was sufficient volume for each sample to run duplicate pseudotype neutralisation assays. This highlights another major advantage of using pseudotypes as surrogate viruses in neutralisation assays, only small serum vol-
The introduction of the LBV, MOKV and DUVV G as binding antigens into our pseudotype platform, coupled with existing CVS-11, EBLV-1 and EBLV-2 pseudotypes, means that serum can now be screened to detect VNAbs against six different lyssavirus genotypes. This is of particular importance because large proportions of these assays are performed on sera from bats, from which only small volumes of serum are available.

At a time when emerging infectious disease outbreaks are becoming more frequent and have the potential to spread faster through international travel, technological advances in infectious disease research are making assays more rapid facilitating more automation. However, countries where the outbreaks are most likely to occur are often unable to utilise these new techniques. We have addressed these limitations by developing an assay that can be used for infectious disease serosurveillance and for monitoring vaccine responses within low-containment laboratories. Neutralisation assays based on pseudotypes as a source of target antigen are a useful platform to use at the start of a new outbreak, not only for rabies, but also other enveloped RNA viruses such as SARS coronaviruses, influenza and Ebola viruses. Incorporating the lyssaviral envelope protein (G) into the pseudotype platform creates an assay that allows rapid screening and vaccine evaluation to be performed within weeks of the start of an epidemic.

Acknowledgements

We thank Denise Marston for help with DNA sequence analysis, Nigel Temperton for constructive discussion and Olivia Avdis and the Viral Transmission Dynamics Project for assistance with sample collection. We are indebted to Tanzania Government ministries,
Tanzania National Parks, Tanzania Wildlife Research Institute and Tanzania Commission for Science and Technology for permission to undertake this research. This research was supported by the UK Medical Research Council (grant number G0801176), the UK Department for Environment, Food and Rural Affairs (grant number SEV3500), partial funding from EU FP7 Epizone project, the Royal College of Veterinary Surgeons Trust (grant number GR000 683), the Wellcome Trust, the University of Edinburgh Development Trust and Lincoln Park Zoo.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.vaccine.2009.09.024.

References