Isolation and characterization of cross-reactive human monoclonal antibodies that potently neutralize Australian bat lyssavirus variants and other phylogroup 1 lyssaviruses

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Abstract: Australian bat lyssavirus (ABLV) is a rhabdovirus that circulates in four species of pteropid bats (ABLVp) and the yellow-bellied sheath-tailed bat (ABLVs) in mainland Australia. In the three confirmed human cases of ABLV, rabies illness preceded fatality. As with rabies virus (RABV), post-exposure prophylaxis (PEP) for potential ABLV infections consists of wound cleansing, administration of the rabies vaccine and injection of rabies immunoglobulin (RIG) proximal to the wound. Despite the efficacy of PEP, the inaccessibility of human RIG (HRIG) in the developing world and the high immunogenicity of equine RIG (ERIG) has led to consideration of human monoclonal antibodies (hmAbs) as a passive immunization option that offers enhanced safety and specificity. Using a recombinant vesicular stomatitis virus (rVSV) expressing the glycoprotein (G) protein of ABLVs and phage display, we identified two hmAbs, A6 and F11, which completely neutralize ABLVs/ABLVp, and RABV at concentrations ranging from 0.19-3.12 µg/mL and 0.39-6.25 µg/mL respectively. A6 and F11 recognize overlapping epitopes in the lyssavirus G protein, effectively neutralizing phylogroup 1 lyssaviruses, while having little effect on phylogroup 2 and non-grouped diverse lyssaviruses. These results suggest A6 and F11 could be effective therapeutic and diagnostic tools for phylogroup 1 lyssavirus infections.

Keywords: bat, monoclonal antibodies, lyssaviruses, neutralization, glycoprotein, ABLV, rabies, RABV, phage display

1. Introduction

Australian bat lyssavirus (ABLV) was first isolated in 1996 from a grounded black flying fox (Pteropus alecto) found near Ballina, Australia [1]. Since then, ABLV has been isolated from all four mainland species of flying foxes (Pteropodidae family) as well as the yellow-bellied sheath-tailed bat (Saccolaimus flaviventris), with two genetically distinct lineages circulating in frugivorous (genus Pteropus, ABLVp) [2] and insectivorous (genus Saccolaimus, ABLVs) [3] Australian bat populations. Before the discovery of ABLV, Australia was thought to be devoid of endemic lyssaviruses. Biosurveillance projects over the
years have drastically expanded the number of known ABLV isolates and provided serological evidence of ABLV infection in a variety of Australian microbat populations [4]. While the prevalence of ABLV antigen, indicative of active infection, is <1% in wild bat populations, increased prevalence is observed in wounded, sick, and orphaned bats [4,5]. Indeed, a recent study found that flying fox pups are a uniquely vulnerable group that is potentially at an heightened risk for mass infection [6]. ABLV can be transmitted to humans from a scratch or bite originating from an infected animal. Historically, there have been three documented human ABLV cases [7-10], all of which manifested as fatal acute encephalitis that presented after variable periods of incubation following the exposure event (5 weeks to 2 years) (reviewed in [11]). In addition to the documented human infections, ABLV was also isolated from two fatal horse infections in Australia in 2013 [12].

Taxonomically, ABLV is a rhabdovirus that belongs to the genus Lyssavirus, a group of 17 viral species with the majority having ancestral origins in bats (order Chiroptera). All lyssavirus species are capable of causing fatal neurological disease with symptomatic presentation and disease progression that is indistinguishable from clinical rabies. Phylogenetic analyses have enabled the subdivision of lyssavirus isolates into at least two phylogroups and several ungrouped viruses [13,14]. Phylogroup I includes the prototype lyssavirus, rabies lyssavirus (RABV), ABLV, Duvenhage lyssavirus (DUVV), Aravan lyssavirus (ARAV), Bokeloh bat lyssavirus (BBLV), Irkut lyssavirus (IRKV), Khujand lyssavirus (KHUV), Gannoruwa bat lyssavirus (GBLV) and European bat lyssaviruses, type 1 and 2 (EBLV-1 and EBLV-2). Shimoni bat lyssavirus (SHIBV), Lagos bat lyssavirus (LBV), and Mokola lyssavirus form phylogroup II. Finally, the most genetically divergent lyssaviruses are ungrouped and include West Caucasian bat lyssavirus (WCBV), Ikoma lyssavirus (IKOV), and Lleida bat lyssavirus (LLBV). Taiwan bat lyssavirus (TWBLV) and Kotalachi bat lyssavirus (KBLV) currently remain unclassified [14]. While genetically and serologically distinct from one another, all lyssaviruses are enveloped bullet-shaped viruses with 12 kb negative-sense single-stranded RNA genomes that encode five major polypeptides: nucleoprotein (N), phosphoprotein (P), matrix (M), glycoprotein (G), and viral RNA polymerase (L) [15]. Lyssavirus G monomers organize in trimers on the virion surface, mediating viral attachment to host cell receptors and facilitating the subsequent clatherin-dependent fusion of viral and host cell membranes during viral entry [16-18]. As a surface-expressed viral protein, G is typically the sole target of neutralizing antibodies against lyssaviruses [19,20]. Despite this fact, cross-neutralization between lyssavirus phylogroups is limited, likely due to the high genetic diversity of lyssavirus G sequences [13,21-23].

Following any lyssavirus exposure event, prompt administration of the RABV post-exposure prophylaxis (PEP) protocol is highly recommended. PEP consists of thorough cleansing of the wound area followed by administration of the rabies vaccine and rabies immunoglobulin (RIG) (reviewed in [24]). Currently, there are two species of RIG available for post-exposure management: human RIG (HRIG) and equine RIG (ERIG). While HRIG is safe and effective when included in PEP, supply limitations and high production costs have made this resource widely inaccessible. ERIG is occasionally used to replace HRIG in PEP, however the high immunogenicity of this therapeutic is the cause of substantial safety concerns [25,26], with documented cases of ERIG-associated serum sickness [27]. The absence of a safe, well-sourced passive immunization component in PEP has led many to propose the replacement of RIG with virus-neutralizing human monoclonal antibodies (hmAbs) [28,29]. Here, we developed hmAbs specific for ABLV by using a recombinant vesicular stomatitis virus (rVSV) in which VSV G was replaced by G from ABLVs. This virus was employed as the capture antigen for panning of a naïve human antibody fragment (Fab) library. This screen resulted in identification of two antigen binding fragments (Fabs), F11 and A6, with specific binding to ABLV G. These Fabs were further engineered to generate human IgG1 monoclonal antibodies (hmAbs). We report that A6 and F11 are cross-reactive hmAbs that potently neutralize both ABLV variants, RABV, and other phylogroup I lyssaviruses.
2. Materials and Methods

2.1 The Cells and viruses

HEK293T cells were provided by Gerald Quinnan (Uniformed Services University) and were maintained in Dulbecco’s modified Eagle’s medium (DMEM; Quality Biologicals, Gaithersburg, MD) supplemented with 10% cosmic calf serum (CCS) (Hyclone, Logan, UT) and 2 mM L-glutamine (DMEM-10). Recombinant turbo green fluorescent protein (GFP) expressing vesicular stomatitis viruses (rVSV) that express ABLVs G, ABLVp G, Rabies CVS 11 G, and VSV (Indiana) G glycoproteins, and rABLVp-GFP have been previously described [30,31].

2.2 Phage panning

One ml of a naïve phage Fab library (size about 3x 10^{11}; a gift from Dr. Dimiter S. Dimitrov, University of Pittsburgh Medical School) [32,33] was re-amplified for phage activation and star panning at 10^{11}. Antigens (10^6 plaque forming units (PFU) of VSV-ABLVs-G) was coated in 100 µl with PBS pH 7.4 on a high-adsorbing flat bottom 96-well plate - Incubation for overnight at 4 °C. After 3 rounds of biopanning, the recovered phage enrichment was evaluated by ELISA; in brief, antigen (5x 10^4 PFU VSV-ABLVs-G) coated on ELISA plates, out-phage from the 1st, the 2nd and the 3rd rounds were checked at 2x 10^5/well and detection by anti-M13 HRp and ABTS.

After confirming of phage panning enrichment specific to the VSV-ABLVs-G antigen at the 3rd round of panning, about 40-50 colonies on agar plates of the 3rd out-put panning were selected for monoclonal ELISA; in brief, single colony was grown in 150 µl 2YT with 0.02% glucose and ampicillin for 2 hrs. shaker at 37 °C (double plate was made for back up clones and keep aside at 4 °C), 25 µl of M13K07 in 2YT at 10^6 was added into each well, keep plate stand for 30 min at 30 °C, then 25 µl of 2YT containing ampicillin 100 µg/ml and kanamycin 200 µg/ml was added into each well. The plate was continue to grow at 30 °C with shaker at 200 rpm for ON for phage secretion. Next day, the plate was centrifuged at 4,000 rpm for 12 min. The SN was collected for ELISA. The positive (>0.8 OD 405) clones at a back ground were selected for monoclonal ELISA; in brief, an infectious units (IU) rVSV-ABLVs G, rVSV-ABLVp G, or rVSV-VSV-G per well diluted in 1×PBS. Plates were blocked with 1×PBS containing 5% bovine serum albumin (BSA) and 0.05% Tween-20 (BSA-PBST) for 1 hr at 37 °C. Human mAbs were diluted in 1% BSA-PBST in 2-fold series and were assayed in duplicate. Goat anti-human IgG HRP was used for detection. For each step, plates were incubated at 37 °C for 1 hr and subsequently,

2.3 Isolation and Characterization of hmAbs F11 and A6

The hmAbs F11 and A6 were isolated and characterized using methods previously described [88]. Briefly, a recombinant vesicular stomatitis virus (rVSV) encoding the ABLVs G gene from an isolate of ABLV derived from a yellow-bellied sheath-tailed bat [3] was used to screen a naïve human Fab phage display library for ABLVs G-specific Fabs. ELISAs were performed to identify clones of Fabs with a high binding affinity for ABLVs-G. The variable regions (V_H and V_L) of positive clones were sequenced and used to express and purify Fabs. The V_H and V_L gene segments were then cloned into human IgG1 expression vector pDR12 (provided by D. Burton, Scripps Research Institute, La Jolla, CA), yielding the constructs which are used to produce the IgG1 hmAbs, F11 and A6. Stable expression in 293F cells obtained by re-cloning the IgG1 construct into pcDNA3.1 Hygro-B and selection cell lines with Hygromycin at 200 µg/ml. IgG1 A6 and F11 were purified by Protein G Sepharose affinity chromatography from culture supernatants.

To analyze the antigen-binding activity of the purified hmAbs, Immulon 2HB microtiter ELISA plates (Fisher Scientific, Hampton, NH) were coated overnight at 4 °C with 10^4 infectious units (IU) rVSV-ABLVs G, rVSV-ABLVp G, or rVSV-VSV-G per well diluted in 1×PBS. Plates were blocked with 1×PBS containing 5% bovine serum albumin (BSA) and 0.05% Tween-20 (BSA-PBST) for 1 hr at 37 °C. Human mAbs were diluted in 1% BSA-PBST in 2-fold series and were assayed in duplicate. Goat anti-human IgG HRP was used for detection. For each step, plates were incubated at 37 °C for 1 hr and subsequently,
that of untreated controls. Error bars represent the standard error of the mean (SEM). Greater than 90% of untreated control were completed on a separate HEK293T cells were infected with 100,000 PFU of either VSV recombinant reporter viruses and ABLVp-GFP virus. A total of 5 x 10⁴ HEK293T cells were added to each well, incubated at 37 °C for 20 hr (VSV recombinant reporter viruses) or 48 hr (ABLvp-GFP), and then scored for GFP expression. Neutralization titers were recorded as the hmAb concentration where at least one of the duplicate wells showed GFP expression. Single-cell preparations were made and fixed (2% paraformaldehyde in 1× PBS) and GFP expression, indicative of productive infection, was analyzed by a Nexcelom Vision C Cellometer (Nexcelom Bioscience LLC., Lawrence, MA) capable of fluorescence detection. The percent of infected cells was calculated by dividing the number of GFP positive cells by the total number of cells and multiplying by 100. Results are expressed as percent inhibition relative to that of untreated controls. Error bars represent the standard error of the mean (SEM). Greater than 90% of untreated cells were infected after 20 hr.

2.4 Virus neutralization assays

Purified hmAbs were serially diluted in DMEM-10, in duplicate wells, in a 96-well tissue culture plate and mixed with 5 x 10⁴ IU of either VSV-ABLVs-G-FGP, VSV-ABLVP-G-FGP, VSV-RABV G-FGP, VSV-VSV G-FGP, or ABLVp-GFP reporter viruses for 30 min at 37 °C. Dilutions of purified hmAbs started at 25 µg/mL or 10 µg/mL, respectively, for recombinant VSV reporter viruses and ABLVp-GFP virus. A total of 5 x 10⁴ HEK293T cells were added to each well, incubated at 37 °C for 20 hr (VSV recombinant reporter viruses) or 48 hr (ABLvp-GFP), and then scored for GFP expression. Neutralization titers were recorded as the hmAb concentration where at least one of the duplicate wells showed GFP expression. Single-cell preparations were made and fixed (2% paraformaldehyde in 1× PBS) and GFP expression, indicative of productive infection, was analyzed by a Nexcelom Vision C Cellometer (Nexcelom Bioscience LLC., Lawrence, MA) capable of fluorescence detection. The percent of infected cells was calculated by dividing the number of GFP positive cells by the total number of cells and multiplying by 100. Results are expressed as percent inhibition relative to that of untreated controls. Error bars represent the standard error of the mean (SEM). Greater than 90% of untreated cells were infected after 20 hr.

2.5 Competitive ELISA

Biotinylated hmAbs A6 and F11 were prepared using NHS-PEO-biotin bound to a nickel chelated support matrix according to the manufacturer’s directions (Pierce, Rockford, IL). Immulon 2HB microtiter ELISA plates (Fisher Scientific, Hampton, NH) were coated overnight at 4 °C with 10⁴ IU rVSV-ABLVs-G per well diluted in PBS. Plates were blocked with PBS containing 5% bovine serum albumin (BSA) and 0.05% Tween-20 (BSA-PBST) for 1 hr at 37 °C. Unlabeled hmAbs were diluted in 1% BSA-PBST in 2-fold series starting at 16 µg/mL and were assayed in duplicate. Plates were incubated at room temperature for 30 min and, subsequently, 25 µl of biotinylated hmAb (1 µg/mL) was added to wells and incubated at 37 °C for 30 min. Following incubation, 50 µl of HRP conjugated streptavidin (Pierce, Rockford, IL) was added at a final dilution of 1:5,000 in 1% BSA-PBST and plates were incubated for 1 hr at 37 °C. For each step, plates were washed 6 times with PBST. Plates were incubated with ABTS [2,2´-azinobis (3-ethylbenzthiazolinesulfonic acid)] substrate (Roche, Indianapolis, IN) (100 µl per well) for 30 min with shaking at room temperature. The absorbance was measured for each well at 405 nm, and the average value was calculated from duplicates.

2.6 Lyssavirus neutralizations

The in vitro neutralization activity of hmAb A6 and F11 were determined using a modified micro-neutralization test [34,35]. The test was performed in a humidity chamber on 8-well cell culture slides (Marienfeld, Germany). Briefly, 1.75 µl of A6 (0.71 mg/mL) and F11 (0.469 mg/mL) were 5-fold serially diluted in 7 µl of Dulbecco’s Modified Eagle Medium with Ham’s F12 (DMEM/F12, Gibco, Life Technologies, United States of America) supplemented with 10% fetal bovine serum (FBS, Gibco, Life Technologies, United States of America) and 1% antibiotics (100 U/mL Penicillin, 100 µg/mL Streptomycin and 0.25 µg/mL Amphotericin, Lonza, Switzerland) in each well of the 8-well slides. Wells represented dilutions from 1:10 through 1:781250. To each well, 7 µL of challenge virus [50 focus-forming dose (50FFD₅₀), as determined by titration [36]] was added and incubated at 37 °C with 5% CO₂ for 90 min. Back titration of the challenge viruses and cell-only control were completed on a separate 8-well cell culture slide. After incubation, 14
μL of mouse neuroblastoma cells (C1300 clone) were added to each well. Slides were incubated at 37 °C with 5% CO₂ for 20 hr. Slides were fixed with cold acetone for 30 min and stained with 1:100 diluted FITC-anti-lyssavirus conjugate (Agricultural Research Council-Onderstepoort Veterinary Institute, South Africa) with 0.2% Evans blue as counterstain. In each well, 10 fields at 200× magnification were scored based on the presence/absence of fluorescent foci. All tests were performed in triplicate. The dilutions reported represent the highest dilution where less than 50% of the observed fields contained infected cells (i.e. the 50% end-point titer or IC₅₀).

2.7 Statistics

One-way ANOVA analysis with Dunnett’s multiple comparisons was performed to evaluate hmAb A6 and F11 binding differences between ABLVp-G and ABLVs-G, and VSV-G. Two-way ANOVA analysis with Tukey’s multiplex comparisons was performed to identify significant differences between A6/F11 and m102.4. Significant inhibitory activity of A6 and F11 against rVSV-GFP G variants was determined via two-way ANOVA and Sidak’s multiple comparisons test of hmAb concentration to 0 µg/mL. Figures and statistical analysis were generated using GraphPad Prism version 7.0.

3. Results

3.1 Identification of phage-displayed Fabs A6 and F11 that are specific for ABLVs glycoprotein

We previously reported the isolation of potent henipavirus-neutralizing hmAb m102 through the screening of a large naïve human Fab library against the soluble HeV G glycoprotein [32,33]. Here, we used the same phage library, which contains over 10¹⁰ phage-displayed human Fabs, to identify Fabs that are specific for ABLVs glycoprotein (ABLVs-G). Since soluble ABLV-G remains unavailable, we used a turbo green fluorescent protein (GFP)-encoding replication competent vesicular stomatitis virus (rVSV-∆G) that expresses ABLVs-G (rVSV-ABLVs-G) as the antigen for screening the Fab library. Three rounds of phage panning against rVSV-ABLVs-G recombinant reporter virus resulted in the enrichment of anti-ABLVs-G Fabs A6 and F11. To probe the binding activity of the identified Fabs, we performed phage ELISAs using rVSV-ABLVs-G coated wells (see Materials and Methods).

Fabs A6 and F11 displayed significant binding to ABLVs-G, with A6 exhibiting stronger binding to ABLVs-G than F11 (Figure 1A). Sequencing revealed both anti-ABLVs-G Fabs have unique amino acid sequences that differ by at least one residue in 5 of the 6 Fab complementarity determining regions (CDRs). Notably, the light chain (LC) CDRs exhibited more variability than the heavy chain (HC) CDRs, with CDR-H1 being identical between F11 and A6, and CDR-H2 and CDR-H3 differing by two and one residues, respectively (Figure 1B).

The DNA sequences encoding the LC and HC of Fabs A6 and F11 were then cloned into the CMVp-driven expression vector, pDR12, for conversion to a whole antibody format and expression as IgG1 hmAbs. To test the antigen-binding activity of the purified hmAbs, we conducted ELISAs using recombinant VSV-∆G viruses expressing G from ABLVs, ABLVp, or VSV as the well-coating antigen.
Figure 1. Identification of ABLVs-G binding Fabs A6 and F11. (A) Fabs specific for ABLVs G were identified by phage ELISA screening as described in the Materials and Methods. Bound phage were detected by anti-FLAG HRP conjugated antibody and the resulting solution absorbance at 405 nm shown. (B) Amino acid sequences of light chain and heavy chain complementarity determining regions (CDR-L1-3 and CDR-H1-3, respectively) for Fabs A6 and F11. Residues which differ between A6 and F11 CDRs are indicated by red and blue font, respectively.

We found that purified A6 binds ABLVs-G and ABLVp-G with similar strength while displaying negligible binding to VSV-G (Figure 2A). Interestingly, unlike A6, slight variation was observed in the maximal binding of F11 to ABLVs and ABLVp G variants (Figure 2B). Overall, however, these data show that both hmAbs A6 and F11 exhibit strong binding to ABLVs G and ABLVp G and minimal binding to VSV-G.

Figure 2. Direct binding of anti-ABLVs-G hmAbs to the glycoproteins of ABLVs and ABLVp. The binding of hmAbs (A) A6 and (B) F11 to ABLVs G (circle), ABLVp G (square), and VSV G (triangle) was quantified using ELISA. The average value from duplicate wells is shown. * P-value α <0.05, one-way ANOVA with Dunnett's multiple comparisons.

Competition ELISAs using rVSV-ABLVs-G coated plates and biotinylated hmAbs revealed that A6 (Figure 3A) and F11 (Figure 3B) recognize overlapping epitopes on ABLV G. These results demonstrated the successful conversion of anti-ABLVs-G Fabs A6 and F11 into functional IgG1 hmAbs that recognize overlapping epitopes on both ABLVs G and ABLVp G.
3.2 Neutralization of recombinant ABLV variants and RABV by hmAbs A6 and F11

To test the neutralization activity hmAbs A6 and F11, we first measured their ability to inhibit the infection of HEK293T cells by an ABLVp-GFP reporter virus (rABLVp-GFP), using a virus neutralization assay (see Materials and Methods).

Results from duplicate experiments showed A6 and F11 inhibited viral infection (as evidenced by prevention of GFP expression) when added to cultures at final concentrations of at 0.31 µg/mL and 0.16 µg/mL respectively (Table 1). m102.4, an antibody known to neutralize Hendra virus (HeV) and Nipah virus (NiV) through binding of surface glycoprotein G [32], did not neutralize rABLVp-GFP infection. While phage display was performed using rVSV-ABLVs-G recombinant virus for Fab selection, these data show that hmAbs A6 and F11 possess the ability to neutralize ABLVp infections in vitro.

Since ABLV is the closest genetic relative to RABV, we next tested the ability of the hmAbs to inhibit the infection of HEK293T cells, using rVSV-ΔG reporter viruses expressing ABLVs, ABLVp, or RABV G (Figure 4).

Table 1. Neutralization of rABLVp-GFP infection by hmAbs A6 and F11

<table>
<thead>
<tr>
<th>µg/mL</th>
<th>A6</th>
<th>F11</th>
<th>m102.4</th>
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<tbody>
<tr>
<td>10</td>
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<td>-</td>
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<tr>
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<tr>
<td>0.08</td>
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<tr>
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<td>++</td>
<td>++</td>
<td>ND</td>
</tr>
<tr>
<td>0.02</td>
<td>+++</td>
<td>+++</td>
<td>ND</td>
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*a Neutralization was performed on HEK293T cells infected with rABLVp-GFP viruses as described in Materials and Methods. -, no GFP expression; (+) to (+++) indicates the relative intensity of fluorescence; ND, not determined. Results from duplicate experiments are shown.
Recombinant GFP virus expressing ABLVs G (rVSV-ABLVs-G) displayed less than 5% inhibition of infection in the presence of 0.10 µg/mL of either hmAb. Indeed, greater than 50% inhibition of ABLVs-recombinant virus required 0.19 µg/mL A6 and 0.39 µg/mL F11 (Figure 4A). Conversely, greater than 50% inhibition of rVSV-ABLVp-G (Figure 4B) and rVSV-RABV-G (Figure 4C) infections was achieved by treatment with 0.05 µg/mL A6 or F11.

Furthermore, as detailed in Table 2, A6 neutralized 100% of recombinant rVSV-∆G reporter viruses expressing ABLVs G (3.12 µg/mL), ABLVp G (0.39 µg/mL), and RABV G (0.19 µg/mL) at lower concentrations than those observed for F11 (6.25, 0.39, and 0.39 µg/mL respectively). Taken together, these results demonstrate that hmAbs A6 and F11 are potent cross-reactive antibodies that can neutralize both known ABLV variants as well as RABV CVS variant, another phylogroup I virus.

Table 2. Neutralization of rVSV-GFP infection by hmAbs A6 and F11

<table>
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<tr>
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<th>RABV G</th>
<th>VSV G</th>
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<td>A6</td>
<td>F11</td>
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<td>25</td>
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<td>1.56</td>
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<td>0.78</td>
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<td>0.39</td>
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<td>0.19</td>
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* Neutralization was performed on HEK293T cells infected with rVSV-GFP viruses as described in Materials and Methods. (.), no GFP expression; (+) to (+++) indicates the relative intensity of fluorescence; ND, not determined. Results from duplicate experiments are shown.

Table 3. Neutralization of diverse lyssaviruses by hmAbs A6 and F11
To more broadly characterize the neutralization activity of A6 and F11, we evaluated the neutralization of a panel of diverse lyssaviruses, including members of phylogroup II (Table 3). Consistent with the above data, ABLV and several distinct RABV isolates were potently neutralized by A6 and F11. Similarly, phylogroup I lyssaviruses EBLV-1, EBLV-2, ARAV, DUVV, KHUV, and IRKV were also neutralized by the anti-ABLV-G hmAbs. In fact, A6 and F11 neutralized all tested phylogroup I lyssaviruses with IC₅₀ values of 0.91 ng/mL and 0.60 ng/mL respectively. As expected, lyssaviruses belonging to phylogroup II (MOKV, SHIBV, and three lineages of LBV) and ungrouped lyssaviruses (IKOV and WCBV) were not neutralized by A6 or F11, indicating that A6 and F11 likely only recognize and neutralize phylogroup I lyssaviruses.

### 4. Discussion

In this study we identified two anti-ABLVs-G hmAbs, A6 and F11, which potently cross-neutralize both ABLV variants as well as other phylogroup I lyssaviruses. While sequencing shows A6 and F11 are genetically distinct, competitive ELISA results suggest they bind overlapping epitopes. Extended viral passaging in the presence of each hmAb, followed by sequencing of the resulting escape mutants could help map the precise location of the epitopes bound by these cross-reactive hmAbs. Furthermore, the in vivo activity...
of A6 and F11 against lyssavirus infections can be further elucidated through their use within our established pre-clinical lyssavirus mouse model [38].

The use of polyclonal RIG as the passive immunization component of PEP has presented a variety of complications in areas ranging from safety to accessibility. Unlike RIG, which is derived from the pooled serum of rabies-immune human donors or horses, recombinant virus-neutralizing hmAbs are produced in human cells thus eliminating safety issues associated with blood- and animal-derived antibodies. Recombinant hmAb preparations can be produced in affordable large-scale quantities and assayed in vitro to ensure minimal variation. Replacement of RIG with hmAbs in the PEP protocol could potentially alleviate supply limitations and extend the availability of a complete PEP protocol to more individuals worldwide.

Various combinations of mAbs can be administered concurrently in antibody cocktails—a treatment that, like RIG, mimics the broad polyclonal antibody response observed in natural infections. The distinct viral epitopes recognized by the different mAbs in a single antibody cocktail presents the unique ability to tailor viral counteraction measures and protect against the emergence of resistance variants. The steady progression of virus-specific mAb cocktail development is well exemplified by work done on Ebola virus (EBOV) [39,40]. Indeed, REGN-EB3, a three-mAb cocktail against EBOV-G was recently approved by the FDA as a treatment for EBOV infection [41]. Similar to EBOV, several mAb cocktails have been developed for use against RABV infections [42-46]. However, since mAb cocktail therapy is still a developing field, it is important to consider potential escape mutants that result from cocktail treatments [47,48]. Ultimately, to achieve the greatest impact, extensive scientific characterization of anti-lyssavirus hmAbs must be met with an international effort to not only produce these reagents in high quantity, but also provide them to the public at minimal cost.

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