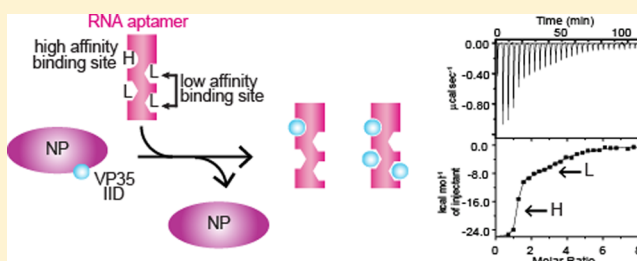


## Development of RNA Aptamers Targeting Ebola Virus VP35

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## S Supporting Information

**ABSTRACT:** Viral protein 35 (VP35), encoded by filoviruses, is a multifunctional dsRNA binding protein that plays important roles in viral replication, innate immune evasion, and pathogenesis. The multifunctional nature of these proteins also presents opportunities to develop countermeasures that target distinct functional regions. However, functional validation and the establishment of therapeutic approaches toward such multifunctional proteins, particularly for non-enzymatic targets, are often challenging. Our previous work on filoviral VP35 proteins defined conserved basic residues located within its C-terminal dsRNA binding interferon (IFN) inhibitory domain (IID) as important for VP35 mediated IFN antagonism and viral polymerase cofactor functions. In the current study, we used a combination of structural and functional data to determine regions of Ebola virus (EBOV) VP35 (eVP35) to target for aptamer selection using SELEX. Select aptamers, representing two distinct classes, were further characterized based on their interaction properties to eVP35 IID. These results revealed that these aptamers bind to distinct regions of eVP35 IID with high affinity (10–50 nM) and specificity. These aptamers can compete with dsRNA for binding to eVP35 and disrupt the eVP35–nucleoprotein (NP) interaction. Consistent with the ability to antagonize the eVP35–NP interaction, select aptamers can inhibit the function of the EBOV polymerase complex reconstituted by the expression of select viral proteins. Taken together, our results support the identification of two aptamers that bind filoviral VP35 proteins with high affinity and specificity and have the capacity to potentially function as filoviral VP35 protein inhibitors.



Negative stranded RNA viruses possess a limited number of open reading frames (ORFs), such that many of the encoded proteins function at multiple stages of the viral replication cycle. The multifunctional nature of these viral proteins potentially provides several opportunities to develop antivirals. However, target validation of nonenzymatic viral proteins is particularly challenging as inhibitors must be identified *de novo*. Innovative methods that combine structure/function information can provide critical target validation prior to the initiation of more effective small molecule-based methods.

Filoviruses are single-stranded, negative sense RNA viruses that cause severe hemorrhagic fever in humans and nonhuman primates.<sup>1</sup> The family *Filoviridae* includes genera *Ebolavirus* and *Marburgvirus* and a proposed genus of *Cuevavirus*.<sup>2</sup> The genus *Ebolavirus* contains five species, which are Zaire Ebola virus (EBOV), Sudan virus (SUDV), Taï Forest virus (TAFV), Reston virus (RESTV), and Bundibugyo virus (BDBV), while the genus *Marburgvirus* has two species, Marburg virus

(MARV) and Ravn virus (RAVV).<sup>3</sup> Among the five species of *Ebolavirus*, RESTV does not cause disease in humans, although it is pathogenic in nonhuman primates.<sup>4,5</sup> However, RESTV was recently isolated from a swine population in the Philippines, whose animal handlers were seropositive suggesting the potential zoonotic nature of filoviruses.<sup>5,6</sup> Together, these observations highlight the potential public health risk posed by filoviruses and the need to develop innovative countermeasures.

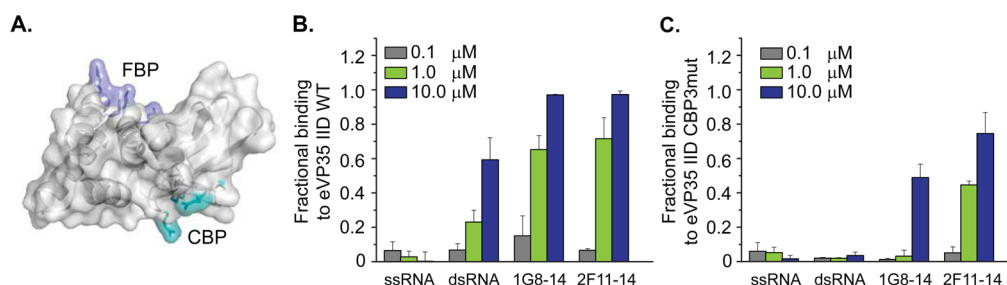
Viral protein 35 (VP35) is a multifunctional immune antagonist that functions at several stages in the filoviral replication cycle.<sup>7–11</sup> VP35 is known to antagonize numerous components in the IFN induction and signaling pathway,<sup>8–17</sup> including RIG-I-like receptors (RLRs),<sup>11,18,19</sup> IFN kinases IKKε

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**Figure 1.** 1G8-14 and 2F11-14 aptamers bind eVP35 IID WT with high affinity. (A) eVP35 IID contains two basic patches. The first basic patch (FBP) contains residues K222, R225, K251, and K248 (purple). The central basic patch (CBP) contains residues R312, K319, and R322 (cyan). eVP35 IID WT and eVP35 IID CBP3mut were used as targets for *in vitro* selection of 1G8-14 and 2F11-14 aptamers, respectively. Filter binding assays were used to assess the ability of (B) eVP35 IID WT and (C) eVP35 IID CBP3mut proteins to bind ssRNA, dsRNA, 1G8-14 aptamer, and 2F11-14 aptamer at protein concentrations of 0.1  $\mu$ M (gray), 1  $\mu$ M (green), and 10  $\mu$ M (blue).

and TBK-1,<sup>20</sup> and RNA-dependent protein kinase (PKR).<sup>13,21</sup> VP35 also functions as an RNAi silencing suppressor,<sup>14,22</sup> a cofactor for the viral polymerase,<sup>23</sup> and a structural protein that is part of the viral nucleocapsid.<sup>24–27</sup> Some, but not all, VP35 mediated functions depend on the ability of VP35 to bind double stranded RNA (dsRNA).<sup>10,11,17,18,23,28</sup> Recent structural and functional studies have identified two conserved basic patches, the first basic patch (FBP) and the central basic patch (CBP), within the C-terminal IFN inhibitory domain (IID) of EBOV VP35 (eVP35) (Figure 1A).<sup>17,18,29,30</sup> The CBP is critical for dsRNA binding and host immune suppression, likely by preventing RNA recognition by RLRs.<sup>18</sup> Viruses containing mutations in the CBP region are replication competent but are attenuated in viral growth. Moreover, prior infection with a guinea pig adapted CBP mutant EBOV protected guinea pigs against subsequent infection by wild type EBOV, further supporting the importance of the VP35 CBP for viral pathogenesis.<sup>28</sup> In contrast, the VP35 FBP is not required for dsRNA binding or inhibiting host immune responses. The filoviral replication complex minimally consists of the viral polymerase L, nucleoprotein (NP), and VP35, with the addition of EBOV viral protein 30 (VP30) being required for viral transcription in EBOV.<sup>31</sup> Previous studies have shown that mutation of residues within the FBP as well as other basic residues that border the CBP (border basic residues) render VP35 defective in its polymerase cofactor function.<sup>23</sup> Mutation of FBP residues leads to loss of critical contacts with the NP protein, an interaction required for replication complex formation.<sup>17,18,23,30</sup> In contrast, mutation of the border basic residues does not result in loss of NP binding.<sup>23</sup> Altogether, these studies highlight the importance of VP35 proteins in virus mediated host immune suppression and viral RNA synthesis, as well as two critical functions which contribute to filoviral pathogenesis.

Despite increasing knowledge of filoviral pathogenesis, there are no approved treatments that target filoviral infections.<sup>1,32–37</sup> Currently, there are several promising vaccine options under investigation;<sup>38,39</sup> however, there are relatively few nonvaccine options available for treatment, particularly for postexposure.<sup>38,39</sup> The multifunctional nature of filoviral VP35 and its relevance to viral pathogenesis makes VP35 a potential target for therapeutic development. Previous efforts using antisense oligonucleotides and small interfering RNAs have been shown to interfere with EBOV replication by targeting VP35.<sup>12,40–42</sup> These studies coupled with recent results, which show that EBOVs containing mutant VP35 are highly attenuated and that guinea pigs infected with these mutant EBOVs are protected

against wild type EBOV infections,<sup>28</sup> suggest that VP35 is a viable target for antiviral development.

Development of inhibitors that target key components in pathogens often requires detailed knowledge of structure and function or screening of large compound libraries with highly sensitive reporter(s). Nucleic acid aptamers provide an alternative method for inhibitor development as well as target validation.<sup>43,44</sup> Aptamer identification and characterization does not require detailed knowledge of the target.<sup>45–48</sup> However, in situations where detailed structural and functional information is available, aptamer development can be tailored to enhance the impact of aptamer binding by targeting aptamer interactions to functionally important regions. Using such an approach, here we report the development of aptamers against eVP35 IID. Previous structural and functional results were utilized to identify functionally relevant regions of VP35 to target for aptamer development. Aptamers were identified using SELEX (systematic evolution of ligands by exponential enrichment<sup>49,50</sup>) methods by targeting WT eVP35 IID and a mutant form of the protein. Two aptamers, representing distinct binding modes, were selected for further characterization. These studies revealed that one aptamer (1G8-14) preferentially binds EBOV and RESTV VP35 IID proteins, while the other (2F11-14) binds both EBOV and MARV VP35 IID proteins. Using *in vitro* studies, we demonstrate the utility of our selected aptamers as potential inhibitors, which can disrupt a critical protein–protein interaction in the filoviral replication complex and the activity of the minigenome system in cells. Taken together, our results suggest that aptamers identified here are valuable reagents for basic research and target validation of filoviral VP35 proteins for antiviral development.

## MATERIALS AND METHODS

**Protein Expression and Purification.** Mutant eVP35 IID proteins were generated by overlap PCR, and the plasmids were sequenced to confirm the mutation(s). Recombinant eVP35 IID WT and mutant proteins were expressed as previously described.<sup>17,51</sup> The integrity of the proteins was assessed by <sup>1</sup>H/<sup>15</sup>N-HSQC NMR spectra.

**Crystallization and Structure Determination.** Initial crystallization conditions for eVP35 IID CBP3mut (where CBP residues R312, K319, and R322 are mutated to alanine) and FBP4mut (where FBP residues K222, R225, K248, and K251 were mutated to alanine) were identified using commercially available crystallization screens (Hampton Research), and in-house optimized native crystals were grown at 25 °C using the hanging-drop vapor diffusion method. 16.4 mg/mL eVP35 IID

CBP3mut was diluted in a 1:1 ratio with the reservoir solution containing 2.2 M Na/K phosphate (pH 4.5). eVP35 IID FBP4mut (24 mg/mL) was diluted in a 1:1 ratio with reservoir solution containing 0.2 M MgCl<sub>2</sub>, 0.1 M HEPES (pH 7.5), and 30% PEG 400. Crystals were soaked in reservoir solution containing 25% glycerol and vitrified in liquid nitrogen. Diffraction data was collected at Advanced Photon Source (Sector 19) at 100 K (Table S1, Supporting Information). One hundred and eighty frames of data were collected with a frame width of 1.0° and detector-to-crystal distance of 300 mm for eVP35 IID FBP4mut. Four hundred fifty frames of data were collected with a frame width of 0.2° and detector-to-crystal distance of 250 mm for eVP35 IID CBP3mut. Diffraction data were indexed, scaled, and merged using HKL-3000.<sup>52</sup> Intensities were converted to structure factors using CCP4.<sup>53</sup> The structures were solved using molecular replacement, with chain B from eVP35 IID crystal structure (PDB ID 3FKE) as the search model using MOLREP.<sup>53</sup> The structures were refined using REFMAC5<sup>54</sup> and PHENIX.<sup>55</sup> Addition of solvent molecules and manual model building was performed using Coot.<sup>56</sup> TLS parameters were refined using the TLMSSD server.<sup>57</sup> Final validation was performed using the MOLPRO-BITY server.<sup>58</sup>

**In Vitro Selection of Aptamers.** *In vitro* SELEX selection of aptamers was carried out using eVP35 IID wild type (WT) and eVP35 IID CBP3mut as targets.<sup>59,60</sup> Oligonucleotide templates and primers were chemically synthesized with standard desalting by Integrated DNA Technology (Coralville, IA). The dsDNA library was generated from a ssDNA library (5'-GCCTGTTGTGAGCCTCCTGTGCGAA(N45)-TTGAGCGTTTATTCTTGTCTCCC-3'). Using the dsDNA library as a template, RNAs were *in vitro* transcribed following established protocols. The transcribed RNA was purified by phenol/chloroform extraction followed by ethanol precipitation and separated out on a 7 M urea–8% PAGE. Purified RNAs were incubated with either eVP35 IID WT or eVP35 IID CBP3mut at 37 °C for 0.5 to 1 h in 10 mM HEPES (pH 7.0), 150 mM NaCl, 2 mM TCEP (tris(2-carboxyethyl)phosphine), and 1 mM MgCl<sub>2</sub> and filtered through a nitrocellulose membrane (Millipore). The protein-bound RNA was eluted with 7 M urea, precipitated by ethanol, and reverse transcribed using SuperScript reverse transcriptase (Invitrogen). The reverse transcribed cDNAs were amplified by polymerase chain reaction (PCR) using oligo-003 (5'-TAATAC-GACTCACTATAGGGAGACAAGAATAAACGCTCAA-3') and oligo-004 (5'-GCCTGTTGTGAGCCTCCTGTGCGAA-3'). *In vitro* transcription, RNA-protein binding, partition, and RT-PCR were repeated for 14 rounds, and the resulting binders at rounds 10 and 14 were subcloned into the pCR-XL-TOPO vector (Invitrogen) for sequencing. Negative selection was carried out against the nitrocellulose membrane at rounds 3, 6, 9, and 12 to reduce nonspecific interactions. The protein concentration for eVP35 IID WT was decreased from 0.4 μM at round 1 to 0.025 μM by round 14 in order to enhance selection stringency. For eVP35 IID CBP3mut, the protein concentration was decreased from 4 μM at round 1 to 0.05 μM by round 14. The concentration of protein used in each selection was adjusted so that we could obtain about 1% retention of aptamers during each selection. Thus, the initial values of 0.4 μM for selection against eVP35 IID WT and 4 μM for selection against eVP35 IID CBP3mut were determined through this process.

**End Labeling of dsRNA and Aptamers.** <sup>32</sup>P 5'-end labeling of RNAs was carried out at 37 °C for 1.5 h in a 20 μL reaction containing 70 mM Tris (pH 7.6 at 25 °C), 10 mM MgCl<sub>2</sub>, 5 mM DTT, 1U/μL T4 polynucleotide kinase (New England Biolabs), 1 μM γ-ATP (Perkin-Elmer), and 5 μM RNA. <sup>32</sup>P 3'-end labeling of RNA was carried out at 16 °C for 18 h in a 30 μL reaction of 50 mM Tris-HCl (pH 7.8 at 25 °C), 10 mM MgCl<sub>2</sub>, 10 mM DTT, 1 mM ATP, 10% DMSO, 0.67 U/μL T4 RNA ligase 1 (New England Biolabs), 2 μM <sup>32</sup>P-cytidine-3',5'-bis-phosphate (<sup>32</sup>pCp), and 5 μM RNA (dsRNA or aptamer).<sup>61</sup> Labeled RNAs were separated on a 7 M urea–8% PAGE and eluted by the crush–soak method.

**Filter Binding Assays for Binding and Competition Assays for the eVP35 IID–Aptamer Interaction.** Filter binding assays were carried out as reported previously.<sup>51</sup> For dissociation constant ( $K_D$ ) measurements, the reaction mixtures contained 1–2 nM <sup>32</sup>P-aptamer or <sup>32</sup>P-dsRNA and eVP35 IID protein ranging from 0 to 100 μM. For single binding point assays, the reaction mixtures contained 10 nM <sup>32</sup>P-aptamer and 0.5 μM eVP35 IID proteins. In the competition assay, the reaction mixtures included 10 nM <sup>32</sup>P-aptamer, 0.5 μM eVP35 IID or 10 nM <sup>32</sup>P-dsRNA, 10 μM eVP35 IID, and unlabeled aptamer or dsRNA ranging from 0 to 20 μM. All reaction mixtures were incubated at 25 °C for 15 min. Binding experiments were carried out in buffer containing 10 mM HEPES (pH 7.0), 150 mM NaCl, 2 mM TCEP, and 1 mM MgCl<sub>2</sub> in 50 μL reactions in a 96-well vacuum filtration apparatus assembled with a nitrocellulose membrane on top of a positively charged nylon membrane. Membranes were washed and air-dried before exposure for 12–16 h along with a calibration standard, scanned on a Typhoon 9410 Variable Mode 1 Imager in phosphor storage mode, and quantified by ImageQuant (GE Healthcare). The ratio of the intensity on the nitrocellulose membrane to that of the sum on nitrocellulose membrane plus nylon membrane was expressed as fractional binding. The binding curves were fit to the equation  $B = B_{\max} [M] / (K_D + [M])$  using ORIGIN 7.0 software (OriginLab), where  $B$  is the fraction bound,  $B_{\max}$  is the maximum fractional binding,  $[M]$  is the protein concentration, and  $K_D$  is the dissociation constant. Competition curves were fitted to the equation  $B/B_0 = 1/(1 + [M]/IC_{50})^n$ , where  $B$  is the fraction of <sup>32</sup>P-labeled aptamers bound to the nitrocellulose membrane at different concentrations of competitors (aptamer or dsRNA),  $B_0$  is the fraction of <sup>32</sup>P-labeled aptamers bound to the nitrocellulose membrane without any competitors,  $[M]$  is the competitor concentration,  $IC_{50}$  is the concentration where 50% competition occurs, and  $n$  (number of binding sites) is a constant.

**Isothermal Titration Calorimetry (ITC).** Quantitative analysis of eVP35 IID protein–aptamer interactions were performed on a VP-isothermal titration calorimeter (VP-ITC) (Microcal). Protein samples were dialyzed against 500 mL of buffer containing 10 mM HEPES (pH 7.0), 150 mM NaCl, 1 mM MgCl<sub>2</sub>, and 2 mM TCEP for 12 h at 25 °C. Aptamer stocks were diluted >200-fold with dialysis buffer. The syringe contained eVP35 IID proteins at 100 to 150 μM, and the cell contained aptamers (1G8-14, 2F11-14) at 3 to 5 μM. ITC titrations were carried out using a reference power of 5 μcal/s. The resulting ITC data were processed and fit to a one-site binding model or a two-site binding model to determine  $n$  (number of binding sites) and  $K_D$  (dissociation constant) using ORIGIN 7.0 software. Both one- and two-site binding models were considered for each data set, but selection of the model



was based on the quality of the fits. For two-site binding, both sequential and independent binding modes were also considered. The two-site binding model used in this study assumes two thermodynamically independent binding events.

**Pulldown Assays.** Pulldown assays were performed in buffer containing 10 mM HEPES (pH 7.0), 150 mM NaCl, 1 mM MgCl<sub>2</sub>, and 5 mM 2-mercaptoethanol at 25 °C. MBP-His tagged eVP35 IID WT or mutant proteins were immobilized on amylose resin. Resin bound MBP-His tagged eVP35 IID was incubated with purified His-tagged NP protein and subsequently washed. For aptamer competition assays, either 100  $\mu$ M aptamer or 300  $\mu$ M dsRNA was incubated with MBP-His tagged eVP35 IID prior to incubation with NP. The level of protein used in the assay reflects our efforts to ensure proper detection of the pulldown experiment. Pulldown samples were separated on SDS-PAGE and analyzed by Western blot using mouse anti-His antibody (Santa Cruz biotechnology), followed by horseradish peroxidase (HRP) conjugated goat antimouse antibody (Bio-Rad). Membranes were developed using Millipore Immobilon Western Chemiluminescence HRP substrate and recorded on a ChemiDoc (Bio-Rad).

**EBOV Transcription/Replication Assays.** The plasmids used in the EBOV transcription/replication assays were described previously.<sup>18,23,31</sup> NP, VP35, VP30 and L proteins were cloned into pTM1 with only the cDNA sequence (no virus-derived UTRs were present). Aptamers were cloned into a modified pSilencer plasmid,<sup>62</sup> and the corresponding plasmids were sequenced to verify the accuracy of the aptamers.

**Quantitation of Total Aptamer Copy Number in Transfected Cells by qRT-PCR.** Genome equivalent (GEq) copy numbers (total copies) of aptamers in the minigenome assay were measured using primer/probes targeting the aptamers. Briefly, real-time PCR (RT-PCR) was performed using Superscript II RT-PCR kits (Invitrogen). All RT-PCR mixtures contained 5  $\mu$ L of RNA eluate, and master mixes were set up following the manufacturer's protocols. Standards and test samples were assayed in triplicate using the CFX96 detection system (Bio-Rad) with the following cycle conditions: 50 °C for 10 min, 95 °C for 5 min, and 40 cycles of 95 °C for 10 s, and 59 °C for 30 s. Threshold cycle (CT) values representing aptamers were used to determine Total copies. To create the GEq standard, DNA from aptamers stocks (100 ng) was used, and the number of genomes was calculated using Avogadro's number and the total molecular weight of the plasmid with the aptamer.

**Aptamer Secondary Structure Prediction.** The computational program RNAfold from Vienna package v.2.0<sup>63</sup> was used to predict the secondary structures for RNA aptamers 1G8-14 and 2F11-14. Default parameters on the RNAfold server were used except for the Turner Model 2004,<sup>64</sup> which was used as the RNA parameter source. Results were assessed for the minimum free energy (MFE) structure prediction as well as thermodynamic ensemble predictions, including centroid structure<sup>65</sup> and base-pairing probabilities using dot plot analysis<sup>66</sup> as implemented in RNAfold. Resulting secondary structures for the MFE and the centroid were visualized with the RNA graphics program VARNA.<sup>67</sup>

**Circular Dichroism (CD).** CD data were collected for 1G8-14 and 2F11-14 at 10  $\mu$ M in a quartz cuvette with a 1 mm path-length using a Chirascan CD spectrometer. CD spectra were acquired from 200 to 320 nm at 5 and 95 °C at an interval of 1 nm and time-per-point of 2 s. Temperature melting of aptamers was performed by monitoring the ellipticity at 260 nm from 5

to 95 °C with a 2 °C increment at a 1 °C/min gradient. ORIGIN 7.0 software was used to plot CD spectra and to calculate first derivatives from the temperature melting plots.

## RESULTS

**Mutation of Conserved Basic Residues Does Not Perturb the Overall Fold of eVP35 IID.** The CBP and FBP contain highly conserved basic residues in VP35 IID that are functionally important as mutations of either the CBP or the FBP and impair several VP35 mediated functions.<sup>7</sup> Therefore, a series of structural studies were carried out to characterize the impact of these mutations on the integrity of the structure of eVP35 IID. These studies resulted in crystal structures of eVP35 IID CBP3mut (PDB code 4IJE) and FBP4mut (PDB code 4IJF), which were solved to a resolution of 1.9 and 2.5 Å, respectively using the eVP35 IID WT (PDB code 3FKE) structure as the search model for molecular replacement (Table S1, Supporting Information). These structures revealed minimal changes in the overall protein fold when compared to those of eVP35 IID (Figure S1, Supporting Information). The most notable difference between the structures was observed in a short helix ( $\alpha$ 5) located within the  $\beta$ -sheet subdomain, in which the CBP3mut and FBP4mut structures make one turn of the helix instead of two. Comparison of the eVP35 IID WT, CBP3mut, and FBP4mut structures revealed that eVP35 IID CBP3mut and eVP35 IID FBP4mut structures display RMSD values of 0.68 and 0.46 Å, respectively, which further suggest minimal structural changes upon mutating out the CBP and FBP. Analysis of surface electrostatics, however, showed significant differences among the three structures (Figure S2, Supporting Information). As expected, generation of the CBP3mut resulted in a loss of the basic charge at the CBP, but the FBP was relatively unaffected, while the structure of FBP4mut revealed that the basic charge at the FBP was lost with the CBP unaffected.

### SELEX Identifies RNA Aptamers That Bind eVP35 IID.

In order to generate different classes of aptamers for eVP35 IID, aptamer selection was carried out against two targets, eVP35 IID WT and eVP35 IID CBP3mut proteins using SELEX (Figure 1A). By using eVP35 IID WT and CBP3mut proteins, we biased the aptamer selection to different conserved regions of eVP35 IID. We expected that the dsRNA binding site formed by CBP residues would dominate aptamer selection using eVP35 IID WT as a target due to the highly basic charge and dsRNA binding properties in this region. In contrast, by mutating the CBP (in eVP35 IID CBP3mut), we reduced the basic nature of the CBP region with the intent that other charged regions, preferably the FBP, could compete during aptamer selection. The initial RNA pool contained RNA species of approximately 90 bases, with 45 nucleotides in the variable region flanked on either side by a 5' and a 3' common region. This initial pool bound eVP35 IID WT with a  $K_D$  of  $2900 \pm 200$  nM and bound eVP35 IID CBP3mut with a  $K_D$  of  $36000 \pm 8000$  nM. After several rounds of SELEX, each aptamer pool displayed enhanced affinity and specificity for eVP35 IID WT and eVP35 IID CBP3mut proteins. Our progress curves show >10-fold affinity enhancement for the final aptamer pool, with an affinity of  $300 \pm 100$  nM at round 14 for eVP35 IID WT (Figure S3A, Supporting Information) and  $2800 \pm 600$  nM at round 10 for eVP35 IID CBP3mut (Figure S3B, Supporting Information). We also extended our selection against eVP35 IID CBP3mut up to round 14. Although this pool displayed a high degree of nonspecific

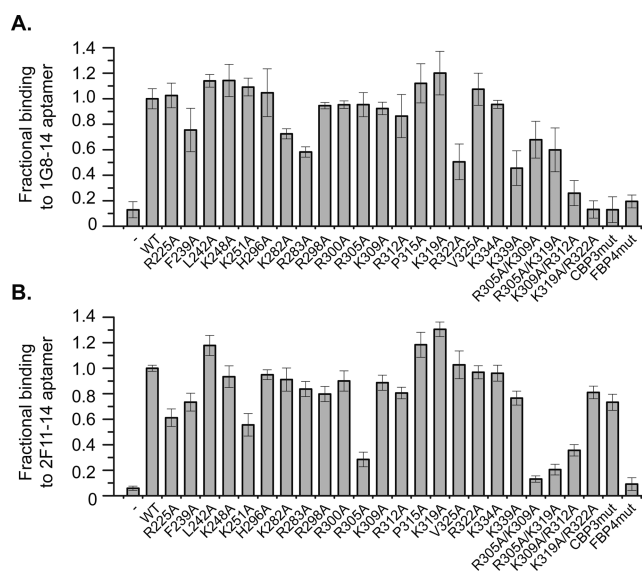
binding to the nitrocellulose membrane, we were able to identify select individual aptamers from rounds 10 and 14. As discussed below, we were able to further characterize these select aptamers for affinity measurements and ranking.

For our initial SELEX studies, we observed significant enrichment of select sequences from a total of 96 clones. Using ClustalW multiple sequence alignments, sequences with variations less than 5% of the total aptamer length (i.e., 4 nucleotides or less that includes the constant/primer regions and the variable region) were grouped together and represented by a single member. We first selected 9 aptamers from the 1-series and 3 aptamers from the 2-series (Figure S3C, Supporting Information). Subsequently, we identified representative members from each group based on their affinity for eVP35 IID WT and their ability to inhibit the interaction between VP35 IID and NP (see below). These selected aptamers were subjected to further characterization and ranking based upon affinity measurements with the top binders from each series selected for further analysis. Aptamer 1G8-14 was chosen to represent the 1-series aptamers, which were selected against eVP35 IID WT, and aptamer 2F11-14 was chosen to represent the 2-series aptamers, which were selected against eVP35 IID CBP3mut. In order to further examine RNA structural determinants, we used ViennaRNA structural prediction software to predict the RNA secondary structure of 1G8-14 and 2F11-14. The resulting secondary structure predictions for the MFE and the centroid were visualized with the RNA graphics program VARNA,<sup>67</sup> and base-pairing probabilities for the MFE and ensemble are shown using dot plot analysis (Figure S4, Supporting Information).<sup>54</sup> On the basis of the structure predictions, the 1G8-14 MFE and centroid structure displayed 15.12 ensemble diversity, and the MFE structure represented 5.12% of the ensemble compared to the 2F11-14 MFE and centroid structure with 13.46 ensemble diversity and 4.23% of the ensemble represented by the MFE. However, the free energy of the ensemble for 1G8-14 was  $-22.63$  kcal/mol compared to  $-12.75$  kcal/mol for 2F11-14 suggesting that the 1G8-14 aptamer may adopt a more stable structure. Our CD data show differences in the corresponding spectra providing some experimental evidence that indicate structural differences between 1G8-14 and 2F11-14 aptamers (Figure S5, Supporting Information). However, additional studies are required to appropriately characterize the actual structural differences between 1G8-14 and 2F11-14 aptamers. As an initial step in our efforts to define the optimum aptamer, we carried out boundary determination studies and tested several shorter aptamers based on the results of our boundary determination experiments. These results are described in the section entitled Primer Binding Region of 1G8-14 Is Important for High Affinity VP35 IID Binding (see below). These studies supported the use of full length aptamers in our present studies as discussed below.

Aptamers 1G8-14 and 2F11-14 bound eVP35 IID WT with affinities higher than dsRNA. At  $1 \mu\text{M}$  protein, the fractional binding for both 1G8-14 and 2F11-14 was  $>0.5$ , while the fractional binding for dsRNA was  $<0.3$  (Figure 1B). eVP35 IID CBP3mut bound to 2F11-14 with high affinity and to 1G8-14 only at higher protein concentrations ( $10 \mu\text{M}$  protein) (Figure 1C). These binding studies provided an initial indication that we had identified aptamers that bind to eVP35 IID differently, with the CBP residues contributing to 1G8-14 aptamer binding. The ability of eVP35 IID WT and eVP35 IID CBP3mut to bind ssRNA and dsRNA was also assessed. As previously shown,

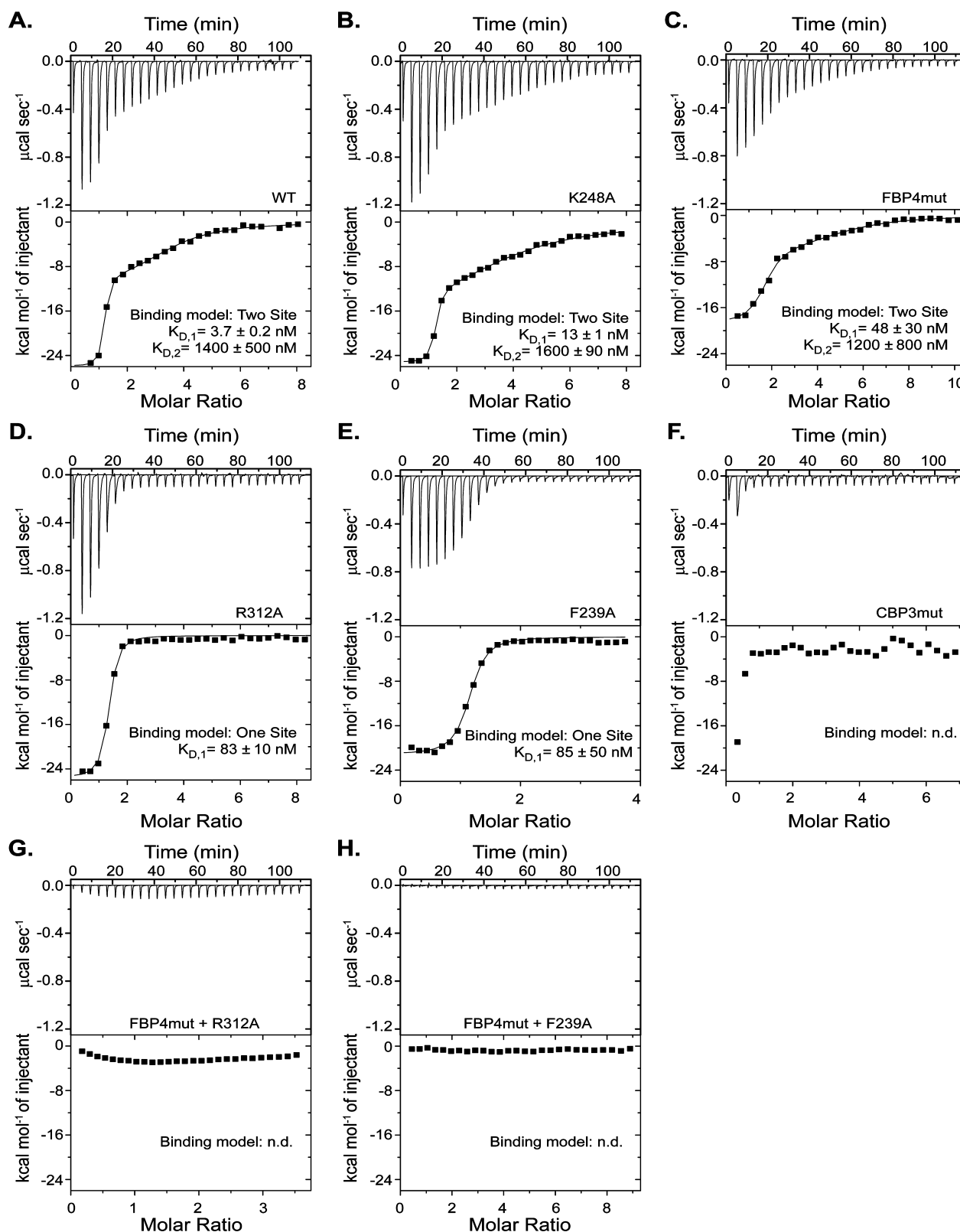
eVP35 IID WT binds to dsRNA but not ssRNA (Figure 1C). eVP35 IID CBP3mut does not bind to dsRNA due to mutations at key dsRNA binding residues, R312, K319, and R322 (Figure 1C).<sup>16,28</sup> Previous studies have shown that eVP35 binds dsRNA containing a 5'ppp moiety with higher affinity than blunt end dsRNA.<sup>18,29</sup> Therefore, we tested if a 5'ppp or 5'OH affected aptamer binding to eVP35 IID. The resulting data suggest that the 5' phosphorylation state of the aptamer does not influence eVP35 IID binding (Figure S6, Supporting Information). Additionally, this observation supports the notion that the eVP35–aptamer interactions are distinct from those observed between eVP35 IID and dsRNA.<sup>11,18</sup>

**Aptamers Recognize Partially Overlapping Sites in eVP35 IID.** We and others have shown that eVP35–dsRNA interactions are mediated through the CBP region of eVP35 IID.<sup>11,16,18,29,30</sup> Therefore, a series of basic residues in eVP35 IID were mutated to determine the contribution of these residues, if any, to 1G8-14 and 2F11-14 aptamer binding. Representative single point binding data for 1G8-14 and 2F11-14 aptamers revealed distinct binding patterns. 1G8-14 aptamer binding primarily depends on a subset of residues that are also important for dsRNA binding and IFN inhibition. Single mutation of residues F239, R305, K309, R312, R322, and K339 to alanine had little effect on 1G8-14 aptamer binding ( $<30\%$  relative to WT), whereas double/triple mutations (K309A/R312A, K319A/R322A, CBP3mut) significantly diminished binding to 1G8-14 ( $>70\%$  relative to WT) (Figure 2A).



**Figure 2.** Mutational analysis reveals differences in eVP35 IID–aptamer binding sites. Fractional binding of aptamers (A) 1G8-14 and (B) 2F11-14 to eVP35 IID WT and mutant proteins measured by filter binding assay. Average fractional binding normalized to eVP35 IID WT are from two independent experiments, each with four repeats for individual eVP35 IID variants.

Mutations of the FBP residues (eVP35 IID FBP4mut) also impair binding to 1G8-14; however, further experiments revealed that this decrease in 1G8-14 binding to eVP35 IID FBP4mut is largely due to a decrease in the maximum fractional binding and not due to a substantial change in binding affinity (Figure S7, Supporting Information). Moreover, these aptamers readily competed with dsRNA in competition assays (Figure S8, Supporting Information). These results are different from



**Figure 3.** CBP is important for high affinity binding of the 1G8-14 aptamer to eVP35 IID. ITC raw data and corresponding binding isotherms for 1G8-14 binding to eVP35 IID (A) WT, (B) K248A, (C) FBP4mut, (D) R312A, (E) F239A, (F) CBP3mut, (G) FBP4mut + R312A, and (H) FBP4mut + F239A.

those observed for dsRNA binding, where select single point mutation of residues in the CBP abolished dsRNA binding.<sup>18</sup> Collectively, these results suggest that aptamer binding involves multiple residues within a given eVP35 IID surface, with the

CBP residues providing the largest contribution toward 1G8-14 aptamer binding.

For the 2F11-14 aptamer, we observe a set of residues different from those important for 1G8-14 binding, including

**Table 1. Measured Binding Affinities between Aptamers and WT or Mutant eVP35 IID Proteins by ITC<sup>a</sup>**

	1G8-14				2F11-14			
	$n_1$	$K_{D,1}$ (nM)	$n_2$	$K_{D,2}$ (nM)	$n_1$	$K_{D,1}$ (nM)	$n_2$	$K_{D,2}$ (nM)
WT	1.1 ± 0.1	3.7 ± 0.2	2.7 ± 0.4	1400 ± 500	1.2 ± 0.0 <sup>b</sup>	7.1 ± 0.1	5.1 ± 0.1	380 ± 40
K248A	1.1 ± 0.0 <sup>c</sup>	13 ± 1	3.3 ± 0.4	1600 ± 90	1.7 ± 0.0 <sup>d</sup>	36 ± 7	5.3 ± 0.1	830 ± 300
FBP4mut	1.4 ± 0.2	48 ± 30	3.2 ± 0.6	1200 ± 800	—	—	—	—
R312A	1.2 ± 0.1	83 ± 10	—	—	1.4 ± 0.0 <sup>e</sup>	52 ± 6	—	—
F239A	1.1 ± 0.0 <sup>f</sup>	85 ± 50	—	—	1.4 ± 0.0 <sup>g</sup>	100 ± 8	—	—
CBP3mut	—	—	—	—	1.3 ± 0.1	56 ± 7	—	—
FBP4mut + R312A	—	—	—	—	n.d.	n.d.	n.d.	n.d.
CBP3mut + K248A	n.d.	n.d.	n.d.	n.d.	1.6 ± 0.2	180 ± 20	n.d.	n.d.

<sup>a</sup>n.d. = not determined; — = no binding. <sup>b</sup> = 1.2 ± 0.04. <sup>c</sup> = 1.1 ± 0.03. <sup>d</sup> = 1.7 ± 0.03. <sup>e</sup> = 1.4 ± 0.01. <sup>f</sup> = 1.1 ± 0.02. <sup>g</sup> = 1.4 ± 0.04.

**Table 2. Summary of ITC Results for Aptamers Binding to WT or Mutant eVP35 IID Proteins<sup>a</sup>**

	dsRNA	1G8-14		2F11-14	
		$K_{D,1}$	$K_{D,2}$	$K_{D,1}$	$K_{D,2}$
WT	++++	++++	++++	++++	++++
K248A	++++	++++	++++	++++	++++
FBP4mut	n.d.	++	++++	—	—
R312A	—	+	—	+++	—
F239A	—	+	—	++	—
CBP3mut	—	—	—	+++	—
FBP4mut + R312A	n.d.	—	—	n.d.	n.d.
FBP4mut + F239A	n.d.	—	—	n.d.	n.d.
CBP3mut + K248A	n.d.	n.d.	n.d.	+	—

<sup>a</sup>++++ = WT; +++ = 5–9-fold decrease in  $K_D$ ; ++ = 10–19-fold decrease in  $K_D$ ; + = >20-fold decrease in  $K_D$ ; — = no binding; n.d. = not determined.

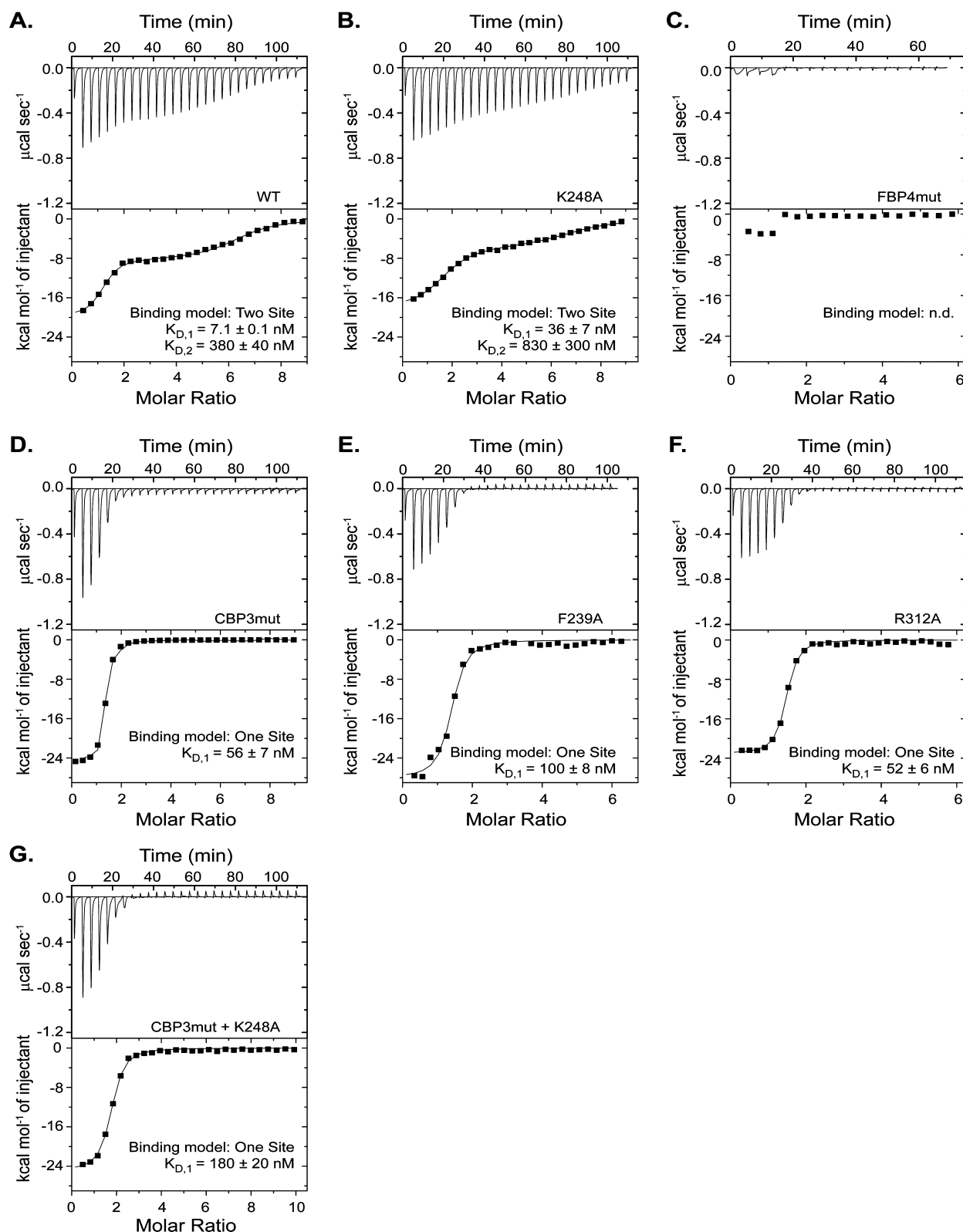
R225, K251, and F239, as mutation of these residues to alanine result in some loss of aptamer binding (<40% relative to WT) (Figure 2B). Most significantly, the R305A mutant alone can drastically decrease binding to 2F11-14 (>70% relative to WT). In contrast to 1G8-14, eVP35 IID CBP3mut had limited impact on 2F11-14 binding, consistent with the single point data (Figure 1C). Residues R305 and K309 are important for both aptamers, suggesting that the aptamer binding sites are partially overlapping at or near these residues.

**Residues in eVP35 IID CBP and FBP Contribute Differentially to Aptamer Binding.** To further characterize eVP35–aptamer interactions, we measured the binding affinities of aptamer to eVP35 IID WT and mutant proteins using ITC. eVP35 IID WT bound the 1G8-14 aptamer with high affinity compared to dsRNA (Figure 3A).<sup>18,23</sup> One- and two-site binding models were considered, ultimately resulting in the ITC data being fit using a two-site binding model, indicating that there are two independent binding modes for eVP35 IID on 1G8-14. The first binding mode is of high affinity ( $K_{D,1}$  = 3.7 ± 0.2 nM;  $n_1$  = 1.1 ± 0.1), which we term the aptamer binding mode, whereas the second binding mode showed moderate affinity ( $K_{D,2}$  = 1400 ± 500 nM;  $n_2$  = 2.7 ± 0.4) (Figure 3A and Table 1). The affinity and the multiple bindings displayed by the second binding mode suggest that this is the dsRNA binding mode. Mutation of residues in the FBP (e.g., K248A and FBP4mut) had little effect on binding (Figure 3B–C). When we test the binding of the aptamer to eVP35 IID proteins containing mutations of residues important for dsRNA binding (e.g., R312A and F239A), we find that the second binding mode is eliminated and that the data can be fit to a one-site binding model ( $K_{D,1}$  values of 83 ± 10 nM and 85 ± 50 nM, respectively) (Figure 3D–E). Furthermore, mutation of three CBP residues together (i.e., R312A/K319A/R322A

triple mutant or CBP3mut) or in combination with mutations of the four FBP residues (i.e., FBP4mut/R312A and FBP4mut/F239A) in eVP35 IID results in complete loss of 1G8-14 aptamer binding (Figure 3F–H). For the second binding mode, these data are consistent with previously reported descriptions of the dsRNA–eVP35 interaction.<sup>18</sup> Altogether, our data suggest that the first binding mode is the high affinity aptamer–eVP35 IID interaction (aptamer mode) and that the second binding mode is eVP35 IID binding to double stranded regions (dsRNA mode) in the aptamer (Table 2).

The ITC binding data for the 2F11-14 aptamer to eVP35 IID were fit to a two-site binding model, where  $K_{D,1}$  = 7.1 ± 0.1 nM ( $n_1$  = 1.2 ± 0.0) and  $K_{D,2}$  = 380 ± 40 nM ( $n_2$  = 5.1 ± 0.1) (Figure 4A and Table 1). A single point mutation in the FBP, K248A, leads to slightly diminished binding with an approximately 5-fold increase in  $K_{D,1}$  and a 2.5-fold increase in  $K_{D,2}$  (Figure 4B). Mutation of four residues in the FBP (eVP35 IID FBP4mut) leads to near complete loss of 2F11-14 binding (Figure 4C), emphasizing a critical role for the FBP in 2F11-14 aptamer binding. Mutation of residues in the CBP eliminates the second binding event with  $K_{D,1}$  = 56 ± 7 nM,  $K_{D,1}$  = 100 ± 8 nM, and  $K_{D,1}$  = 52 ± 6 nM for eVP35 IID CBP3mut, F239A, and R312A, respectively (Figure 4D–F). Interestingly, the mutation of K248A in combination with the CBP3mut leads to a ~25-fold increase in  $K_{D,1}$  compared to eVP35 IID WT (Figure 4G) but does not abolish binding altogether. Thus, in contrast to 1G8-14, residues in the FBP are critical for high affinity 2F11-14 binding. Similar to the results observed for 1G8-14 binding, residues in the CBP contribute to the second binding event, presumably through contacts with double stranded regions of the 2F11-14 aptamer. Our results are summarized in Table 2.



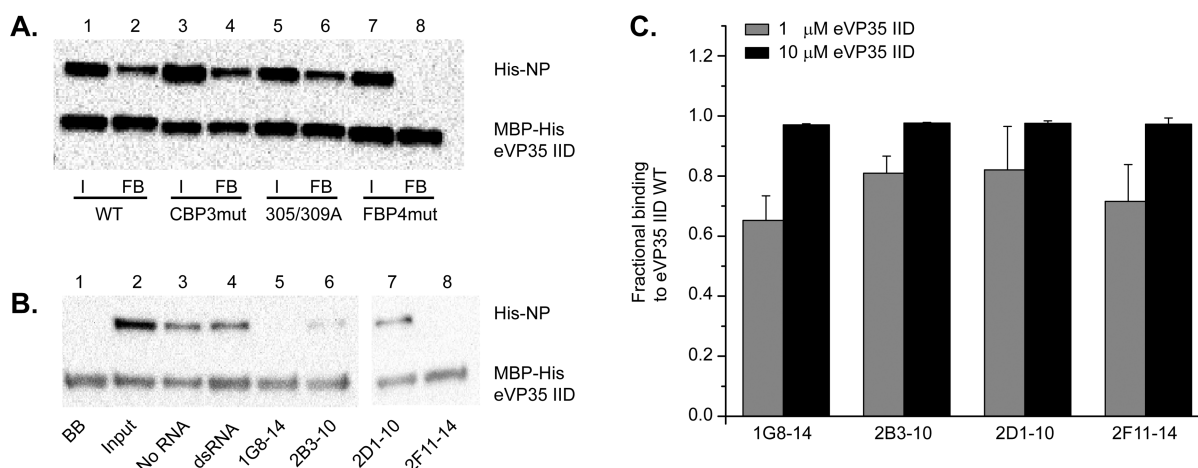


**Figure 4.** FBP is important for high affinity binding of the 2F11-14 aptamer to eVP35 IID. ITC raw data and corresponding binding isotherms for 2F11-14 binding to eVP35 IID (A) WT, (B) K248A, (C) FBP4mut, (D) CBP3mut, (E) F239A, (F) R312A, and (G) CBP3mut + K248A.

In order to better define the binding interface for eVP35 IID-1G8-14 and eVP35 IID/2F11-14, and how these aptamers are oriented with respect to eVP35 IID, we used a filter binding competition assay to see if 1G8-14 could compete with 2F11-14 for eVP35 IID binding and vice versa. As expected,

unlabeled 1G8-14 was able to compete off  $^{32}$ P-labeled 1G8-14. Unlabeled 1G8-14 can also compete off  $^{32}$ P-labeled 2F11-14 (Figure S9, Supporting Information). However, unlabeled 2F11-14 could only compete off  $^{32}$ P-labeled 2F11-14 but not  $^{32}$ P-labeled 1G8-14. Since 1G8-14 was selected against WT





**Figure 5.** 1G8-14 and 2F11-14 aptamers disrupt eVP35 IID-NP interaction. MBP-His tagged WT or mutant eVP35 IIDs were immobilized on amylose resin and incubated with His-tagged NP proteins in the (A) absence or (B) presence of dsRNA or aptamer. (A) Lanes 1, 3, 5, and 7 are input (I) samples for MBP-His eVP35 IID WT, CBP3mut, 305A/309A, and FBP4mut, respectively. Lanes 2, 4, 6, and 8 are final bead (FB) samples for MBP-His eVP35 IID WT, CBP3mut, 305A/309A, and FBP4mut, respectively. (B) Lane 1 shows MBP-His tagged eVP35 IID WT bound to the amylose resin. Lane 2 shows the presence of both His-NP and MBP-His eVP35 IID WT bound to resin, and lanes 3–8 are final bead samples for pull-downs done in the absence (lane 3) or presence of dsRNA (lanes 4) and aptamers 1G8-14 (lane 5), 2B3-10 (lane 6), 2D1-10 (lane 7), and 2F11-14 (lane 8). (C) eVP35 IID WT binds aptamers 1G8-14, 2B3-10, 2D1-10, and 2F11-14 at protein concentrations of 1  $\mu$ M (gray), and 10  $\mu$ M (black).

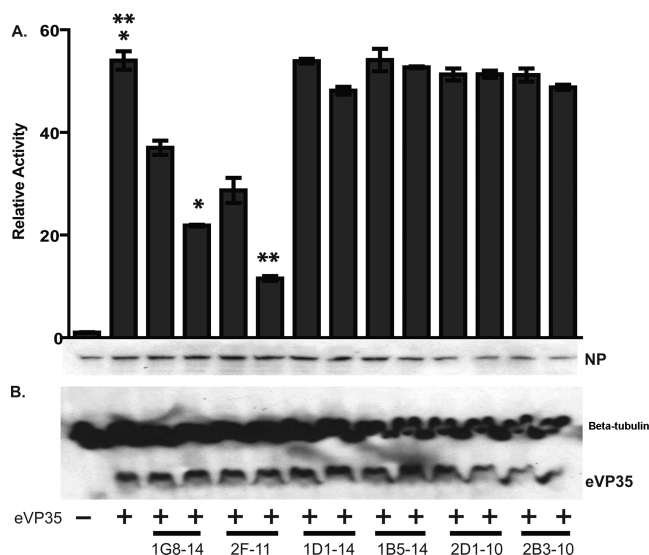
eVP35 IID, it is conceivable that the 1G8-14 aptamer would make contacts with both basic patches. This notion is supported by our ITC data (described above), showing that the CBP is the primary binding site for 1G8-14 but that the FBP also contributes to 1G8-14 binding and that mutations to a CBP residue in combination with mutations to the FBP region (FBP4mut + R312A) results in the loss of interaction. In contrast, the 2F11-14 aptamer, which was selected against a mutant protein with only the FBP present, may predominantly interact with the FBP surface. Altogether, these data are consistent with the notion that distinct classes of high affinity aptamers can be developed using eVP35 IID WT and mutant proteins as SELEX targets.

**Primer Binding Region of 1G8-14 Is Important for High Affinity VP35 IID Binding.** The structure predictions for the 1G8-14 aptamer (Figure S4, Supporting Information) suggested that the primer binding regions may be important for secondary structure. To further evaluate the contributions from secondary structure of the 1G8-14 primer regions, we performed binding site boundary measurements to identify the nucleotides of 1G8-14 important for binding to eVP35 IID (Figure S10, Supporting Information). The resulting data suggests that the region between nucleotides 36 and 65 is important. However, given the nature of the experiment, it was not clear if this is due to binding of the structural element shown in Figure S10B (Supporting Information) or if binding was primarily driven by the sequence between nucleotides 36 to 65 (indicated by arrows in Figure S10C, Supporting Information). In this assay, we used partially digested aptamers under native conditions and isolated aptamer fragments that bound eVP35 IID WT. RNA bound eVP35 IID proteins were subsequently subjected to denaturing gel electrophoresis for sequence mapping. In order to differentiate between the two potential aptamer fragments that form the high affinity binding, we generated the minimum RNA sequence (nucleotides 36–65) and subjected that sequence to structure predictions and binding studies. In addition, we also tested RNA fragments corresponding to 1–65 and 36–90 nucleotides, and the

minimal free energy (MFE) structure predictions are shown in Figure S11 (Supporting Information). These predictions revealed that bases 36–65 form a stem-loop, which is not a substructure predicted for the full-length 1G8-14 structure (compare Figure S11A with Figure S11B, Supporting Information). We performed ITC experiments in order to compare the full-length and truncated 1G8-14 constructs in their ability to bind eVP35 IID WT and mutant proteins. Truncated 1G8-14 36–65 bound to eVP35 IID WT with a two-site binding mode similar to that of full-length 1G8-14. However, the truncated 1G8-14 36–65 aptamer was unable to bind eVP35 IID R312A (Figure S12, Supporting Information; see Figure S12F vs Figure S12C, Supporting Information). The sensitivity to the eVP35 IID R312A mutation was also observed previously for dsRNA binding. We also conducted ITC experiments for truncated 1G8-14 1–65 and 1G8-14 36–90 to assess the importance of the 5' constant region and 3' constant region, respectively (Figure S13, Supporting Information). Truncated 1G8-14 1–65 bound eVP35 IID WT with a two-site binding mode but similar to truncated 1G8-14 36–65 was unable to bind eVP35 IID R312A. Additionally, truncated 1G8-14 1–65 binding to eVP35 IID WT appears to be cooperative, and this observation will require additional experiments in order to fully explore this behavior. However, it is evident that there are significant differences between WT and the 1–65 truncated aptamer. In contrast, truncated 1G8-14 36–90 bound eVP35 IID WT with a two-site binding mode similar to that of the full-length aptamer, and more importantly, this truncated aptamer bound eVP35 IID R312A with high affinity (Figure S13I, Supporting Information). Together, these data suggest that the variable region of 1G8-14 is important for binding to eVP35 IID, but additional contributions from the constant regions are required for the optimal aptamer formation. On the basis of these observations, we used full-length aptamers in our studies described here.

**Aptamers Disrupt eVP35 IID–NP Interactions and Inhibit EBOV Replication/Transcription Activity.** In order to test the utility of the 1G8-14 and 2F11-14 aptamers, we

assessed their ability to inhibit interactions between eVP35 IID and NP proteins. This interaction is important for the function of the filoviral replication complex.<sup>23,31</sup> Specifically, residues in the FBP of eVP35 IID are important for eVP35 IID–NP interactions and minigenome activity (Figure 5A).<sup>23</sup> Mutation of all four residues in the FBP (eVP35 IID FBP4mut) leads to loss of eVP35 IID–NP binding in a pulldown assay. In contrast, mutation of residues in the CBP (eVP35 IID R305A/K309A and eVP35 IID CBP3mut) does not impact eVP35–NP binding. Since the FBP appears to be important for both 1G8-14 and 2F11-14 binding, we tested the ability of these aptamers to compete with NP for binding to eVP35 IID. Both 1G8-14 and 2F11-14 aptamers were able to inhibit eVP35 IID–NP complex formation, while dsRNA and aptamers 2B3-10 and 2D1-10 are unable to disrupt the eVP35 IID–NP interaction (Figure 5B). Aptamers 2B3-10 and 2D1-10, selected against eVP35 IID CBP3mut, are of similar length to 1G8-14 and 2F11-14 aptamers, and they display similar high affinity binding to eVP35 IID (Figure 5C). Together, our data demonstrate the ability of 1G8-14 and 2F11-14 aptamers to specifically disrupt the eVP35–NP interaction (Figure 6). We also tested the



**Figure 6.** 1G8-14 and 2F11-14 aptamers inhibit EBOV replication/transcription activity in a dose dependent manner. A replication/transcription assay was performed in which plasmids encoding the different aptamers (250, 500 ng, left and right, respectively) were cotransfected with the plasmids required for minigenome replication and transcription (plasmids expressing EBOV L, NP, VP30 and VP35 along with plasmid expressing T7 promoter dependent EBOV minigenome, which encodes a Renilla luciferase reporter gene). Also cotransfected was a plasmid that expresses firefly luciferase from an RNA polymerase II promoter. Renilla luciferase activity was normalized to firefly luciferase activity. Minigenome reporter activation was expressed as relative activity by setting the negative control (without VP35) to a value of 1. The error bars indicate standard deviation of three independent replicates, \* $p = 0.001$  and \*\* $p = 0.0005$ . The Western blot shows the expression of NP and eVP35.

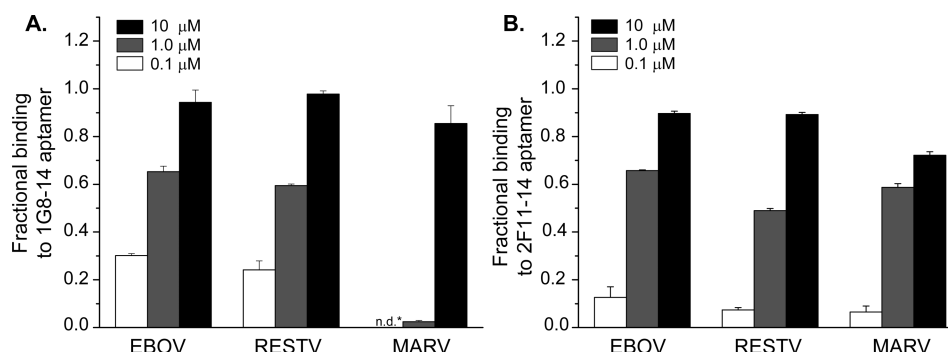
ability of aptamers 1G8-14 and 2F11-14 to inhibit RNA synthesis in a replication/transcription assay, where the viral replication complex was reconstituted by expression of select viral proteins.<sup>31</sup> Resulting data show that 1G8-14 and 2F11-14 can antagonize the replication/transcription activity in a dose dependent manner (Figure 6). In contrast to these two specific aptamers, additional aptamers with similar length and affinity

show little to no inhibition, despite all six aptamers being present at comparable levels as judged by qRT-PCR (Figure S14, Supporting Information). Altogether, these results demonstrate that 1G8-14 and 2F11-14 aptamers can potentially function as competitive inhibitors of the filoviral replication complex by disrupting eVP35 IID–NP interactions. However, it is worth noting that the aptamers can interact with NP at high protein concentrations ( $>100 \mu\text{M}$ ); thus, it is not clear if additional rounds of aptamer optimization with negative selection against NP will be necessary prior to the use of these aptamers in virally infected cells.

**Aptamers Differentially Recognize Closely Related Filoviral VP35 IID Proteins.** Filoviral VP35 proteins share high sequence similarity in the IID region (97.6% similar and 88.9% identical for RESTV aligned with EBOV; 81.7% similar and 22.9% identical for MARV aligned with EBOV) (Figure S15, Supporting Information). Therefore, we tested the ability to the 1G8-14 and 2F11-14 aptamers to differentiate between closely related VP35 proteins. The resulting data showed that the 1G8-14 aptamer preferentially binds to eVP35 IID proteins (EBOV and RESTV VP35 IID proteins) (Figure 7A). When binding was tested at protein concentrations  $<1 \mu\text{M}$ , EBOV and RESTV showed a fractional binding  $>0.6$ , while  $<0.1$  fractional binding was observed for MARV. In contrast, the 2F11-14 aptamer bound to all three filoviral VP35 IID proteins with  $>0.5$  fractional binding at protein concentrations  $>1 \mu\text{M}$ . (Figure 7B). The ability of 1G8-14 to differentiate between eVP35 IID and MARV VP35 IID proteins is likely due to sequence differences between EBOV and MARV VP35 proteins near the CBP. For example, in EBOV VP35 IID positions 319 and 322 are K319 and R322, respectively, while corresponding positions in MARV VP35 IID are T309 and K312 (Figure 2 and Figure S15, Supporting Information). Thus, it appears that the relative sensitivity to different filoviral species is entirely dependent on the relative contributions of the different basic patches.

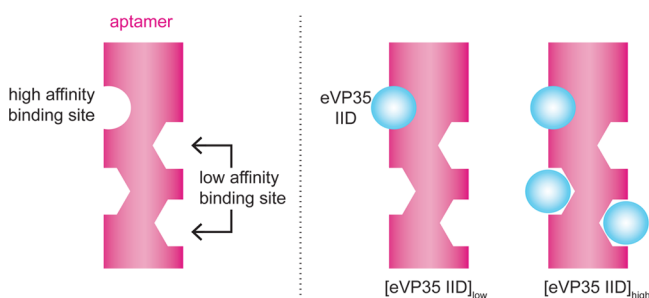
## DISCUSSION

In this study, we have identified and characterized two aptamers against eVP35 IID that bind with low nanomolar affinity using SELEX technology. These aptamers were generated by using two targets, eVP35 IID WT and eVP35 IID CBP3mut, with the goal of identifying different classes of aptamers that interact with distinct sets of residues within filoviral VP35 proteins. Target selection was in large part motivated by available structure–function data, and our goal was to drive the selection process to different functionally important regions of this multifunctional nonenzymatic protein. On the basis of the data described above, our selected aptamers display binding affinities that are 20–100-fold higher than dsRNA and the 92 nucleotide long random RNA pool that was used in the initial SELEX, supporting the specific enrichment of aptamers. From the initial groups of aptamers, 1G8-14 and 2F11-14 aptamers were chosen to represent two aptamer classes as they bound eVP35 IID with high affinity, disrupted the eVP35IID–NP complex, and displayed significant differences at the nucleotide level ( $\sim 40\%$  difference in the aptamer variable region). Biochemical and functional mapping of select aptamers revealed that 1G8-14 and 2F11-14 aptamers bind eVP35 IID primarily through interacting with residues from the CBP and the FBP, respectively, although it is apparent that additional interactions between eVP35 IID and aptamers are also formed due to the highly basic nature of the eVP35 IID surface.<sup>30</sup> Our ITC data



**Figure 7.** Aptamers differentially bind to filoviral VP35 IID proteins. eVP35 IID from EBOV, RESTV, and MARV were tested in their ability to bind (A) 1G8-14 and (B) 2F11-14 aptamers at varying protein concentrations shown as fractional binding.

revealed two independent binding modes: a high affinity aptamer binding mode and a the dsRNA binding mode (Figure 8). Each aptamer has a single high affinity binding site and



**Figure 8.** 1G8-14 and 2F11-14 aptamers have multiple binding modes. VP35 aptamers display two binding modes that include a single high affinity binding site and multiple dsRNA binding sites. At low eVP35 IID WT concentrations or when binding to key dsRNA binding mutant R312A, only the high affinity site is occupied. At higher concentrations of eVP35 IID WT, additional binding events occur via the dsRNA binding mode.

multiple dsRNA binding sites. These two binding events are energetically independent, at least in the context of eVP35 IID. Interestingly, mutations to CBP residues or key residues for dsRNA binding resulted in the presumable loss of the dsRNA binding mode. Therefore, at low eVP35 IID concentrations, we observe the high affinity aptamer binding event with a eVP35 IID:aptamer ratio of 1:1. As we increase eVP35 IID concentrations, eVP35 IID is able to make additional interactions with double stranded regions of the aptamers due to its inherent dsRNA binding function, therefore, allowing multiple eVP35 IID proteins to interact with a single aptamer.

This work demonstrates how integrating available structural and functional data to tailor SELEX selection can result in aptamers that target different functional regions and potentially generate multivalent aptamers. The 1G8-14 aptamer was selected against eVP35 IID WT and binds primarily to residues in the CBP with additional contacts to the FBP region. Despite the fact that the CBP is important for both 1G8-14 aptamer and dsRNA binding, there are significant differences in how eVP35 IID binds these two ligands. Most notably, a single point mutation within the CBP completely abolishes dsRNA binding.<sup>16,18,28</sup> However, single point mutations have only marginal effects on 1G8-14 aptamer binding (Figure 2). In contrast, double and triple mutation of CBP residues significantly affect the ability of the 1G8-14 aptamer to bind

eVP35 IID, which supports the notion that the aptamers establish multiple contact sites at the binding interface(s). In addition to the CBP, our data shows that mutations within the FBP affect 1G8-14 aptamer binding (increases  $K_{D,1}$ ). Mutation of the FBP along with a single mutation in the CBP region abolishes aptamer binding altogether. These results further support the view that the 1G8-14 aptamer recognizes several surfaces across eVP35 IID. In contrast, the main site of interaction on eVP35 IID for the 2F11-14 aptamer appears to be near the FBP, with the CBP providing additional contact points. Our results clearly suggest that both 1G8-14 and 2F11-14 aptamers interact with multiple copies of eVP35 IID and that therefore there is a high potential for these aptamers to inhibit multiple VP35-mediated functions *in vivo*.

In this work, we were able to target the CBP or the FBP of eVP35 IID and show that 1G8-14 and 2F11-14 aptamers have distinct binding footprints through biochemical mapping. Moreover, we are able to distinguish that the CBP is the primary binding site for 1G8-14, while 2F11-14 uses the FBP as its primary binding site. As a multifunctional protein, eVP35 IID inhibits IFN responses to viral infection, in part through dsRNA sequestration, and supports RNA viral synthesis through structural bridging of NP and the viral polymerase L. We have shown that aptamers bind eVP35 IID using multiple points of contact, a mode that is distinct from dsRNA binding. Furthermore, aptamers can compete with dsRNA for eVP35 IID binding, indicating that aptamers have the potential to disrupt the IFN inhibitory function of eVP35. Here, we show that aptamers disrupt the eVP35–NP interaction and replication/transcription activity, indicating their potential to inhibit eVP35 polymerase cofactor function. It is interesting to note that selected aptamers can potentially inhibit both the eVP35 immune suppressor function and the polymerase cofactor function.

Aptamers are well known for their high affinity and specificity against their targets. The high specificity of aptamers can be used to distinguish among viral subtypes, which is relevant for the development of aptamers against viral proteins. However, aptamers with broad specificity can potentially serve as pan-antivirals. One of our selected aptamers, 1G8-14, can distinguish between EBOV and MARV VP35 IIDs, binding to EBOV and RESTV VP35 IIDs with  $K_D$ 's 20-fold lower (at 1 μM concentration of protein; see Figure 5A) than that of MARV VP35 IID. The other selected aptamer, 2F11-14, can bind EBOV, RESTV, and MARV VP35 IIDs with comparable affinities. Thus, we have demonstrated that it is feasible to develop both highly specific and pan-antifiloviral aptamers using VP35 as the SELEX target molecule, and our success in



selecting both highly specific and pan-antifiloviral aptamers indicates that there are highly conserved regions of VP35, which are recognized by 2F11-14, as well as species unique regions in VP35, which allows differential binding of 1G8-14 to EBOV and MARV VP35 IIDs.

Filoviruses cause high rates of fatalities during outbreaks and are potential agents of bioterrorism. Although filoviral vaccine options are under investigation, no approved vaccines are available at this time. Furthermore, other therapeutics to counter disease are largely lacking at present. Work here, using available structure–function data, supports the development of VP35 specific aptamers that can facilitate validation of multifunctional VP35 as an antiviral target. Thus, inhibition of VP35 function using aptamers will not only provide a path to develop antivirals but also provide novel target sites and mechanisms for antiviral development.

## ■ ASSOCIATED CONTENT

### ■ Supporting Information

Data collection, structure solution, and refinement statistics for eVP35 IID CBP3mut and eVP35 IID FBP4mut; alignment of crystal structures and electrostatic surface representation for zVP35 IID WT, CBP3mut, and FBP4mut; enrichment of aptamer populations using SELEX; secondary structure prediction for 1G8-14 and 2F11-14; CD spectra show that 1G8-14 and 2F11-14 have different secondary structures; aptamer binding is not affected by the 5′-ppp moiety; mutations in the FBP of zVP35 IID result in overall lower maximum fractional binding; aptamers compete with dsRNA for eVP35 IID binding; 1G8-14, but not 2F11-14, can compete with 1G8-14 for binding to eVP35 IID; determination of the minimal binding region of 1G8-14 with eVP35 IID; structure predictions and binding studies for truncated 1G8-14; similar amounts of aptamer RNA in transfected cells as judged by qRT-PCR; and sequence alignment of EBOV, RESTV, and MARV IIDs. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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### Notes

The content of the information does not necessarily reflect the position or the policy of the federal government, and no official endorsement should be inferred.

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## ■ ABBREVIATIONS

EBOV, Ebola virus; RESTV, Reston virus; MARV, Marburg virus; IFN, interferon; IID, IFN inhibitory domain; VP35, viral protein 35; eVP35 IID, EBOV VP35 IID; RLRs, RIG-I-like receptors; CBP, central basic patch; FBP, first basic patch; SELEX, systematic evolution of ligands by exponential enrichment; NP, nucleoprotein; ITC, isothermal titration calorimetry; dsRNA, double stranded RNA; IKKε, IkappaB kinase epsilon; TBK-1, TANK-binding kinase 1; PKR, RNA-dependent protein kinase; CD, circular dichroism; TCEP, tris(2-carboxyethyl)phosphine; HRP, horseradish peroxidase; MFE, minimum free energy; RMSD, root-mean-square deviation

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