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dsRNA binding characterization of full length recombinant wild type and mutants Zaire ebolavirus VP35

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ABSTRACT

The Ebola viruses (EBOVs) VP35 protein is a multifunctional major virulence factor involved in EBOVs replication and evasion of the host immune system. EBOV VP35 is an essential component of the viral RNA polymerase, it is a key participant of the nucleocapsid assembly and it inhibits the innate immune response by antagonizing RIG-I like receptors through its dsRNA binding function and, hence, by suppressing the host type I interferon (IFN) production. Insights into the VP35 dsRNA recognition have been recently revealed by structural and functional analysis performed on its C-terminus protein. We report the biochemical characterization of the Zaire ebolavirus (ZEBOV) full-length recombinant VP35 (rVP35)–dsRNA binding function. We established a novel in vitro magnetic dsRNA binding pull down assay, determined the rVP35 optimal dsRNA binding parameters, measured the rVP35 equilibrium dissociation constant for heterologous in vitro transcribed dsRNA of different length and short synthetic dsRNA of 8 bp, and validated the assay for compound screening by assessing the inhibitory ability of auryntri-carboxylic acid (IC50 value of 50 μg/mL). Furthermore, we compared the dsRNA binding properties of full length wt rVP35 with those of R305A, K309A and R312A rVP35 mutants, which were previously reported to be defective in dsRNA binding-mediated IFN inhibition, showing that the latter have measurably increased Kd values for dsRNA binding and modified migration patterns in mobility shift assays with respect to wt rVP35. Overall, these results provide the first characterization of the full-length wt and mutants VP35–dsRNA binding functions.

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1. Introduction

Ebolaviruses (EBOVs) constitute a group of five species of filamentous, enveloped, non-segmented single- and negative-stranded RNA viruses belonging to the family of Filoviridae, order of Mononegavirales (Barrette et al., in press; Kuhn et al., 2010). EBOVs are causative agents of highly lethal hemorrhagic fevers in humans and non-human primates and they are classified as category A pathogens, for which treatments and vaccination are currently lacking (Falzarano et al., 2011; Feldmann and Geisbert, 2011). Although human outbreaks are rare and generally confined to sub-Saharan African regions, the risk of infection due to travel-imported cases or their misuse as bioterrorism agents place EBOVs among the highest public health threats (Hartman et al., 2010; Leroy et al., 2011). In 2008, the discovery in the Philippines of the presence of viruses belonging to the Reston ebolavirus (REBOV) species in infected domestic swine, gave rise to worldwide concern due to EBOVs theoretical capability of entering the human food chain (Barrette et al., 2009). More recently, susceptibility of pigs to EBOVs infection was assessed, showing that the virus is able to replicate with high titers in their respiratory tract and to shed to naïve hosts from the oronasal mucosa (Kobinger et al., 2011). Furthermore, the EBOVs capability of inducing disease following aerosol inhalation exposure has been recently demonstrated for three species of non-human primates (Reed et al., 2011).

In humans, EBOV hemorrhagic fever often results in very high mortality rates, reaching 90% for the most lethal species such as Zaire ebolavirus (ZEBOV). As recently observed in ZEBOV human fatal cases (Wauquier et al., 2010), as well as in previously described studies with experimentally infected animal models, death is associated with a markedly impaired innate immunity reaction, a strong production of pro-inflammatory cytokines and a profound immunosuppression, resulting in peripheral T lymphocyte apoptosis and lack of adaptive immunity (Mahanty and Bray, 2004; Mohamadzadeh et al., 2007; Zampieri et al., 2007). By contrast, survivors to EBOVs infection seem to develop an effective immune response (Becquart et al., 2010; Wauquier et al., 2009). This disparity suggests that events early in the EBOV infection may influence the patients’ ability to activate an effective immune response (Basler and Amarasinghe, 2009; Mohamadzadeh, 2009).

In general, pathogen associated molecular patterns (PAMPs) are recognized, early in infection, by PAMP recognition receptors (PRRs) (Randall and Goodbourn, 2008) which activate the host
in innate immune response (Koyama et al., 2008). Among PAMPS, dsRNA is a unique viral product, very effectively detected by cellular PRRs such as retinoic acid-inducible gene 1 (RIG-I) and melanoma differentiation-associated gene 5 (MDA-5) proteins (Loo and Gale, 2011; Yoneyama and Fujita, 2010) which, once recognized, initiate signaling cascades that activate interferon regulatory factor 3 (IRF-3), leading to production of interferon α/β (IFN-α/β) which, in turn, activates the antiviral responses (Sadler and Williams, 2008). To circumvent viral PAMPs recognition by cellular PRRs, viruses evolved different strategies to block IFN-α/β activation (Bowie and Unterholzner, 2008; Conzelmann, 2005; Katze et al., 2008). In particular, EBOVs code for two viral proteins, VP24 and VP35, which display innate immune antagonism (Basler and Amarasinghe, 2009).

VP35 is a multifunctional protein that is indispensable for EBOVs replication. In fact, VP35 is an essential cofactor of the EBOV RNA polymerase complex (Boehmann et al., 2005; Mühlberger et al., 1998, 1999), it serves as a viral assembly factor (Huang et al., 2002; Johnson et al., 2006), it is a RNAi suppressor in mammalian cells (Haasnoot et al., 2007; Fabozzi et al., 2011) and, moreover, it counteracts the host innate immune response by blocking, at many different steps, the production of IFNs-α/β. In this respect, it has been reported that VP35 (i) binds to dsRNA, suppressing RIG-I helicase signaling cascade triggered by dsRNA recognition (Cárdenas et al., 2006); (ii) impairs IKK-α and TBK-1 kinases functionality (Prins et al., 2008); (iii) enhances PIAS1-mediated SUMOylation of IRF-7, decreasing IFN induction (Chang et al., 2009); (iv) abrogates IRF-3 phosphorylation–imerization and nuclear translocation (Basler et al., 2000, 2003; Hartman et al., 2006, 2008a) and (v) inhibits dsRNA-induced activation of PKR (Feng et al., 2007; Schümann et al., 2009). In particular, among VP35 functions, dsRNA binding seems to be the most important in the context of EBOV pathogenesis, since mutations in its dsRNA binding domain (RBD) (i) abolish its IFN-antagonism properties (Cárdenas et al., 2006; Hartman et al., 2004); (ii) result in greatly attenuated viral growth rate and virulence loss (Hartman et al., 2006, 2008a; Prins et al., 2010); (iii) abrogate VP35 capability to suppress RNA silencing (Haasnoot et al., 2007).

VP35 contains an N-terminal coiled-coil domain that is important for its homo-oligomerization (Möller et al., 2005; Reid et al., 2005) and a C-terminal RBD (Cárdenas et al., 2006; Hartman et al., 2004; Leung et al., 2009, 2010b,c). The coiled-coil domain is required for several VP35-mediated functions such as viral replication (Möller et al., 2005; Reid et al., 2005) nuclecapsid formation (Huang et al., 2002; Johnson et al., 2006; Shi et al., 2008) and immune suppression (Feng et al., 2007; Jin et al., 2010; Reid et al., 2005), while the VP35 RBD is involved in dsRNA binding, mainly linked to immune suppression (in fact, at our best knowledge, the VP35–dsRNA binding properties seems to be not involved in the replication complex function), even though, when the sole VP35 RBD is expressed into cells, it is not able to suppress the type I IFN activation at the same level than the full length protein (Basler and Amarasinghe, 2009; Cárdenas et al., 2006; Feng et al., 2007; Reid et al., 2005). Currently, the complete tertiary structure of the full length EBOV VP35 is not yet available. However, the crystallographic resolution of the RBD C-terminal portion (amino acid residues 221–340) has been recently solved for the ZEBOV and REBOV species, either alone, or bound to short dsRNA molecules (Kimberlin et al., 2010; Leung et al., 2009, 2010c,d). EBOV VP35 RBD presents a unique fold among the dsRNA binding proteins, different from the αβββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββββbeta
protocols, other two linearized DNA templates to in vitro transcribe the 150 and 50 bp dsRNA as above. All in vitro transcribed (IVT) dsRNA molecules were purified from transcription reaction with the MEGAClear kit (Ambion) or with Quick Spin G25 columns (Roche), and quantified by spectrophotometry. dsRNA oligos of 8 bp in length (sequence 5′-GCGUACGC-3′) bearing 5′ phosphate and 5′ hydroxyl ends were purchased from Metabion International AG (Germany). The integrity of DNA templates, IVT dsRNAs and synthetic dsRNA molecules was assessed by agarose-gel electrophoresis.

2.4. Electrophoresis mobility shift assay (EMSA)

Purified IVT dsRNA molecules of different length were incubated at a concentration of 40 nM with 0.3 μg rVP35 in a reaction volume of 50 μL containing 50 mM sodium phosphate pH 7.5, 100 mM NaCl, 20 mM MgCl2 and 1 μg tRNA for 30 min at 23 °C under gentle rotating agitation (20 rpm). Five microliters of 6× non-denaturing gel loading buffer (37% glycerol, 0.025% bromophenol blue, 0.025% xylene cyanol, 20 mM Tris–HCl pH 8.0) were then added to the reaction and the samples were loaded on a 1.7% non-denaturing agarose gel. Electrophoresis was allowed to proceed in 1X TBE running buffer at 4 °C for 2.5 h at a constant voltage (5.5 V/cm). Gels were stained with 1X SYBR Green II RNA gel stain (Invitrogen) and migration profiles of reaction products were visualized against a fixed amount of labeled 500 bp3H-dsRNA (0.1 Ci/mL). In both cases, increasing amounts of unlabeled dsRNA were titrated against a fixed amount of labeled 500 bp 3H-dsRNA (0.1 Ci/mmoles). Raw data from at least three independent binding experiments were used to calculate Kd value utilizing non-linear regression fitting model provided by Prism software (GraphPad).

2.5. Magnetic pull down assay

The rVP35–dsRNA complex formation was assessed exploiting the properties of the TALON paramagnetic Dynabeads (Invitrogen). Firstly, 1 μg of rVP35 was conjugated to 50 μL TALON beads in a volume of 700 μL of binding buffer (50 mM sodium phosphate pH 7.5, 150 mM NaCl, 0.05% Tween-20) for 15 min at 23 °C under gentle rotating agitation (20 rpm). Unconjugated rVP35 was removed by magnetic field application, supernatant removal and further washing with binding buffer. Pellets with conjugated rVP35 were re-suspended in a 100 μL volume binding buffer containing 20 mM MgCl2 and 1.5 mM 500 bp 3H-dsRNA (0.1 Ci/mmoles) and incubated for 60 min at 37 °C (20 rpm). Unbound 3H-dsRNA was separated by the conjugated rVP35–dsRNA complex by magnetic field application and supernatant removal. A further washing step was performed to completely remove unbound 3H-dsRNA. 3H-dsRNA elution was performed by incubation of the pellets in 300 μL elution buffer (binding buffer plus 1 M imidazole pH 7.5) for 10 min at 23 °C (20 rpm), subsequent magnetic field application and supernatant removal. The supernatant was transferred to vials and its radioactivity determined with a Beckman LS 6500 beta-counter.

2.6. Determination of biochemical parameters for optimal dsRNA binding

The determination of the equilibrium dissociation constants (Kd), was performed by competition binding studies using the homologous competition approach for the 500 bp dsRNA and the heterologous competition for the 150 and 50 bp dsRNAs, as described (Bylund and Murrin, 2000; Motulsky and Neubig, 2010). In both cases, increasing amounts of unlabeled dsRNA were titrated against a fixed amount of labeled 500 bp 3H-dsRNA (0.1 Ci/mmoles). Raw data from at least three independent binding experiments were used to calculate Kd value utilizing non-linear regression fitting model provided by Prism software (GraphPad).

2.7. Thermodynamic Van’t Hoff analysis

The effect of temperature on rVP35 binding affinity for dsRNA was evaluated using a comparative thermodynamic Van’t Hoff analysis, as described (Chung et al., 2010). Briefly, triplicate 500 bp dsRNA homologous–competition binding experiments were conducted at 23 °C, 30 and 37 °C and titration curves were generated for Kd determination. The Kd values obtained at each temperature were plotted as ln(Kd) versus 1/temperature (T; expressed in Kelvin), and the simplified Van’t Hoff equation ln(Kd) = [(AH/R) – (1/T)] – AS/R was used to estimate enthalpy and entropy, by assuming that these two parameters are constant within the temperature range assayed. ln(Kd) is the natural log of the equilibrium dissociation constant, AH and AS are the enthalpy and entropy and R is the molar gas constant (8.314 kJ·mol⁻¹).

3. Results

3.1. rVP35 shifts the electrophoretic migration of an heterologous 500 bp IVT dsRNA

It has been previously shown that full length EBOV VP35 expressed in mammalian cells can bind to poly-IC, poly-AU as well as IVT dsRNA molecules ranging from 200 to 1000 bp (Cárdenas et al., 2006). When truncated, E. coli-expressed VP35 proteins were tested, different dsRNA binding abilities were observed. In fact, while a VP35 truncated version containing the amino acid residues 169–340 was still able to bind poly-IC dsRNA (Cárdenas et al., 2006), the protein containing only the 201–340 amino acid residues was not able to co-precipitate with the poly-IC substrate (Feng et al., 2007). However, more recently, a shorter VP35 truncated protein containing the amino acid residues 221–340 was found to bind dsRNA as short as 8 bp in length (Kimberlin et al., 2010; Leung et al., 2009, 2010a,b; Prins et al., 2010). Within this context, we previously reported the expression and purification of a full-length rVP35 protein and demonstrated that it co-precipitates with agarose-beaded poly-CG in a standard pull down assay, thereby assessing the rVP35 ability to bind dsRNA, similarly to the native VP35 (Zinzula et al., 2009). Therefore, in order to further characterize the full-length rVP35–dsRNA binding ability, we first asked whether the rVP35 protein could effectively bind to heterologous IVT dsRNA molecules obtained using a T7-promoter-driven DNA–plasmid. To answer this question, we performed an EMSA using IMAC purified ZEBOV rVP35 (Fig. 1A) and IVT dsRNAs.
showing that rVP35 binds to the 500 bp dsRNA and leads to a shift in its electrophoretic migration (Fig. 1B, compare lanes 3 to 4). In addition to the shifted dsRNA signal into the gel (second band from the top of the gel), a further fluorescent signal could also be observed stuck in the well of the gel, which was possibly ascribed to higher molecular mass rVP35–dsRNA specific aggregates (Fig. 1B, lanes 4 first band from the top). In this respect, it is important to note that (i) the migration of rVP35 alone was unable to produce any UV fluorescent signal in the gel (Fig. 1B, lane 2), therefore excluding the possible presence of bacterial RNA bound to rVP35; (ii) the addition of an excess of tRNA in the mixture did not alter the rVP35–dsRNA bands pattern (Fig. 1B, lane 5); (iii) heat-denatured rVP35 did not display any form of rVP35–dsRNA aggregates (Fig. 1B, lane 7). Moreover, EMSA performed using dsRNA substrates of 150 and 50 bp in length led to similar results (data not shown), in agreement with the length-independent modality proposed for VP35 binding to dsRNA (Kimberlin et al., 2010; Leung et al., 2010c). Altogether, EMSA experiments showed that bacterially expressed rVP35 is able to modify the migration profile of IVT dsRNAs of 50–500 bp in length. Noteworthy, even though the two distinct bands observed in the gel are probably due to specific rVP35–dsRNA complexes of different weight, their functional significance is still not determined. Overall, these data demonstrate the suitability of the heterologous IVT dsRNA molecules as effective substrates for quantitative studies on VP35–dsRNA binding.

3.2. rVP35 binds to a 3H-labeled 500 bp dsRNA in a magnetic pull down assay

With the aim of establishing a quantitative method to measure the rVP35 binding to IVT dsRNA we exploited the biochemical properties of the TALON Dynabeads which are paramagnetic beads that specifically bind His-tagged recombinant proteins. Purified rVP35 was first coated to Dynabeads through its N-terminal His-tag to form stable complexes of conjugated bead–proteins. The use of excess rVP35 assured that all beads were conjugated with rVP35 and the application of magnetic field to the solution allowed the washing out of the non-conjugated rVP35 (data not shown). The conjugated rVP35 was incubated with a 500 bp IVT 3H-dsRNA, allowing the binding to dsRNA. The unbound 3H-dsRNA was washed out by magnetic field application while the rVP35–3H-dsRNA complex was subsequently eluted with imidazole and quantified. Data showed that dsRNA was specifically retained by conjugated rVP35, in fact, the 3H-dsRNA binding to the empty beads (background signal) was very low as compared to the rVP35–3H-dsRNA binding (data not shown). Optimal binding conditions were determined to be at pH 7.5, 100 mM NaCl and 20 mM phosphate group.

It has been reported that, as measured by isothermal titration calorimetry (ITC), the RBD of VP35 (aminocids 221–340) binds to a synthetic 8 bp dsRNA with a $K_d$ value of 500 nM, while it binds to an IVT 8 bp dsRNA with higher affinity, showing a $K_d$ value of 30 nM (Leung et al., 2010c,d). Hence, we wanted to estimate the full length rVP35 equilibrium dissociation constant using more natural and longer substrates utilizing the magnetic pull down assay. Homologous competition binding experiments, in which the IVT 500 bp dsRNA, 3H-labeled and unlabeled, were used as ligand and competitor, respectively, were firstly performed (Fig. 3A). Data showed that the full-length rVP35 $K_d$ value for IVT 500 bp dsRNA binding at 37 °C temperature was 2.8 ± 0.1 nM.

Since our EMSA studies performed with IVT dsRNA of different length suggested that rVP35 could bind to IVT dsRNA of 500, 150 and 50 bp in length with a similar mode, we wanted to determine whether, despite this similarity, any difference would exist in the rVP35 binding affinity constant for IVT 500, 150 and 50 bp dsRNAs. Heterologous competition binding curves were performed at 37 °C temperature using 500 bp IVT 3H-dsRNA as ligand and unlabeled IVT 150 and 50 bp dsRNA molecules as competitors, and the $K_d$ values were calculated to be 2.4 ± 0.3 and 3.2 ± 0.5 nM, respectively (Fig. 3A). Furthermore, given that it was reported that the VP35 RBD is able to bind with high affinity dsRNA molecules as short as 8 bp in length, we also tested the full-length rVP35–dsRNA binding function towards such short substrates. To this end, heterologous competition binding curves were performed using 500 bp IVT 3H-dsRNA as ligand and two synthetic 8 bp dsRNA as unlabeled competitors. One of the 8 bp dsRNA competitors terminated with a 5′-phosphate, the other with a 5′-hydroxyl group. Data showed that rVP35 bound to the 5′-phosphate 8 bp dsRNA with high affinity, showing a $K_d$ value of 64 ± 9 nM, while it bound to the 5′-hydroxyl 8 bp dsRNA with a lower affinity. A $K_d$ value of 1.1 ± 0.2 μM. Overall, these data show that rVP35 binding affinity for IVT dsRNA in the magnetic pull down assay is independent from dsRNA length, at least for substrates ranging from 50 to 500 bp. In addition, rVP35 demonstrates a lower affinity for a 5′-phosphate 8 bp dsRNA, and a dramatic affinity reduction for a 8 bp dsRNA molecule lacking the 5′ phosphate group.

In the process of the magnetic pull down assay optimization, we observed that rVP35 binding ability to dsRNA was slightly dependent on temperature (data not shown). Therefore, we wanted to precisely determine the effect of the temperature on the rVP35 binding by carrying out homologous competition curves with IVT 500 bp dsRNA also at 30 and 23 °C (Fig. 4A). Results showed that, as temperature decreases, the rVP35 binding affinity for dsRNA increases. In fact, the rVP35 $K_d$ values for the binding to IVT 500 bp dsRNA were equal to 1.5 ± 0.5 and 1.1 ± 0.1 nM, at 30 and 23 °C, respectively, indicating that the rVP35–dsRNA binding at these temperatures does not change significantly. The effect of temperature on the rVP35–dsRNA binding affinity was also examined using the Van’t Hoff analysis which allows to determine $\Delta G$, $\Delta H$ and $\Delta S$ values (Chung et al., 2010). The final Van’t Hoff plot was linear (Fig. 4B) and the thermodynamic parameters were calculated to be $\Delta H = -14.9$ kcal/mol, $\Delta S = -009$ eu and $\Delta G = -12.12$ kcal/(mol K) at 37 °C. These values are in agreement with the thermodynamic parameters calculated for other dsRNA binding proteins such as the protein kinase PKR, which were reported to be from $-41.7$ to $-26.3$ kcal/(mol K) for $\Delta G$ and from $-21.1$ to $-7.3$ kcal/(mol K) for $\Delta H$ (Zheng and Bevilacqua, 2000).

3.4. rVP35–dsRNA interaction is inhibited by ATA

The EBOV VP35 RBD has a unique fold among the dsRNA binding proteins, which is different from the αβββββ-fold of the canonical cellular dsRNA binding proteins (Kimberlin et al., 2010; Leung et al., 2009, 2010c). Thus, given the important role of the VP35 inhibition of the IFN-α/β activation following EBOV infection for viral pathogenesis (Leung et al., 2010a), the VP35–dsRNA binding ability represents a promising therapeutic target for viral inhibition. Therefore, we wanted to validate our newly established magnetic pull down assay also as a method for the screening of molecules that may inhibit the interaction between dsRNA and VP35. In the absence of a specific VP35–dsRNA binding inhibitor, we used a known inhibitor of nucleic acid–protein interactions such as the ATA (Gosh et al., 2009; González et al., 1980) (Fig. 5). Results showed that ATA inhibited rVP35–dsRNA binding.
in a dose–response manner with an IC50 of 50 μg/mL. However, to be suitable for the identification of antivirals by screening compound libraries, a biochemical assay is required to be up-scalable to a medium or high-throughput system. To satisfy such requirement, we also tested the magnetic pull down assay in a 96-wells format, which is cost-effective and amenable of automation. In this scale-up, ATA was found to inhibit the rVP35–dsRNA binding with an IC50 of 61 μg/mL (data not shown), confirming that the rVP35 magnetic pull down assay is an effective in vitro biochemical assay for the screening of potential antiviral agents.

3.5. R305A, K309A and R312A rVP35 mutants display modified migration profiles on EMSA and bind to IVT 500 bp dsRNA with lower affinity with respect to wt rVP35

It has been reported that some amino acid residues in the RBD are important for dsRNA binding. In particular, the R312A mutation severely impaired the VP35–dsRNA binding ability and its IFN antagonist activity in cell culture, without affecting the VP35 function as part of the viral polymerase complex (Cárdenas et al., 2006; Hartman et al., 2004; Hartman et al., 2008a,b; Leung et al., 2009). More recently, the 221–340 amino acid R312A VP35 truncated protein was shown to lose the dsRNA binding ability, as measured by EMSA (Leung et al., 2009) and ITC analysis (Leung et al., 2010c). Similarly, the K309A mutation has also been reported to greatly impair the VP35–dsRNA binding ability and its IFN antagonist activity in cell culture, without altering the VP35 function in the viral polymerase complex (Cárdenas et al., 2006; Hartman et al., 2004; Leung et al., 2009) and the 221–340 amino acid K309A VP35 mutant was unable to bind dsRNA, as measured by EMSA analysis (Leung et al., 2009). In addition, the R305A mutation was recently reported to reduce by 3-fold the 221–340 amino acid VP35 truncated protein $K_d$ value for dsRNA without affecting the viral replication (Leung et al., 2010c). Within this picture, we were interested in assessing the impact of these mutations on the full-length rVP35–dsRNA binding ability. Therefore, we performed in vitro site-directed mutagenesis of the wt plasmid construct to express the R305A, K309A and R312A rVP35s in E. coli and purified these mutants as full length, recombinant, His-tagged proteins. Next, we assessed the ability of the R305A, K309A and R312A rVP35 mutants to bind to heterologous dsRNAs in EMSA studies. Using the IVT 500 bp dsRNA, the rVP35 mutants, when compared to wt rVP35, displayed a diminished ability to shift the dsRNA into the gel, even though the shifted band was only scarcely visible (Fig. 6A and B). Differently, all mutants maintained their ability to form the higher order aggregates, which were retained into the wells. Similar results were obtained using dsRNA substrates of 150 and 50 bp in length (data not shown).

Subsequently, given that the EMSA analysis was merely qualitative, we wanted to quantify the differences in dsRNA binding observed for the different rVP35 mutants using the magnetic pull down assay. Therefore, we performed homologous competition binding curve using the IVT 500 bp dsRNA as ligand at 37°C temperature for the three rVP35 mutants (Fig. 6C). Results showed that all three rVP35 mutants were able to bind the IVT 500 bp dsRNA but showed measurable increased $K_d$ values with respect to wt rVP35. In particular, the R305A rVP35–dsRNA binding affinity appeared to be only slightly lower than wt rVP35 affinity, showing a $K_d$ value dsRNA of 3.85 ± 0.6 nM (with a $p$ value equal to 0.0418 with respect to wt rVP35). Interestingly, the $K_d$ value for R309A rVP35 was 4.99 ± 1.0 nM (with a $p$ value equal to 0.0219 with respect to wt rVP35) and the $K_d$ value for R312A rVP35 was 10.60 ± 1.7 nM (with a $p$ value equal to 0.0015 with respect to wt rVP35). In addition, given that $K_d$ value for wt rVP35 binding to the 5'-phosphate 8 bp dsRNA was roughly 25-fold higher than the $K_d$ values observed for 50–500 bp dsRNAs, we wanted to assess whether a comparable decrease in dsRNA binding affinity could be observed for the R312A–rVP35 binding. As shown by heterologous
competition binding curve (Fig. 6D), the $K_d$ value calculated for the R312A-rVP35 binding to the 5'-phosphate 8 bp dsRNA was 1.76 ± 0.2 nM, around 166-fold higher than the value shown for the 500 bp dsRNA.

4. Discussion

At nearly 40 years from their identification, EBOVs are still highly infectious pathogens without available therapy and, despite the vast knowledge achieved in the last two decades on their biology and pathogenesis, the molecular basis for their extreme lethality remain to a large extent unraveled (Feldmann and Geisbert, 2011). However, the impairment of the innate immunity, and particularly the suppression of the IFN-α/β response, has been proved to be critical in supporting an efficient EBOV replication (Basler and Amarasinghe, 2009). Recently, the multifunctional virally-coded VP35 has been proposed to be the key determinant of the EBOVs virulence (Jin et al., 2010), and its ability to antagonize the IFN-α/β response has been ascribed mainly to its capacity to bind and sequestrate dsRNA, thereby suppressing RIG-I signaling cascade triggered by this viral nucleic acid (Cárdenas et al., 2006; Hartman et al., 2004, 2006). In fact, EBOVs bearing mutations in the VP35 dsRNA RBD, which reduce or suppress VP35–dsRNA binding, display greatly attenuate viral growth rate or are avirulent in animal models (Hartman et al., 2008b; Prins et al., 2010). Until now, all the
in vitro information on the VP35–dsRNA binding were referred to data obtained using a truncated version of the VP35 protein, including only the RBD, and dsRNA molecules as short as 8–18 bp [Kimberlin et al., 2010; Leung et al., 2009, 2010c,d]. Therefore, it was of interest to perform the first characterization of the full-length rVP35 binding to heterologous IVT dsRNA as long as 50–500 bp.

Data from the EMSA study we performed to demonstrate that the IVT dsRNA could be used for subsequent quantitative studies, clearly indicate that full length rVP35 binds to 50–500 bp IVT dsRNAs. rVP35 determined a shift in their mobility, forming two main VP35–dsRNA complexes regardless of the dsRNA length, which seem to be both specific. The precise composition of the two complexes is not known yet, however, it is worth to note that gel filtration studies performed on the full length VP35 suggest that VP35 forms dimers, tri-tetramers and higher order aggregates (Zinzula et al., 2009; Reid et al., 2005; Möller et al., 2005; Leung et al., 2010c). Therefore, it is possible to hypothesize that the VP35 dimers or tri-tetramers may bind dsRNA forming the low order complex that can be seen into the gel, while the VP35 high order aggregates are still able to bind IVT dsRNAs but are too big to enter the gel. The specificity of such interactions is also supported by the fact that a C-terminal His-tagged rVP35, when tested in the same EMSA, totally fails to shift the IVT dsRNA, forming neither low nor high order complexes (data not shown). Given that rVP35 C-terminal residues such as I340 and R339 have been reputed critical for stabilization of the RBD (Leung et al., 2010a,c), the presence of an His-tag at the C-terminus could be sufficient to create some hurdle in the interaction between VP35 and dsRNA. The functional role of such low and high order VP35–dsRNA complexes is not clear, however, the fact that the R305A, K309A and R312A rVP35 mutants, which reduced or abrogated the EBOV IFN-α/β inhibition (Leung et al., 2010c), display a diminished ability to form low order rVP35–dsRNA aggregates, while they still formed high order aggregates, may suggest that the low order VP35–dsRNA complex may have an important functional role in the VP35 IFN-antagonistic function.

The fact that the disruption of IFN-α/β production is essential for the efficient EBOV propagation in vivo and that the loss of VP35–dsRNA binding capability strongly correlates with the loss of EBOV virulence, highlights the importance of VP35–dsRNA binding as an attractive and promising target for antiviral development. Therefore, we established a novel, straightforward and reproducible biochemical in vitro assay to measure the VP35–dsRNA binding, we used it to quantitatively characterize this interaction and validated it as a tool for compound screening. Firstly, we showed that full-length rVP35 can be stably conjugated to paramagnetic beads and that these conjugates specifically retain a radiolabeled dsRNA probe in a linear dose-dependent manner. Secondly, we determined the optimal conditions for the in vitro reaction at 37 °C, describing the VP35–dsRNA binding dependency by pH, ionic strength, Mg2+ concentration, temperature, time of incubation and amount of beaded-protein. Interestingly, such conditions were comparable to those observed in a different biochemical assay developed to target the dsRNA binding function of the influenza virus NS1 protein [Maroto et al., 2008], a known viral IFN-antagonist that shares several functions with EBOVs VP35. Thirdly, we showed that the full-length rVP35 binds to IVT dsRNA of 50–500 bp in length with high affinity, since the Kd values ranged from 1 to 3 nM according to the temperature used. These Kd values were comparable, although determined with different techniques, with those calculated for other RNA binding proteins, such as EBOV VP30 (Kd = 61 nM) [John et al., 2007], HIV-1 Tat (Kd = 1 nM) [Slice et al., 1992], Hantaan virus N (Kd = 14 nM) [Severson et al., 2005], influenza virus NS1 (Kd = 1 µM) [Chien et al., 2004; Maroto et al., 2008] and the cellular antiviral protein PKR (Kd = 15 nM at one binding site) [Lemaire et al., 2008]. Noteworthy, the EBOV rVP35 Kd value for dsRNA was lower than the Kd values of all these proteins. In particular, it is interesting to compare the affinity for dsRNA between VP35 and the cellular protein RIG-I. In fact, inhibition of the IFN-α/β response due to viral antagonism primarily occurs through the inhibition of the RIG-I-dependent signaling, since this cellular helicase is a crucial sensor of infection through the recognition of viral dsRNA. In this regard, RIG-I stimulation and overexpression has been demonstrated to attenuate EBOVs.
replication in cell cultures (Spiropoulou et al., 2009) and it is likely
that an effective suppression of the IFN-α/β response by EBOVs
would require an efficient sequestration of dsRNA by VP35. Deter-
mination by ITC of the RIG-I dissociation constant for a 12 bp
substrate with a 5’-triposophosphate (5’-ppp) terminus showed K_d
values from 30 to 214 nM, while the K_d value was 1 μM for the
same dsRNA with 5’-OH substrate (Wang et al., 2010). Other
studies performed by surface plasmon resonance at 25 °C with a
14 bp dsRNA with a 5’-ppp terminus yielded a K_d value of 0.3 nM,
while the K_d value was 5 nM for the same dsRNA with 5’-OH
ends (Lu et al., 2010). Using an 8 bp dsRNA we determined
that VP35 K_d values was around 70 nM and 1 μM for 5’-p and
5’-OH oligos, respectively. Therefore, our results show that VP35
could effectively compete with RIG-I for dsRNA binding in the
infected cell. Furthermore, the full length rVP35 demonstrated a
comparable affinity for dsRNA with respect to its truncated C-termi-
nal RBD. In fact, when the VP35 RBD has been studied by ITC, it
showed K_d comparable affinity for dsRNA with respect to its truncated C-ter-
minal RBD. In fact, when the VP35 RBD has been studied by ITC, it
showed K_d value for IVT 18 bp
dsRNA of 890 nM (Kimberlin et al., 2010). Both reported
values are in substantial agreement with those obtained in our
magnetic pull down assay using similar short dsRNA substrates. However, it
is worth to note that full-length rVP35 showed a 25-fold increase in
dsRNA binding affinity for slightly longer IVT dsRNA (50 bp in
length). Hence, it is possible to speculate that VP35 domains other
than RDB may efficiently contribute to the establishment of a
stronger or more stable dsRNA–protein complex with longer
dsRNAs, leading to a higher binding affinity. In addition, it is
important to note that, while a bimodal strategy has been recently
proposed for VP35–RBD–dsRNA binding in which two RBD
monomers cooperatively undertake a dimeric assembly to bind dsRNA
(Kimberlin et al., 2010; Leung et al., 2010c), our data analysis used
a single site binding approach because the rVP35 that conjugates to
the magnetic beads, and binds to the IVT dsRNA, is a tri-tetrameric
homo oligomer (Zinzula et al., 2009 and data not shown).
Therefore, at least under our experimental conditions, the oligo-
merization of the full length rVP35 proteins takes place independ-
ently of the dsRNA presence and rVP35 probably binds to
dsRNA as a tri-tetrameric homo oligomer.
Two recently solved crystallographic structures of the VP35
RBD bound to short RNA duplexes describe a binding mode
to dsRNA where a patch of conserved basic residues interacts with
the end-proximal phosphate backbone and a pocket of hydropho-
bic residues end-caps the dsRNA blunt ends (Kimberlin et al.,
2010; Leung et al., 2010c). According to this model, the simulta-
neous recognition of both strands of the RNA double helix leads to
the formation of a VP35–RBD dimer that mimics the strategies
employed by PRRs to detect viral dsRNA (Kimberlin et al., 2010;
Leung et al., 2010c). Our results agree with a model for which
dsRNA binding is not dependent on the dsRNA sequence or length.
In fact, the full-length rVP35 shows no significant differences in
dsRNA binding affinity for heterologous IVT molecules ranging
from 500 to 50 bp. Moreover, they also agree with an end-capping
binding modality. In fact, the full-length rVP35 binding affinity is
affected by the nature of dsRNA ends, since a 14-fold lower K_d
value is observed for 5’-phosphorylated versus 5’-unphosphorylated
8 bp dsRNAs. In addition, the fact that, in the EMSA studies, the
intensity of the low order wt VP35–dsRNA complexes obtained
with the three IVT dsRNAs is comparable, is also in agreement
with this model.
It has been shown that VP35 RBD has a relatively thermal stabil-
ity when tested in ThermoFluor assays, yielding Tm values of 57
and 63 °C for the ZEBOV and REBOV proteins, respectively (Leung
et al., 2010b). As indicated by Van’t Hoff analysis, at increasing
temperatures the rVP35 affinity to dsRNA diminishes linearly with
a relatively shallow slope, suggesting that the binding reduction is
likely due to changes in the rVP35–dsRNA complex stability fol-
lowing the thermodynamics of the interaction rather than in any
loss in rVP35 binding ability. Noteworthy, even though higher
affinities were shown to be 23 and 30 °C, we selected the more bio-
logically relevant 37 °C for all our studies.
The three basic residues R305, K309 and R312 lie in the VP35
central basic patch that was originally proposed to be required for
dsRNA binding on the basis of its sequence homology with the
RBD of the influenza virus NS1 protein (Hartman et al., 2004).
Alanine substitution of these residues was demonstrated to be critical
for dsRNA binding and to impair VP35 suppression of IFN-α/β
production, even though on a differential extent (Cárdenas et al.,
2006; Hartman et al., 2006, 2008a,b). In particular, in cell culture,
R305A and K309A VP35s suppressed IFN-β activation slightly less
than wt VP35 (Leung et al., 2010c), and K309A VP35 failed to bind piC
beads in a pull down assay (Cárdenas et al., 2006). In addition, while in silico modeling predictions have attributed to R305 only limited interactions with dsRNA, in vitro the truncated (comprising only the RBD) R305A VP35 mutant K_d
value measured by ITC was 3- to 5-fold higher than wt VP35 (Leung
et al., 2010c). Furthermore, the R312A VP35 was shown to be
severely impaired in its IFN-inhibition capability in cell culture
(viruses bearing such mutation were attenuated and avirulent)
(Hartman et al., 2008a,b), the structural data on the VP35 RBD
indicated that R312 is critical for VP35–dsRNA binding (Kimberlin
et al., 2010; Leung et al., 2010a,c,d) and the truncated VP35
R312A mutant showed a total loss of dsRNA binding by ITC, using
an 8 bp IVT dsRNA (Leung et al., 2010c). Our data on full
length rVP35 mutants are in substantial agreement with these
observations. However, our data on full length rVP35 mutants
describe a somehow more subtle gradient in 500 bp dsRNA binding
ability. In fact, all mutants retained a major dsRNA binding capacity,
even though their K_d values were significantly higher than the
one observed for wt rVP35 (a 4-fold difference was observed
between wt and R312A–rVP35 binding affinities). Importantly
however, when compared to wt rVP35, the full length rVP35
R312A showed a 25-fold reduction in binding affinity for the 5’-
phosphorylated 8 bp dsRNA. Furthermore, the comparison of the
full length rVP35 R312A K_d values for the 500 bp IVT dsRNA
and for the 5’-phosphorylated 8 bp dsRNA shows a 166-fold
difference, indicating that the length of the dsRNA severely affects the
mutant binding ability.
Taking together these data, it is possible to hypothesize that, on
the one side, when other VP35 domains are present in addition to
the sole RBD, the VP35 interaction with dsRNA takes place in a
more complex, and/or stable, modality than the one described so
far. In this hypothesis, the contribution of the N-terminal domain
may represent a stabilizing factor. On the other side, it is possible
that these single VP35 mutations may affect only one of the exist-
ing functional modes through which VP35 binds dsRNA. Given
that other VP35 residues have been suggested to be critically impli-
cated in dsRNA binding-mediated IFN inhibition (Leung et al.,
2010a,c; Prins et al., 2010), further mutagenesis studies on the
full-length rVP35 might help to elucidate these aspects.
Among the strategies adopted to interfere with EBOVs propaga-
tion in animal models, two successful attempts based on RNA-
interference approach and the use of phosphorodiamidate mor-
pholino antisense oligomers have been recently reported (Enter-
lein et al., 2006; Geisbert et al., 2010; Warfield et al., 2006).
Interestingly, both studies included the silencing of VP35 gene to
knockdown virus replication. According to this vision, the most
suitable EBOVs countermeasure would be to impair viral replica-
tion and to allow the innate and adaptive immune responses to
clear the infection (Feldmann and Geisbert, 2011). This strategy
further validates the choice of targeting the VP35–dsRNA interaction. With the aim of finding a molecule that would inhibit this interaction and would validate the magnetic pull down reliability for compound screening, we assayed ATA, which was originally recognized as a non-specific inhibitor of protein interactions with nucleic acids (Gonzáles et al., 1980), including the HIV-1 integrase–dsDNA interaction (Fabozzi et al., 2011; Cushman and Sherman, 1992; Tramontano et al., 1998). More recently, ATA has been identified to be a potent and selective inhibitor of viral SARS-CoV and HCV RNA polymerases (Chen et al., 2009), and it has been shown to be 10–100 times more potent than IFN-α/β in inhibiting SARS-CoV replication in infected cells (He et al., 2004; Yap et al., 2005). In our in vitro assay, ATA displayed a dose dependent inhibitory effect on the formation of rVP35–dsRNA complexes, with an IC₅₀ value equal to 50 μg/mL, which is in agreement with similar results obtained for other RNA binding proteins. Importantly, the presently described magnetic pull down assay can be easily scaled up to a high-throughput format and, in fact, studies in this directions are currently underway.

5. Conclusions

In summary, in the present study we performed the first characterization of the full length rVP35 wt and mutants dsRNA binding function that brought novel insights into the VP35–dsRNA interaction. Moreover, we validated a novel and straightforward biochemical in vitro assay for the screening of antiviral agents targeted to the interaction between rVP35 and dsRNA. Finally, given that EBOV VP35 is a paradigm protein for potent viral dsRNA binding-dependent IFN-suppression, the described method can be further applied to other viral proteins that are implicated in the IFN-antagonist inhibition of the innate immunity by dsRNA binding.

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