Interaction of Transportin-SR2 with Ras-related Nuclear Protein (Ran) GTPase*

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Background: Transportin-SR2 (TRN-SR2) is a karyopherin implicated in nuclear import of the HIV-1 preintegration complex.

Results: RanGTP can displace HIV-1 integrase and induces large scale structural changes in TRN-SR2.

Conclusion: Structural and functional analysis of TRN-SR2 supports its role in nuclear import.

Significance: Characterization of TRN-SR2 in the nuclear and cytoplasmic states allows further insights into its function during nuclear import.

The human immunodeficiency virus type 1 (HIV-1) and other lentiviruses are capable of infecting non-dividing cells and, therefore, need to be imported into the nucleus before integration into the host cell chromatin. Transportin-SR2 (TRN-SR2, Transportin-3, TNPO3) is a cellular karyopherin implicated in nuclear import of HIV-1. A model in which TRN-SR2 imports the viral preintegration complex into the nucleus is supported by direct interaction between TRN-SR2 and HIV-1 integrase (IN). Residues in the C-terminal domain of HIV-1 IN that mediate binding to TRN-SR2 were recently delineated. As for most nuclear import cargoes, the driving force behind HIV-1 preintegration complex import is likely a gradient of the GDP- and GTP-bound forms of Ran, a small GTPase. In this study we offer biochemical and structural characterization of the interaction between TRN-SR2 and Ran. By size exclusion chromatography we demonstrate stable complex formation of TRN-SR2 and RanGTP in solution. Consistent with the behavior of normal nuclear import cargoes, HIV-1 IN is released from the complex with TRN-SR2 by RanGTP. Although in concentrated solutions TRN-SR2 by itself was predominantly present as a dimer, the TRN-SR2-RanGTP complex was significantly more compact. Further analysis supported a model wherein one monomer of TRN-SR2 is bound to one monomer of RanGTP. Finally, we present a homology model of the TRN-SR2-RanGTP complex that is in excellent agreement with the experimental small angle x-ray scattering data.

The human immunodeficiency virus type 1 (HIV-1) and other lentiviruses have the capacity to infect non-dividing cells such as macrophages through an active nuclear import mechanism (1, 2). Nuclear import is particularly important in the pathogenesis of HIV-1 because non-dividing cells are a key reservoir of virus in infected individuals. After viral entry and partial uncoating the reverse transcriptase produces a double-stranded DNA copy of the viral RNA genome. During its migration to the nucleus, the reverse transcription complex is gradually transformed into the preintegration complex (PIC).7 Capsid (CA) proteins remain at least partially associated with the PIC during its journey to the nuleopore (3). Upon arrival at the nuclear membrane, the PIC has to overcome the formidable challenge of crossing the nuclear membrane (1, 2). The nuclear membrane is composed of a double lipid bilayer. The outer and the inner nuclear membrane are joined at nuclear pore complexes (NPCs) that serve as entry gates. It is believed that γ-retroviruses wait for the breakdown of the nuclear membrane to access the chromatin, whereas lentiviruses use cellular import pathways for the transport of the PIC through the nuleopore. Different viral signals have been implicated in HIV-1 nuclear import (DNA-flap, IN, Vpr, Matrix), but no single one is accepted as the dominant nuclear import factor (1, 2). The ability of HIV-1 PICs to cross an intact nuclear envelope during interphase implicates the involvement of active cellular transport machineries. Involvement of the classic importin α/β pathway and importin 7 has been proposed by several authors (4–9) and a role for nuclear pore proteins has been described as

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7 The abbreviations used are: PIC, preintegration complex; CA, capsid; NPC, nuclear pore complex; TRN-SR2, Transportin-SR2 (Transportin-3, TNPO3); IN, integrase; SR proteins, serine/arginine-rich proteins; CPSF, cleavage and polyadenylation specificity factor subunit 6; SEC, size-exclusion chromatography; SAXS, small angle x-ray scattering; DSF, differential scanning fluorometry; Imp13, Importin-13.
well (10–12). Also tRNA was proposed as potential nuclear import factor (13).

Using yeast-two-hybrid, we identified Transportin-SR2 (TRN-SR2, Transportin-3), encoded by the TNPO3 gene, as a cofactor of HIV-1 IN (14). TRN-SR2 was independently discovered as a host factor of HIV replication in two large scale siRNA screens (15, 16). The direct interaction between HIV IN and TRN-SR2 has been confirmed independently (17, 18). The karyopherin TRN-SR2 is known to shuttle essential splicing factors, serine/arginine-rich proteins (SR-proteins), between the nucleus and the cytoplasm and is involved in the regulation of mRNA splicing (19). Recognition of SR-proteins by TRN-SR2 mainly relies on the conserved RS-domain and requires phosphorylation. Alternative cargoes lacking an RS domain have been identified as well, indicating that other interactions are possible (20, 21). Transient (siRNA) depletion of TRN-SR2 strongly hampers HIV-1 but not MLV infection. Knockdown of TRN-SR2 in primary macrophages likewise interferes with HIV-1 replication, demonstrating its requirement for productive infection of non-dividing cells. Using quantitative PCR we could pinpoint the block in replication to an event after reverse transcription but before integration and could exclude that TRN-SR2 knockdown affects later steps in the replication cycle. The reduction in the number of two long-terminal repeat circles (14) was confirmed by some (22–24) but not all groups (25–27). In any case the consistent reduction in integration is never accompanied by an increase in two long-terminal repeat circles, suggesting a defect in nuclear import. Interestingly, TRN-SR2 depletion appears to affect integration site selection (28). Using enhanced GFP-labeled IN, a defect in HIV nuclear import upon TRN-SR2 depletion was shown (14).

Some HIV capsid mutations (e.g. N74D CA) reduce the dependence of HIV replication on TRN-SR2 (18, 29). Whether this implies a direct and specific interaction between TRN-SR2 and capsid or capsid core particles remains controversial (29–34). Because many capsid mutations are known to affect uncoating, a plausible explanation for the observed phenotype is the requirement for capsid uncoating before direct interaction between TRN-SR2 and IN can take place (33). The N74D CA mutant was originally selected to overcome restriction by the C-terminally truncated fragment of cleavage and polyadenylation specificity factor subunit 6 ( CPSF6) (29). Full-length CPSF6, a cellular protein involved in splicing, contains an RS domain at its C terminus. Because CPSF6 binds TRN-SR2 and interacts with CA through its N-terminal domain, TRN-SR2 depletion may lead to cytoplasmic accumulation of CPSF6 that in turn may restrict HIV replication at the uncoating step (24, 35). In contrast, spreading replication of N74D CA HIV (as compared with single round transduction) remained highly sensitive to TRN-SR2 depletion (33), suggesting that CPSF6 accumulation does not explain the full phenotype of TRN-SR2 depletion.

Both a peptide-based approach and mass spectrometry-based protein footprinting revealed hot spots for the interaction with TRN-SR2 in the C terminus of HIV-1 IN (30, 36). The cargo domain of TRN-SR2 is required for nuclear import of HIV (22).

TRN-SR2 is a 923-amino acid protein consisting entirely of stacked HEAT repeats (two antiparallel α-helices connected by a small turn linker) (37). These create a curving structure with a high degree of flexibility, allowing binding to different types of cargo and regulatory proteins (38). Importins bind their cargo in the cytoplasm either directly or through the adaptor importinα (39). After docking at the NPC on the cytoplasmic side, the importin-cargo complex moves through the nucleopore channel via its interactions with nucleoporins (Nups) (40). RanGTP binding induces conformational changes in importins leading to the release of the cargo in the nucleus (41). The complex of the importin and RanGTP moves back through the NPC to the cytoplasm where GTP is hydrolyzed, and the import factor is available for a new round of nuclear transport (39). The 25-kDa Ran (Ras-related nuclear protein), a small GTPase, is a key modulator of protein interactions of importins and the motor behind nuclear transport (42). The direction of nuclear transport (import/export) is controlled by its gradient (GTP/GDP-bound forms) across the NPC. RanGTP is enriched in the nucleoplasm, and RanGDP is enriched in the cytoplasm. The nature of the bound nucleotide (GTP or GDP) modulates the interaction between Ran and importins (38, 39).

Because nuclear import is generally believed to be a bottleneck during HIV infection and precedes the integration of the proviral DNA into the host genome, the interaction of IN and TRN-SR2 holds promise as a potential target for anti-HIV therapy. Clear understanding and structural analysis of the interaction of TRN-SR2 with various cargoes is essential before efficient drug development. Here we present biochemical and structural biology studies on the interactions of TRN-SR2 with RanGTP.

**EXPERIMENTAL PROCEDURES**

**Purification of His6–TRN-SR2—Cultures of Escherichia coli**

strain Rosetta (DE3) transformed with pET19b–TRN-SR2 in LB medium were induced with 0.5 mM isopropyl β-D-thiogalactoside at A600nm ~ 0.6 and incubated 6 h at 30 °C. The cultures were harvested by centrifugation for 15 min at 4000 rpm and 4 °C, and pellets were washed with STE buffer (100 mM NaCl, 10 mM Tris-HCl, pH 7.3, 0.1 mM EDTA), centrifuged again, and then stored at −20 °C until purification. The frozen cultures were thawed and resuspended in lysis buffer (50 mM Tris-HCl, pH 7.3, 0.5 mM NaCl, protease inhibitors (Complete, EDTA-free, Roche Applied Science), 2 units DNase/10 ml, 5 mM dithiothreitol (DTT)). Cells were lysed by a French press and sonication on ice, and the lysate was centrifuged at 15,000 rpm at 4 °C for 15 min. The soluble lysate containing His6-fused protein was loaded onto a Ni2+–affinity column equilibrated with binding buffer (20 mM Tris-HCl, pH 7.5, 250 mM NaCl, 5 mM DTT) using an AKTA purifier system. The protein was eluted with a linear gradient of imidazole (0–1 M), and 1-ml fractions were collected. Fractions with a peak absorption at 280 nm were pooled, concentrated by centrifugal concentrators (Vivaspin 6 50,000 MWCO PES, Sartorius Stedim Biotech), and dialyzed overnight against buffer A (20 mM Tris-HCl, pH 7.5, 5 mM DTT) before ion exchange chromatography. A gradient from 50 mM to 1 M NaCl in buffer A was run using a HiTrap Q HP 5-ml column. The 1-ml fractions containing His6–TRN-SR2 pro-
tein were pooled and concentrated with Vivaspin. The last step in the purification procedure was size-exclusion chromatography (SEC) using a HiLoad 16/60 Superdex 200 prep grade column. SEC was performed in 10 mM Tris-HCl, pH 7.5, 150 mM NaCl, 5 mM DTT. The pooled fractions with His9-TRN-SR2 were stored at 4 °C.

**Purification of GST-TRN-SR2**—Cultures of E. coli strain BL21 transformed with pGEX-6P2-TRN-SR2 in LB medium substituted with 1 M d-sorbitol and 2.5 mM trimethylglycine were induced with 0.5 mM isopropyl β-D-thiogalactoside at 30 °C and incubated overnight at 28 °C. The cultures were harvested by centrifugation for 10 min at 4000 rpm at 4 °C, and pellets were washed with STE buffer, centrifuged again, and stored at −20 °C until purification. The frozen cultures were thawed, resuspended in lysis buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, protease inhibitors, 2 units DNase/10 ml, 5 mM DTT), and lysed by a French press, and the lysate was centrifuged at 15,000 rpm at 4 °C for 30 min. The soluble lysate containing GST-fused protein was loaded onto a glutathione-Sepharose column equilibrated with binding buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 5 mM DTT). The protein was eluted with the elution buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 5 mM DTT, 20 mM reduced glutathione), and 1-ml fractions were collected. Fractions exhibiting a peak absorption at 280 nm were pooled and dialyzed overnight against 100 mM Tris-HCl, pH 7.5, 150 mM NaCl, protease inhibitors, 5 mM DTT, and 10% glycerol. The pooled fractions with His9-TRN-SR2 were stored at −80 °C.

**Purification of GST-ASF/SF2**—Cultures of E. coli strain BL21 transformed with pGEX-2TK-ASF/SF2 in LB medium were induced with 0.5 mM isopropyl β-D-thiogalactoside at 30 °C and incubated for 6 h at 25 °C. The cultures were harvested by centrifugation for 10 min at 4000 rpm at 4 °C, and pellets were washed with STE buffer, centrifuged again, and stored at −20 °C until purification. The frozen cultures were thawed, resuspended in lysis buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1 mM PMSF, 2 units DNase/10 ml, 10 μM/ml RNase A, 5 mM DTT), and lysed by sonication, and the lysate was centrifuged at 15,000 rpm at 4 °C for 30 min. The soluble lysate containing GST-fused protein was loaded onto a glutathione-Sepharose column equilibrated with binding buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 5 mM DTT). The protein was eluted with the elution buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 5 mM DTT, 20 mM reduced glutathione), and 1-ml fractions were collected. Fractions exhibiting a peak absorption at 280 nm were pooled and dialyzed overnight against 100 mM Tris-HCl, pH 7.5, 150 mM NaCl, 5 mM DTT, 10% glycerol) and stored at −80 °C.

**Purification of His6-HIV-1 IN**—N-terminally His6-tagged HIV-1 IN was purified as described previously (43).

**Purification of His9-Ran and Ran**—Cultures of E. coli strain BL21 (DE3) transformed with pET-3d-His9-HRV3C-RanQ69L in LB medium were induced with 0.5 mM isopropyl β-D-thiogalactoside at 30 °C and incubated overnight at 28 °C. The cultures were harvested by centrifugation for 10 min at 6000 rpm at 4 °C, and pellets were washed with STE buffer, centrifuged again, and then stored at −20 °C until purification. The frozen cultures were thawed, resuspended in lysis buffer (20 mM Tris-HCl, pH 7.5, 0.25 M NaCl, 1 mM PMSF, 20 mM imidazole, 4 mM MgCl2, 2 units DNase/10 ml, 5 mM DTT), and lysed by sonication on ice, and the lysate was centrifuged at 15,000 rpm and 4 °C for 30 min. The soluble lysate containing His9-fused protein was loaded onto nickel column equilibrated with binding buffer (20 mM Tris-HCl, pH 7.5, 250 mM NaCl, 50 mM imidazole, 4 mM MgCl2, 5 mM DTT). The protein was eluted with elution buffer (20 mM Tris-HCl, pH 7.5, 250 mM NaCl, 250 mM imidazole, 4 mM MgCl2, 5 mM DTT), and 1 ml fractions were collected. Fractions exhibiting peak absorption at 280 nm were pooled and dialyzed overnight against 100 mM volume of dialysis buffer (20 mM Tris-HCl, pH 7.5, 250 mM NaCl, 4 mM MgCl2, 5 mM DTT, 10% glycerol) or used to cleave off the His tag. The dialyzed protein was stored at −80 °C.

**Nucleotide Loading of Ran**—To load recombinant Ran with nucleotides (GDP or GTP), maximally 100 μM Ran was incubated for 30 min at 30 °C with 1 mM GDP or GTP and 20 mM EDTA. After 30 min, the reaction was stopped by adding 50 mM MgCl2. The buffer was exchanged on a PD10 desalting column to the Ran dialysis buffer (20 mM Tris-HCl, pH 7.5, 250 mM NaCl, 4 mM MgCl2, 5 mM DTT, 10% glycerol). The peak fractions were pooled and used immediately or stored for later use at −80 °C.

**Complex Formation and Analytical Size Exclusion Chromatography**—For complex formation, His9-Ran2 and RanGTP/GDP were mixed in a 1:3 molar ratio and incubated on ice for at least 1 h. SEC runs were performed on a Superdex 200 10/300 GL column attached to an AKTA purifier system (GE Healthcare) at a flow rate of 0.5 ml/min at 4 °C in SEC buffer containing 10 mM Tris-HCl, pH 7.3, 150 mM NaCl, 5 mM MgCl2, and 5 mM DTT. Proteins were detected by absorbance at 280 nm. The column was calibrated with the following proteins: ferritin (440 kDa), catalase (232 kDa), aldolase (158 kDa), conalbumin (75 kDa), ovalbumin (43 kDa), carbonic anhydrase (29 kDa), ribonuclease A (13.7 kDa), and aprotinin (6.5 kDa) (Gel Filtration Calibration kits HMW & LMW; GE Healthcare). The fractions were analyzed by SDS-PAGE and silver-stained following the manufacturer’s instructions.

**AlphaScreen Interaction Assays**—The AlphaScreen assays (Amplified Luminescent Proximity Homogeneous Assay, ALPHA; PerkinElmer Life Sciences) were optimized for use in 384-well OptiPlate microplates (PerkinElmer Life Sciences) with a final volume of 25 μl. Recombinant proteins (GST- or His9-tagged TRN-SR2, GST-ASF/SF2, His9-tagged or untagged Ran charged with GDP or GTP and His9-IN) were diluted to 5× working solutions in assay buffer (25 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1 mM DTT, 1 mM MgCl2, 0.1% (v/v) Tween 20, 0.1% (w/v) bovine serum albumin). First, 5 μl of buffer or diluted RanGTP was pipetted into the wells followed by 5 μl of each of the interacting protein dilutions. The plate was sealed and left to incubate for 1 h at 4 °C, allowing an equilibrium to be established. Next, 10 μl of a mix of Ni2+ chelate acceptor and glutathione donor AlphaScreen beads (PerkinElmer Life Sciences) was added, bringing the total volume to 25 μl and establishing
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final concentrations of 10 μg/ml for each of the beads. The plate was then placed at room temperature and incubated for one more hour before being read in an EnVision Multilabel Reader (PerkinElmer Life Sciences). Background AlphaScreen counts were subtracted, and data were analyzed in Prism 5.0 (GraphPad).

The cargo dissociation experiments were performed with final concentrations of 10 nm GST-TRN-SR2 or His6-Tr-mTRN-SR2 and 40 nm His6-LN or 40 nm GST-ASF/SF2, respectively. All titrations (RanGDP/GDP) were performed against 10 nm GST-TRN-SR2.

Small Angle X-ray Scattering (SAXS) Measurements—SAXS data were collected using synchrotron radiation at the European Molecular Biology Laboratory X33 beamline of the DORIS III storage ring (DESY, Hamburg, Germany). SAXS curves were measured over the range of momentum transfer 0.006 < q = 4π sin(θ)/λ < 0.63 Å−1, where θ is the scattering angle, and λ = 1.5 Å is the x-ray wavelength. Protein samples were in 20 mM Tris-HCl buffer, pH 7.3, 150 mM NaCl, 5 mM DTT, and 5 mM MgCl2. The ATSAS program package (44, 45) was used for data processing. Guinier plots were used to evaluate the zero angle scattering (I0) and the radius of gyration (Rg). The particle distance distributions were evaluated from SAXS data for q < 0.3 Å−1 using SASSMoW (47).

The fits between the experimental scattering curves and the theoretical scattering from atomic models were calculated using CRYSOIL (48).

Dynamic Light Scattering—Measurements were made in small droplets with the SpectroSize 300 instrument at the scattering angle of 150° and at 20 °C. As with the SAXS measurements, protein samples were in 20 mM Tris–HCl buffer, pH 7.3, 150 mM NaCl, 5 mM DTT, and 5 mM MgCl2.

Different Scanning Fluorimetry—His6-Tr-mTRN-SR2 WT or the E145Q, V149A, E152Q, E153Q mutant at 1 μM final concentration was mixed with 1× SYPRO Red dye (Invitrogen), and the corresponding dilution of Ran was loaded with either GDP or GTP (3-fold dilution series from 3 to 0.33 μM). Mixtures were left for 15 min at room temperature before 25 μl was transferred to two wells of a 96-well plate (Bio-Rad). The plate was sealed with optical flat 8-cap strips (Bio-Rad), and differential scanning fluorimetry (DSF) melting curves were obtained on a Bio-Rad iCycler equipped with an iQ5 real-time PCR detection system. The raw fluorescence data were analyzed with Excel (Microsoft), whereas Prism 5.0 (GraphPad) was used to fit the transitions with a Boltzmann sigmoidal equation and extract melting temperatures.

Homology Modeling—The TRN-SR2 amino acid sequence was retrieved from the UniProt Knowledge base (identifier: Q9Y5L0–2) and used as a query on the HHpred server (49). The Research Collaboratory for Structural Bioinformatics Protein Data Bank (RCSB PDB) was hence searched with the profile Hidden Markov Model-Hidden Markov Model (HMM-HMM) comparison through HHblits (8 iterations maximum, secondary structure scoring on), and the resulting query-template alignments were realigned with the Maximum ACCuray alignment (MAC) algorithm. The best templates were selected, and a multiple alignment was generated. These alignments pointed to Importin-13 (Imp13) as the highest quality template for homology modeling of TRN-SR2 (E values of 0 for different HMMs). The relatively low sequence homology (23% identical, similarity around 38%), however, necessitates careful validation of the final model quality. The optimal alignment to Imp13, corresponding to a maximal TM-score of 0.9194, was subsequently used for model building in MODELLER 9.8 (50). Three crystal structures of Imp13 are available in the PDB, 2XWU, 2X19, and 2X1G, which represent two complexes of human Imp13 with UBC9 or RanGTP and one of Drosophila Imp13 with Mago and Y14, respectively. For the RanGTP-bound model of TRN-SR2, structure 2X19 was used as a template, and the RanGTP structure (Saccharomyces cerevisiae) from 2X19 was set as an environment for induced fit after it was modified to its human counterpart.

RESULTS

Functionality of the TRN-SR2-RanGTP Interaction—We first studied the direct interaction between TRN-SR2 and Ran loaded with either GDP or GTP (RanGDP or RanGTP, respectively). Ran is a crucial regulator of nuclear transport, and the regulation is executed through conformational changes in the Ran switch I and switch II loops depending on whether guanosine di- or triphosphate is bound (51, 52), altering the affinity for different binding partners. To ensure that GTP is not hydrolyzed to GDP during our experiments, a GTPase-deficient Ran mutant, RanQ69L (henceforth simply referred as Ran), was used that is characterized by a severely reduced hydrolysis of GTP to GDP (53). As expected, RanGDP and RanGTP bound TRN-SR2 to a different extent, as evidenced by AlphaScreen (Fig. 1). We determined an apparent Kd = 4.7 ± 1.2 nM for the TRN-SR2-RanGTP interaction, whereas no binding to RanGDP could be detected under tested conditions.

Using DSF, we provided further support for this specific interaction and assessed the thermostability of the established complex. The addition of RanGDP to TRN-SR2 did not significantly affect the observed melting temperature of TRN-SR2 (supplemental Fig. S1A). In contrast, the addition of increasing amounts of RanGTP markedly increased the melting temperature. In the presence of a 3× molar excess of RanGTP, the melting temperature of TRN-SR2 was 56.6 °C compared with
45.7 °C for TRN-SR2 alone (supplemental Fig. S1B). These data clearly point to the formation of a stable TRN-SR2-RanGTP complex in solution.

Structural analysis of proteins in vitro gains in relevance if the purified recombinant proteins are able to perform their physiological reactions. After evaluating the interaction with RanGTP, we determined whether recombinant TRN-SR2 binds its natural cellular cargo and, more importantly, whether cargo can be released from the complex upon the addition of RanGTP. 20 nM His9-TRN-SR2 (henceforth referred to as TRN-SR2) was incubated with 80 nM GST-tagged ASF/SF2 (SRSF1), one of the known cellular cargoes of TRN-SR2 (19) or 80 nM His6-HIV-1 IN and increasing amounts of RanGTP. When the TRN-SR2-ASF/SF2 interaction was probed with AlphaScreen, a clear RanGTP-dependent dissociation could be observed, demonstrating the functionality of the recombinant protein (Fig. 2A). Similarly, with HIV-1 IN, the viral cargo was efficiently released from TRN-SR2 by increasing the concentrations of RanGTP (Fig. 2B). Indirectly this finding indicates that the TRN-SR2 basic functionality is likely independent of eukaryotic post-translational modifications.

**TRN-SR2 and Its Complex with Ran Characterized by DLS and SAXS**—We next used DLS to obtain further size characteristics of the proteins. We measured samples of TRN-SR2 alone and the TRN-SR2-RanGTP complex at varying total protein concentrations (2–16 mg/ml). All samples were shown to be monodisperse. DLS data allowed determination of radii of hydration ($R_h$) for the TRN-SR2-RanGTP complex and TRN-SR2 alone, respectively (Table 1 and supplemental Fig. S2).

Subsequently, we performed SAXS to characterize the solution structure of TRN-SR2 and its complex with RanGTP. The measurements were done on samples with varying total protein concentrations (2–16 mg/ml) (Fig. 5A). The SAXS data permitted the calculation of the radius of gyration ($R_g$), the maximum dimension ($D_{max}$) (Table 2), and pair-distance distributions $P(r)$ for each sample (Fig. 5B). All studied solutions of both
TRN-SR2 alone and the complex yielded scattering with linearity in the Guinier region, indicating no considerable aggregation. All samples were monodisperse, with little dependence of the normalized SAXS curve on protein concentration (Table 2). For free TRN-SR2 at 8 mg/ml, the measured $R_g$ was 4.5 nm, and the maximum particle dimension $D_{\text{max}}$ was 15.5 nm, which is close to the values reported earlier for TRN-SR2 dimers (30). Moreover, the apparent molecular mass estimated from the SAXS data at this concentration is 202 kDa, which is a close match to the theoretical mass of the dimer (214 kDa). Interestingly, comparison of the intraparticle distance distributions for TRN-SR2 alone and for the TRN-SR2-RanGTP complex (Fig. 5B) clearly suggests a smaller particle in the latter case. For the complex at 8.1 mg/ml, the $R_g$ was 3.6 nm, and $D_{\text{max}}$ was 10.7 nm (Table 2). The apparent molecular mass for the complex was also clearly smaller (131 kDa) than for TRN-SR2 alone. The fact that the TRN-SR2-RanGTP complex in solution is smaller than TRN-SR2 alone is also evident from comparing the lowest q parts of the scaled scattering curves (Fig. 5A), which clearly

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**TABLE 1**

Solution species parameters derived from dynamic light scattering

| Sample            | $R_h$ (nm) |
|-------------------|------------|
| TRN-SR2           | 4.78 ± 0.10 |
| TRN-SR2-RanGTP    | 3.92 ± 0.17 |

**TABLE 2**

Solution species parameters derived from small-angle x-ray scattering

| Sample            | Concentration | $I_0$ (a.u.) | $R_g$ (nm) | $D_{\text{max}}$ (nm) | $M_r$ (kDa) |
|-------------------|---------------|--------------|------------|------------------------|-------------|
| TRN-SR2           | 1.8           | 118.4        | 4.37       | 15.3                   | 181.0       |
|                   | 4             | 115.2        | 4.37       | 15.3                   | 200.5       |
|                   | 8             | 119.4        | 4.51       | 15.5                   | 202.3       |
|                   | 16.3          | 133.0        | 4.86       | 17.2                   | 219.8       |
| TRN-SR2-RanGTP    | 2.1           | 81.2         | 3.60       | 11.3                   | 144.9       |
|                   | 4             | 85.7         | 3.66       | 11.2                   | 140.8       |
|                   | 8.1           | 80.7         | 3.61       | 10.7                   | 130.5       |
|                   | 15.8          | 88.1         | 3.65       | 10.3                   | 134.3       |
point to smaller extrapolated zero angle scattering ($I_0$) for the complex. Importantly, the measured mass of 131 kDa is very close to the expected mass of a complex containing one TRN-SR2 molecule and one RanGTP molecule (a total of 132 kDa). The values for $R_g$ correspond well to the hydrodynamic radii ($R_g$) (Table 1) as obtained in the DLS experiments, considering the most ordered first solvation shell to be 0.3 nm thick (54).

**Homology Model of the TRN-SR2-RanGTP Complex**—At the moment there is no high resolution structure reported for the TRN-SR2 protein described. However, the structure of its closest parologue, Imp13, has been solved. In 2010 Bono et al. (55) reported the structures of the complexes of human Imp13 with the S. cerevisiae Ran homologue, GTP binding nuclear protein GSP1/CNR1, and of the Drosophila melanogaster Imp13 (Cadmus) with Mago and Y14 homologues (Protein mago nashi and RNA-binding protein 8A, respectively, in fruit fly). Later the structure of the complex of human Imp13 with sumo-conjugating enzyme Ubc9 was published (56). Since the original duplication event, TRN-SR2 and Imp13 have diverged significantly, resulting in a present day low sequence identity. Nonetheless, we could produce alignments of human TRN-SR2 to both human and Drosophila Imp13 using HHpred, the most optimal of which is shown in supplemental Fig. S3 (22% sequence identity, probabilities = 100% and E values = 0) (49). This alignment was used to guide modeling of the human TRN-SR2-RanGTP complex (Fig. 6A).

Importantly, we have calculated a theoretical SAXS curve from the obtained homology model of the TRN-SR2-RanGTP complex and compared it to the experimental data. A good match was obtained (goodness of fit $\chi = 1.846$, Fig. 6B) which confirms that in solution TRN-SR2 and RanGTP indeed form a 1:1 complex. For further validation, we have introduced E145Q,V149A,E152Q,E153Q mutations into TRN-SR2 (referred to as TRN-SR2EVEE), which the modeling predicts to be at the interaction interface with RanGTP (supplemental Fig. S4A). Indeed, although DSF showed that the stability of free TRN-SR2 was unaffected by the substitutions ($T_m = 45.7 \pm 0.5 ^\circ C$ and $47.1 \pm 0.9 ^\circ C$ for TRN-SR2WT and TRN-SR2EVEE, respectively), the temperature shift upon the addition of 3 $\mu$m RanGTP was significantly smaller for TRN-SR2EVEE compared with TRN-SR2WT ($\Delta T = 5.8 \pm 0.9 ^\circ C$ versus $11.8 \pm 0.3 ^\circ C$, respectively, supplemental Fig. S4B). The smaller shift indicates that a complex can still be formed between RanGTP and TRN-SR2EVEE, likely due to the multiple interfaces between RanGTP and TRN-SR2 (see below), but that the affinity is significantly reduced. Together, these results support the accuracy of the proposed model despite the relatively low sequence identity of the template used (below 30%, the so-called “twilight zone”).

**Detailed Interactions between TRN-SR2 and RanGTP in the Complex**—TRN-SR2 is an all-helical protein (Fig. 6A and supplemental Fig. S3), as can be expected for a member of the karyopherin-β family. In the case of TRN-SR2 these helices are arranged into 19 typical HEAT repeats with A (inside) and B (outside) helices and one C-terminal capping 3-helix HEAT repeat (A, B, and C helices). On a larger scale, the HEAT repeats stack on top of one another into a flexible toroid shape. When binding to RanGTP, the TRN-SR2 toroid wraps around the smaller Ran protein, most likely compressing its toroid shape and bringing the N and C terminus closer together (Fig. 6A). Despite contacts with the TRN-SR2 middle and C-terminal parts (see below), RanGTP mainly occupies the inside of the N-terminal half of the toroid. This N-terminal RanGTP binding is a general observation for members of the karyopherin family, which is reflected in a higher degree of sequence conservation in this region (supplemental Fig. S3).

The small GTPase Ran has been extensively studied and characterized (57–59). Upon exchanging GDP for GTP or vice versa, two stretches of this protein undergo a conformational change, adequately called the switch I and II regions (amino acids Thr-32–Val-45 and Thr-66–Tyr-80 respectively) (60).
These switches are responsible for specific recognition of Ran in its GTP-bound state by importins and hence for import cargo displacement in the nucleus (59, 61). In our model, like with Imp13 and exportin Crm1, switch I establishes interactions with HEAT repeats 16 to 19 (Fig. 7, right zoom). Notably, two lysines on RanGTP (Lys-37 and Lys-38) are predicted to engage in electrostatic interactions with residues Asn-744, His-745, and Asp-747 from HEAT repeat 17 and Asp-787 and His-788 from repeat 18. As evident from the alignment (supplemental Fig. S3), these interactions are largely conserved between TRN-SR2 and Imp13 (TRN-SR2 Asp-787 can potentially assume the role of Imp13 Glu-830 here). The Ran switch II region also shows electrostatic complementarity (Fig. 7, bottom zoom), as a salt bridge is formed between Lys-68 on the TRN-SR2 HEAT 2 B-helix and Asp-77 on Ran. Meanwhile, the hydrophobic Leu-74 is buried in a superficial groove lined by conserved Leu-18, Tyr-19, Leu-34, Gln-38, and Phe-62 side chains and located between HEAT 1 and 2 B-helices (Fig. 7, bottom, supplemental Fig. S3). The area next to switch II on RanGTP is also in close contact and further contributes to electrostatic interactions with the TRN-SR2 N terminus; two Ran arginine residues (Arg-106, Arg-110) balance the charge of two conserved glutamic acids (Glu-152, Glu-153) located on the TRN-SR2 HEAT 4 B-helix (Fig. 7, left zoom, supplemental Fig. S3). A final interaction site was formed in the middle of TRN-SR2 involving an acidic stretch encompassing Glu-391, Glu-392, and Asp-394 on HEAT repeat 9 and its basic counterpart on Ran involving residues Lys-159, Arg-166, and Lys-167 (Fig. 7 left, supplemental Fig. S3). This central interaction is found in other karyopherins as well, albeit with small differences concerning the exact position of the acidic stretch (56). Overall, mainly electrostatic interactions seem to steer complex formation between TRN-SR2 and RanGTP.

**DISCUSSION**

Although TRN-SR2 is generally accepted to act as a cofactor for HIV replication, its exact role remains unclear. According to one of the (non-exclusive) hypotheses, HIV nuclear import is mediated by the direct interaction between TRN-SR2 and HIV integrase present in the PIC (14). All karyopherin-mediated nuclear import is directed by a gradient of RanGTP/GDP (41). As a step in the elucidation of the nuclear import mechanism of
HIV, here we studied the biochemical and structural aspects of the interaction between TRN-SR2 and RanGTP.

First we demonstrated in AlphaScreen that Ran binds TRN-SR2 in its GTP-loaded form only (Fig. 1). This result was confirmed with DSF, which showed increased thermostability upon binding to RanGTP, also pointing to stable complex formation (supplemental Fig. S1). RanGTP is further able to displace recombinant HIV integrase from TRN-SR2, much alike its well known cellular cargo ASF-SF2 (Fig. 2). This result is consistent with the model in which nuclear import is mediated by the direct interaction between TRN-SR2 and HIV IN. According to this model RanGTP will dissociate the complex after its arrival in the nucleus.

Next, we performed SEC (Figs. 3 and 4), DLS (supplemental Fig. S1, Table 1), SAXS (Fig. 5, Table 2), and mutagenesis and homology modeling (Fig. 6) to study the structure of TRN-SR2 and its complex with RanGTP in solution (Fig. 7). Importins in general show high conformational flexibility and variability (62, 63) allowing them to adapt their conformations in response to their cargo and the highly dynamic environment of the NPC. The original view of the accommodation of different binding partners via induced-fit mechanisms was replaced by a concept of population shift between preexisting alternative conformations (64). This makes studies of conformational states of these flexible helicoids very demanding, especially at the level of interpretation of the obtained data. The elastic behavior of solenoid proteins like TRN-SR2 is dominated by non-polar interactions between HEAT repeats (65). In particular, the hypothesis of a soft nanospring has arisen with a molten globule-like hydrophobic core (65), where small changes between HEAT actions between HEAT repeats (65). In particular, the hypothesis of a soft nanospring has arisen with a molten globule-like hydrophobic core (65), where small changes between HEAT repeats distort the karyopherin pathway in human immunodeficiency virus type 1 nuclear import. J. Virol. 70, 1027–1032

In line with a recent report, our SAXS data for TRN-SR2 alone indicate that the protein predominantly exists as a dimer in solution (30). For TRN-SR2-RanGTP, SEC, SAXS, and homology modeling data strongly support a 1:1 stoichiometry of the complex. The remarkable increase in the SEC elution volume ($V_e$) as well as the decreased values of $R_g$ (SAXS) and $R_h$ (DLS) may hence result from dissociation of TRN-SR2 dimers as well as the conformational changes in monomeric TRN-SR2 while wrapping around RanGTP. As we are likely observing two processes at the same time, dissociation of TRN-SR2 dimers and formation of the 1:1 TRN-SR2-RanGTP complex, it is difficult to make conclusions on conformational changes within monomeric TRN-SR2 itself upon RanGTP binding. For importin $\beta$, which stayed monomeric in solution, both $R_g$ and $D_{max}$ decreased upon binding of RanGTP. In contrast, for TNPO1 the $R_g$ value increased and $D_{max}$ remained the same upon binding of Ran (62). The fact that importin $\beta$ and TNPO1 are more related in the sequence-based phylogenetic tree and distant from TRN-SR2 (66) suggests that the structural changes may be dictated rather by the higher order structural elements (HEAT repeats) than by the primary sequence. Potentially extensive conformational changes in TRN-SR2 versus TRN-SR2-RanGTP are in accordance with SAXS measurements available for importin $\beta$ and TNPO1 (62). A more compact and rigid conformation of the complex with RanGTP in comparison with the free karyopherin was also demonstrated by (non)equilibrium molecular dynamics simulations for importin $\beta$ (66). However, to fully understand the role of preexisting alternative conformations (such as TRN-SR2 dimers), additional studies have to be conducted. Due to the intrinsic flexibility of TRN-SR2, it will also be of interest to perform molecular dynamics simulation of the contacts involved in the interaction of the proteins.

Last, because of the good fit between experimental and model back-calculated SAXS curves for the TRN-SR2-RanGTP complex and the additional support from mutagenesis and DSF, we can consider the molar ratio of 1:1 between both protein partners as corroborated. Furthermore, this validated model provides detailed insights into the complex (Fig. 7). RanGTP establishes elaborate contacts with TRN-SR2; while mainly occupying the N-terminal part of the TRN-SR2 toroid, Ran also contacts several central and C-terminal HEAT repeats. The interface between both proteins is characterized by extensive charge complementarity, notably involving the switch I and II regions of Ran, which determine nucleotide-dependent recognition of protein partners. Similar to other importins, mainly electrostatic interactions seem to steer complex formation between TRN-SR2 and RanGTP. These details on the TRN-SR2-RanGTP interaction and on TRN-SR2 itself may provide important insights into its function as a nuclear import factor for HIV-1. Additionally, these structural insights provide a first stepping stone toward modulation of TRN-SR2 function or inhibition of its interaction with HIV-1 IN for therapeutic purposes.

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