Stability of local secondary structure determines selectivity of viral RNA chaperones

Jack P.K. Bravo¹,², Alexander Borodavka¹,²,³,*, Anders Barth³, Antonio N. Calabrese¹,², Peter Mojzes⁴, Joseph J.B. Cockburn¹,², Don C. Lamb³ and Roman Tuma¹,²,⁵,*

¹Astbury Centre for Structural Molecular Biology, University of Leeds, Leeds, UK, ²School of Molecular and Cellular Biology, University of Leeds, Leeds, UK, ³Department of Chemistry, Center for NanoScience (CeNS), Nanosystems Initiative Munich (NIM) and Centre for Integrated Protein Science Munich (CiPSM), Ludwig Maximilian University of Munich, Munich, Germany, ⁴Institute of Physics, Faculty of Mathematics and Physics, Charles University, Ke Karlovu 5, CZ-12116 Prague 2, Czech Republic and ⁵Faculty of Science, University of South Bohemia, Ceske Budejovice, Czech Republic

Received April 03, 2018; Revised April 24, 2018; Editorial Decision April 25, 2018; Accepted April 30, 2018

ABSTRACT

To maintain genome integrity, segmented double-stranded RNA viruses of the Reoviridae family must accurately select and package a complete set of up to a dozen distinct genomic RNAs. It is thought that the high fidelity segmented genome assembly involves multiple sequence-specific RNA–RNA interactions between single-stranded RNA segment precursors. These are mediated by virus-encoded non-structural proteins with RNA chaperone-like activities, such as rotavirus (RV) NSP2 and avian reovirus αNS. Here, we compared the abilities of NSP2 and αNS to mediate sequence-specific interactions between RV genomic segment precursors. Despite their similar activities, NSP2 successfully promotes inter-segment association, while αNS fails to do so. To understand the mechanisms underlying such selectivity in promoting inter-molecular duplex formation, we compared RNA-binding and helix-unwinding activities of both proteins. We demonstrate that octameric NSP2 binds structured RNAs with high affinity, resulting in efficient intramolecular RNA helix disruption. Hexameric αNS oligomerizes into an octamer that binds two RNAs, yet it exhibits only limited RNA-unwinding activity compared to NSP2. Thus, the formation of inter-segment RNA–RNA interactions is governed by both helix-unwinding capacity of the chaperones and stability of RNA structure. We propose that this protein-mediated RNA selection mechanism may underpin the high fidelity assembly of multi-segmented RNA genomes in Reoviridae.

INTRODUCTION

Members of the Reoviridae family of double-stranded (ds) RNA viruses encompass important human, animal and plant pathogens (1). Reovirus genomes are distributed between 9 and 12 RNA segments, all of which are essential for virus assembly and replication (2). Multiple sequence-specific RNA–RNA interactions between ssRNA segment precursors are believed to underpin the assembly of a complete multi-segmented viral genome (3–7). The underlying molecular mechanisms of high fidelity RNA selection and accurate genome assembly remain poorly understood (1,3). Segment selection, genome replication and virus assembly occur in cytoplasmic membraneless organelles, termed viral factories or viroplasms (8–11). Non-structural proteins rotavirus (RV) NSP2, or mammalian and avian reovirus (ARV) αNS are major components of viroplasms and viral factories, and are essential for genome replication and virus assembly (8,12–16).

Recently, we have demonstrated that NSP2 can promote selective RNA duplex formation between genomic ssRNAs, acting as an RNA chaperone (17). We have shown previously that ARV αNS exhibits similar non-specific ssRNA binding, helix-destabilizing and strand-annealing activities to NSP2, suggesting that both proteins may play similar roles during genome assembly (18). Despite these apparent similarities, it remains unclear whether these proteins employ similar mechanisms of facilitating specific inter-segment interactions.

Here, using a previously established interaction between RV segments S5 and S11 ssRNAs as a model system, we compared the abilities of NSP2 and αNS to promote inter-segment RNA duplex formation. This allowed us to decouple the RNA chaperone activities of both proteins from their multiple, overlapping roles during virus replication, such as viroplasm assembly, kinase activity of NSP2 and...
interactions with other viral and cellular proteins (8,12–16). While NSP2 is efficient at mediating stable S5: S11 RNA–RNA interactions, σNS completely fails to do so. While both proteins simultaneously bind multiple unstructured ssRNAs with similar affinities, NSP2 has a greater propensity for binding and unfolding structured RNA hairpins. RNA stem-loop unwinding experiments monitored by single-pair Förster resonance energy transfer (FRET) indicates that NSP2 disrupts RNA secondary structure more efficiently, whereas σNS binding induces an ensemble of partially unfolded intermediates. Upon binding multiple RNAs, hexameric σNS undergoes further oligomerization, forming stable octameric ribonucleoprotein (RNP) complexes with increased helix destabilizing activity. Such differences in modes of RNA binding and helix-unwinding between these proteins may explain the failure of σNS to promote a stable inter-segment S5 : S11 RNA–RNA interaction. These results suggest that RNA structural stability can modulate viral RNA chaperone activity, thereby restricting non-cognate RNA duplex formation during segment selection and genome assembly.

MATERIALS AND METHODS

Protein expression and purification

NSP2 and σNS were expressed and purified as previously described in (18). Size exclusion chromatography (SEC) analysis was performed using Superdex200 10/300 column (GE healthcare) equilibrated in SEC buffer (50 mM HEPES pH 8.0, 150 mM NaCl) at 4°C.

RNA production

RV segments S5 and S11 were produced and labeled as previously described in Borodavka et al. (19). AlexaFlour488 dye-labeled 20mer and unlabeled 15mer, 20mer and 40mer RNAs (Supplementary Table S1) were purchased from Integrated DNA technologies (IDT). Cy3- and Cy5-labeled 17mer RNAs (Supplementary Table S1) were derived from the RV segment S11 (nucleotides 49–66, complementary to segment S5 nucleotides 307–324) and ordered from IDT. Cy3- and Cy5-labeled dye-labeled 20mer and unlabeled 15mer, 20mer and 40mer RNAs, hexameric σNS and 20mer RNA (Supplementary Table S1) to ensure complete saturation of 20mer RNA, assuming that a single σNS hexamer binds two RNA molecules. In the case of a higher-order σNS–RNP complex (i.e. octamer bound to 2 RNAs), this stoichiometry also ensures protein saturation with RNA. For light scattering analysis, 25 μM σNS and σNS:20mer complex were run at a flow-rate of 0.4 ml min⁻¹ on a TSKgel G6000PWxl SEC column (Tosoh) with an AKTA pure system (GE Healthcare) connected to a DAWN HELEOS (Wyatt).

Small angle X-ray scattering (SAXS) samples were prepared in a similar manner, whereby σNS was incubated with saturating amounts of 20mer or 40mer RNA (Supplementary Table S1), as described above. SAXS intensity data, I(q) versus momentum transfer q = 4πsinθ/λ, where θ is the scattering angle and λ is the wavelength, were collected using SEC-SAXS on beamline B21 at Diamond Light Source (Didcot, UK) over a range of q of 0.004 < q < 0.442 Å⁻¹. A total of 50 μl of each sample (~175 μM) was loaded onto a 2.4 ml Superdex 200 Increase 3.2 column mounted on Agilent HPLC High Pressure Liquid Chromatography (HPLC) system, and the eluent was flowed through the SAXS beam at 0.04 ml/min. The SEC buffer used as the background was collected after one SEC column volume. SAXS data were collected at 1 s intervals using a Pilatus 2M detector (Dectris, Switzerland) at a distance of 3.9 m and an X-ray wavelength of 1 Å. Guinier plot fit and real space inversions were performed using Prisum (20) and GNOM from the ATSAS software package v.2.8.3 (21,22). Radii of gyration (Rg) were estimated using AUTORIG (21). Low resolution envelopes were determined using the simulated annealing procedures implemented in DAMMIF (23) in slow mode, with no symmetry applied. Each scattering curve generated 20-independent models, which were averaged and filtered using DAMAVER (24) and DAMFILT with a mean normalized spatial discrepancy of 0.820 ± 0.05 (σNS apoprotein) and 0.576 ± 0.03 (σNS–20mer complex). SAXS experimental data together with the relevant experimental conditions and the derived models are available from SASBDB.

Native mass spectrometry

σNS was dialoged into 200 mM ammonium acetate, pH 7.6 overnight at 4°C. RNP complexes were assembled as described above. σNS apoprotein and σNS–RNP complex were diluted to a final concentration of 20 μM. NanoESI–IMS–MS spectra were acquired with a Synapt HDMS mass spectrometer (Waters) with platinum/gold-plated borosilicate capillaries prepared in-house. Typical instrument parameters were: capillary voltage, 1.2–1.6 kV; cone voltage, 40 V; trap collision voltage, 6 V; transfer collision voltage, 10 V; trap DC bias, 20 V; backing pressure, 4.5 mbar; IMS gas pressure, 0.5 mbar; traveling wave height, 7 V; and traveling wave velocity, 250 ms⁻¹. Data were processed with MassLynx v4.1, Driftscope 2.5 (Waters) and Massign (25). Collision cross-sections (CCSs) were estimated through a calibration (26–28) using arrival-time data for ions with known CCSs (β-lactoglobulin A, avidin, concanavalin A and yeast alcohol dehydrogenase, all from Sigma-Aldrich). The CCS values of the lowest observed charge state (and therefore the least affected by Coulombic repulsion (29)) were selected for comparison with SAXS-derived CCS estimates. Theoretical CCS values for SAXS ab initio reconstructions of σNS and σNS–RNP were generated using the calibrated projection approximation method in IMPACT (30). Estimated CCSs for the 20 independently generated dummy atom models were generated and averaged.
Affinity measurements by fluorescence anisotropy

Fluorescence anisotropy measurements with Alexa Fluor 488 dye-labeled RNAs (Supplementary Table S1) were performed at 25°C using a POLARstar Omega plate reader (BMG Labtech) in Greiner 384 well black polypropylene plates. Serial 2-fold dilutions of NSP2 and σNS were titrated into 5 nM RNA in 50 mM Tris–HCl pH 7.5, 50 mM NaCl, 1 mM ethylenediaminetetraacetic acid (EDTA), 0.05% Tween-20 in a total volume of 50 μl and equilibrated at room temperature for 15 min prior to measurements were taken. Where required, buffers were supplemented with 10 mM MgCl₂. Raw Anisotropy (r) values were calculated as follows:

\[ r = \frac{(I∥ - I⊥)}{(I∥ + 2I⊥)} \]  

(1)

Where \( I∥ \) and \( I⊥ \) are the parallel and perpendicular emission signals, respectively. Normalized anisotropy values were plotted as a function of protein concentration and fitted to a Hill equation using OriginPro 9.0.

Electrostatic contributions to free energies of protein–RNA interactions

The dependence of \( K_{\text{obs}} \) on buffer ionic strength can be expressed as:

\[ \log K_{\text{obs}} = -m'\psi \log [M^+] + \log K_{\text{obs,1M}} \]  

(2)

where \([M^+]\) is the monovalent counterion concentration (in this case Na⁺), \(m'\) is the number of ion pairs formed, \(\psi\) is defined as the thermodynamic extent of counterion binding and \(K_{\text{obs,1M}}\) is the non-electrostatic contribution to the dissociation constant, defined as the \(K_{\text{obs}}\) at 1 M NaCl, when the polyelectrolyte effect is minimal (31,32). The slope, \(SK_{\text{obs}}\), of log\(K_{\text{obs}}\) against log\([M^+]\) relates to the number of counterions released upon binding as follows:

\[ SK_{\text{obs}} = -\frac{\Delta \log K_{\text{obs}}}{\Delta \log[M^+]} = m'\psi \]  

(3)

The electrostatic (poly-electrolyte) contribution to free energy of binding, \(\Delta G_{\text{PE}}\), can therefore be determined as:

\[ \Delta G_{\text{PE}} = \Delta G_{\text{obs}} - \Delta G_{\text{aPE}} = -RT(SK_{\text{obs}}\log[M^+]) \]  

(4)

where \(\Delta G_{\text{obs}}\) is the total free energy of binding, and \(\Delta G_{\text{aPE}}\) is the non-electrostatic (non-polyelectrolyte) contribution. As the parameter \(\psi\) varies between different polynucleotides, we estimated the amount of salt bridges using empirically determined values, poly(U) = 0.68 and poly(A) = 0.78 as the upper and lower limits of salt bridges involved in complex formation (31,32). As NSP2 octamers disassemble under high ionic strengths (33), a range of 50–250 mM NaCl was used to determine RNA binding affinities. σNS, however, aggregates at low (<50 mM) NaCl concentrations, but is not affected by higher ionic strength conditions, so a wider range of ionic strength buffers were used to determine RNA binding affinities (up to 500 mM NaCl).

Electrophoretic mobility shift assays (EMSA) and in-gel FRET

A dual-labeled RNA stem-loop was designed with an MFE of folding of −8.9 kcal mol⁻¹, containing 3'-donor (ATTTO532) and 5'-acceptor (ATTTO647N) dye fluorophores as a FRET pair (Supplementary Table S1). Dual-labeled stem-loop was heat-annealed in binding buffer (50 mM Tris–HCl pH 7.5, 50 mM NaCl, 1 mM EDTA) at 75°C for 5 min and cooled to 4°C prior to incubation with increasing amounts of σNS (final RNA concentration 10 nM). A total of 10 μl of each sample was mixed with 2 μl 6x loading buffer (30% v/v glycerol, 5% Ficoll 400, 50 mM NaCl, 10 mM HEPES pH 8, 5 mM EDTA, 0.002% w/v bromophenol blue). Electrophoresis was carried out on a non-denaturing 1.5% agarose gel for 90 min at 100 V in 1× Tris-borate-EDTA buffer at 4°C. Gels were imaged using a fluorescence scanner (Fujifilm FLA-5100) with 532 nm excitation, imaging donor and acceptor wavelengths separately. The 2D densitometry was performed using ImageJ. Apparent FRET efficiencies (\(E_{\text{FRET(app)}}\)) were calculated as follows:

\[ E_{\text{FRET(app)}} = I_{\text{ATTO647N}}/(I_{\text{ATTO532}} + I_{\text{ATTO647N}}) \]  

(5)

Where \(I_{\text{ATTO532}}\) and \(I_{\text{ATTO647N}}\) are donor and acceptor emission intensities, respectively. For probing RNA:RNA interactions between S5f and S11f fragments (Supplementary Table S1), 100 nM of each RNA was individually heat-annealed at 85°C for 5 min in folding buffer (10 mM HEPES pH 7, 1 M NaCl, 10 mM MgCl₂) and snap-cooled to 4°C for 20 min, ensuring complete stem-loop folding prior to co-incubation. A total of 10 nM of S5f and S11f were incubated at 37°C in annealing buffer (10 mM HEPES pH 7, 100 mM NaCl, 1 mM MgCl₂) and immediately analyzed by electrophoresis on a native 15% acrylamide–TBE gel, run at 80 V at 4°C, and stained with 0.01% (w/v) SYBR gold. Gels were imaged using a fluorescence scanner as described above, using 488 nm excitation.

Fluorescence cross-correlation spectroscopy (FCCS)

Equimolar amounts of 18-nt Cy3- and Cy5-labeled non-complementary RNAs (10 nM each) (Supplementary Table S1) were incubated with varying concentrations of NSP2 and σNS in 50 mM NaCl, 20 mM HEPES pH 7.4. Interactions between S5 and S11 RNAs and S5 and Cy5-17mer (Supplementary Table S1) were measured as previously described (17). Briefly, 55 nM of each RNA strand was incubated with 5–10 μM NSP2 or σNS at 37°C for 15 min. Samples were then diluted into 150 mM NaCl, 20 mM HEPES pH 7.4, 0.05% Tween 20, resulting in a final RNA concentration of 1 nM each labeled RNA. NSP2 removal using proteinase K does not significantly reduce the amplitude of cross-correlation, suggesting that the observed cross-correlation is due to strand-annealing (17).

Circular dichroism (CD)

Circular dichroism (CD) experiments were performed on a Chirascan plus spectrometer (Applied Photophysics). Samples were prepared by dialyzing protein solutions against...
10 mM phosphate buffer pH 7.4, 50 mM sodium fluoride. Spectra were recorded over a wavelength range of 190–260 nm, with a bandwidth of 1 nm, step size of 1 nm and a path length of 1 mm. An average of three scans were used for the final spectra. Thermal stability was analyzed by monitoring the CD signal at 222 nm during heating from 20°C to 70°C with a heating rate of 1°C min⁻¹.

**Ensemble FRET**

Dual-labeled stem-loop (Supplementary Table S1) was heat-annealed at 75°C for 5 min and cooled to 4°C. Folded RNA-alone and denatured RNA (in 50% v/v formamide) were initially measured in 100 μl volumes at a final RNA concentration of 10 nM. Serial 2-fold dilutions of NSP2 and σNS from 15 μM were incubated with 10 nM RNA at room temperature for 15 min prior to measurement. Measurements were performed using a Fluorolog spectrofluorimeter (Horiba Jobin-Yvon). Apparent FRET efficiencies were calculated using Equation (5).

**Single-pair FRET measurements**

Dual-labeled RNA stem-loop used for ensemble FRET measurements was heat-annealed as described above and diluted to 10 pM final concentration. σNS and NSP2 were incubated with the stem-loop (25 nM each). Single-pair FRET measurements were performed on a custom-built confocal microscope with multiparameter fluorescence detection and pulsed interleaved excitation (PIE) (34) as described previously (35). Briefly, two picosecond pulsed lasers (532 and 640 nm) were operated at a repetition rate of 26.66 MHz and delayed by 18 ns with respect to each other to achieve rapid alternating excitation of donor and acceptor fluorophore at 100 μW laser power. By diluting the sample to picomolar concentrations, single molecule events were detected as bursts of fluorescence as they diffuse through the confocal volume on the millisecond timescale. Bursts were selected using a sliding time window burst search (36) with a count rate threshold of 10 kHz, a time window of 500 μs and a minimum of 100 photons per burst. Using time-correlated single photon counting and polarized detection, one can calculate for every molecule its FRET efficiency, labeling stoichiometry and the fluorescence lifetime and anisotropy of the donor and acceptor fluorophores (34). To remove molecules lacking the donor or acceptor dye, we used the ALEX-2CDE filter (37) with a time constant of 100 μs and an upper threshold of 10. Accurate FRET efficiencies, E, were calculated from background-corrected photon counts in the donor channel and acceptor channel after donor excitation (FDD/FDA) and acceptor channel after acceptor excitation (FAA) by correcting for crosstalk (α = 0.03), direct excitation (δ = 0.06) and differences in the quantum yield and detection efficiency (γ = 0.65):

\[
E = \frac{F_{DA} - \alpha F_{DD} - \delta F_{AA}}{\gamma F_{DD} + F_{DA} - \alpha F_{DD} - \delta F_{AA}}
\]

Species-selective fluorescence correlation functions were determined as follows: sub-populations of molecules were selected using FRET efficiency thresholds (NSP2: low-FRET E < 0.4, high-FRET E > 0.6; σNS: low-FRET E < 0.15, medium-FRET 0.3 < E < 0.8, high-FRET E > 0.9). For every burst, photons in a time window of 50 ms around the edges of the burst were added. If another single molecule event was found in the time window, the respective burst is excluded from the analysis. Correlation functions were calculated for each individual burst using acceptor photons after acceptor excitation to ensure that the obtained correlation functions are independent of the FRET efficiency. Species-selective correlation functions were then averaged to obtain the burst-selective correlation function (38, 39). All analysis was performed using the PAM software package (40).

**Raman spectroscopy**

Raman spectra of RNA, σNS and RNP and their corresponding buffers were acquired on a modular multi-channel Raman spectrograph Jobin Yvon–Spex 270M in 90° scattering geometry using 532 nm line of a continuous-wave solid-state Nd:YVO₄ laser for excitation (power of 240 mW at the sample), as described in detail elsewhere (41). Raman measurements were performed in a temperature-controlled hermetrical quartz microcell (4 μl volume) at 20°C and 60°C. Final spectra represent averages of 30–720 individually acquired and analyzed scans (depending on the sample type) each of 1 min integration time to notice any spectral changes during laser exposure and to increase signal-to-noise ratio without mathematical smoothing. Wavenumber scales were precisely calibrated (±0.1 cm⁻¹) using the emission spectra of a neon glow lamp taken before and after each Raman measurement. The Raman contribution from corresponding buffer was subtracted, and the spectra were corrected for non-Raman background.

**RESULTS**

σNS is unable to promote inter-segment interactions between RV RNAs

Recently, we have demonstrated that NSP2 can selectively promote RNA–RNA duplex formation between genomic ssRNAs in RVs (17). Both NSP2 and σNS possess helix-destabilizing and strand-annealing activities in vitro (17, 18). Using a previously established RNA–RNA interaction between RV segment S5 and S11 ssRNAs (17), we compared the abilities of NSP2 and σNS to promote inter-segment duplex formation. We employed fluorescence cross-correlation spectroscopy (FCCS) to monitor intersegment RNA–RNA interactions.

While S5 and S11 do not interact in the absence of protein (Figure 1A, magenta), co-incubation of S5 and S11 with NSP2 results in inter-molecular RNA duplex formation (Figure 1A, blue). In contrast, co-incubation of S5 and S11 RNAs with σNS does not promote duplex formation (Figure 1A, black). Binding of both NSP2 and σNS to either RNA results in an increase in apparent diffusion time, confirming that both proteins interact with S5 and S11 and form larger RNP complexes (Supplementary Figure S1). Hence, both proteins can bind to these RNAs, but only...
NSP2 can mediate a sequence-specific interaction between RV genomic ssRNAs.

Given that σNS promotes strand-annealing of short RNA oligonucleotides (18), its failure to promote interactions between full-length genomic ssRNAs may be due to sequestration of complementary sequences within RNA secondary structure. We compared the strand-annealing activities of both proteins using shorter complementary RNA fragments derived from S5 (S5f, nucleotides 299–350) and S11 (S11f, nucleotides 31–77). Incubation of S5f and S11f (10 nM each) resulted in spontaneous hybridization (Supplementary Figure S2). To limit this, we investigated interactions between a full-length S5 and an unstructured S11-derived 17mer complementary to S5 (Supplementary Table S1). NSP2 increases the amount of interacting S5 : 17mer (Figure 1B, blue), while σNS was unable to promote this interaction beyond the level of spontaneous annealing (Figure 1B, black and magenta, respectively).

As RNA annealing activity typically involves simultaneous binding of two RNAs, we next examined whether NSP2 and σNS can bind multiple RNAs in solution. To distinguish strand-annealing from RNA binding, we designed differently-labeled non-complementary 17mer RNAs (Supplementary Table S1). In the absence of protein, these RNAs did not interact (Figure 1C and D, magenta). Incubation of either NSP2 or σNS with an equimolar mixture of distinct RNAs at 2:1 to 4:1 protein oligomer : RNA ratio resulted in the largest fraction of co-diffusing oligomer-bound ssRNAs, indicating that both NSP2 and σNS can bind multiple unstructured RNAs. Having established that both proteins bind multiple RNAs with similarly high affinity, the observed failure of σNS to assist annealing of the 17mer to S5 ssRNA could be explained by its inability to remodel the target RNA sequence and increase its accessibility.

σNS undergoes an RNA-induced hexamer-to-octamer transition

Having established that both NSP2 and σNS can bind multiple RNAs per oligomer, we further investigated σNS–RNP complex formation. NSP2 is a stable octamer in the presence or absence of RNA (11,33,42), while the σNS apoprotein is hexameric, although RNP stoichiometry is unknown (18). We incubated σNS hexamer with a stoichiometric excess of 20mer RNA (‘Materials and Methods’ section) and analyzed the RNP complexes using size-exclusion chromatography and dynamic light scattering (DLS). Surprisingly, σNS–RNP eluted earlier and had a greater hydrodynamic radius (R_h) of ~8 nm than the apoprotein (R_h ~ 5 nm) (Figure 2A). This difference in R_h cannot be explained by the binding of multiple 20mer RNA molecules...
RNA binding results in assembly of a larger σNS oligomer. (A) Size-exclusion chromatography elution profiles of σNS apoprotein (blue) and σNS–RNP complex (red). Absorbances at 260 and 280 nm are shown as dashed and continuous lines, respectively. DLS-derived hydrodynamic radii (R_d) are shown for each species. A second, minor peak corresponds to free, excess RNA. (B) SAXS curves for σNS apoprotein (blue) and RNP complex (red), with respective fits shown in black. Scattering profiles are shown as the logarithm of the scattering intensity, I, as a function of the momentum transfer, q = 4πsin(θ)/λ. Radii of gyration (R_g) values of both species are displayed. (C) Normalized pair-wise distance distribution functions, P(r), calculated from the scattering curves of σNS apoprotein (blue) and RNP complex (red) showing an increase in maximum dimension (D_max). alone, suggesting that σNS undergoes a change in conformation or its oligomeric state.

We then examined RNP complex formation by SAXS. Radii of gyration (R_g) values for σNS apoprotein and RNP complex were 5.5 ± 0.03 nm and 7.6 ± 0.05 nm, respectively (Figure 2B). Guinier region analysis suggests that both σNS and σNS–RNP samples are monodisperse, confirming that the observed increase in size is not due to σNS aggregation when bound to RNA (Supplementary Figure S3A). Upon RNA binding, hexameric σNS further undergoes a 2-fold increase in its maximum distance (D_max) value from 19.9 to 41.6 nm (Figure 2C). Further analysis of SAXS data indicates that both NSP2 and σNS are globular, however σNS–RNP complex formation results in a flexible, elongated particle (Supplementary Figure S3B). No further increase in size of the RNP was observed after incubation with a 40mer RNA (Supplementary Figure S3C). The assembled σNS–RNP complex appears to be more stable than σNS apoprotein, while NSP2 exhibited significantly decreased stability upon RNA binding (Supplementary Figure S4), explaining the severe aggregation of NSP2-RNP that precluded its characterization by SAXS. Together, these data suggest that σNS undergoes an RNA-driven oligomerization that is independent of the substrate RNA length.

To analyze the stoichiometries of σNS–RNP complexes, we used native electrospray ionization—ion mobility spectrometry—mass spectrometry (ESI-IMS-MS). A typical ESI-MS spectrum shows σNS hexamers along with additional oligomeric species, in agreement with previous analysis of σNS (18) (Figure 3A and Supplementary Figure S5). In contrast, the RNP contains a large population of octameric species with two RNAs bound (Figure 3B). A small fraction of hexameric species was bound to a single RNA. Together, IMS-MS and SAXS data suggest that σNS undergoes a hexamer-to-octamer transition upon RNP complex formation (Figure 3).

We then used ion-mobility mass-spectrometry to estimate rotationally averaged collision cross-sections (CCS) for each observed species (43). We compared CCSs derived from the ESI-IMS-MS spectra with the CCS values of σNS apoprotein and RNP complex calculated for SAXS-derived models, as described in the ‘Materials and Methods’ section

Figure 3. σNS RNP complex is predominantly octameric. (A and B) Native ESI-MS of σNS apoprotein (A) and RNP complex (B). Averaged collision cross-sections, CCS (Ω), are shown in nm² for each species. Inset: relative abundances of hexameric and octameric σNS oligomers. Smaller protein oligomers observed in both spectra are due to dissociation of higher order species during the ionization process.
Figure 4. σNS undergoes a hexamer-to-octamer transition upon binding RNA. (A and B) SAXS-derived ab initio models of hexameric σNS and σNS–RNP complex. Twenty best models for each σNS apoprotein species (A) and σNS–RNP (B) were generated as described in ‘Materials and Methods’ section, averaged using DAMAVER (light mesh) and filtered using DAMFILT (superimposed dark surface). (C) CCS of σNS oligomers detected by ESI-IMS-MS. σNS apoprotein species are shown in blue and σNS–RNP are shown in red. Dashed horizontal lines denote CCS values estimated for the SAXS models of σNS apoprotein and the RNP complex, shown in (A) and (B), respectively. Masses of each oligomer and their charge states are summarized in Supplementary Tables S4 and 5.

1.0 nm²). Thus, simultaneous binding of two RNAs results in the formation of stable, elongated octameric σNS–RNP complexes.

Octameric σNS disrupts RNA structures more efficiently than its hexameric form

We then investigated the relationship between σNS oligomeric state and its helix-destabilizing activity. We designed a dual-labeled 36mer RNA stem-loop containing 3′-Atto532 and 5′Atto647N fluorophores (see ‘Materials and Methods’ section, Figure 5A), and incubated it with increasing amounts of σNS. Using electrophoretic mobility shift assays (EMSAs) to separate free RNAs from assembled RNP complexes (Figure 5B and C), we compared the helix-destabilizing activities of different σNS oligomers by estimating the FRET efficiencies of each band-shift (44,45) (Figure 5D and Supplementary Figure S7). Titrations of σNS into this stem-loop produced three shifts that sequentially occurred at higher protein concentrations, corresponding to multiple oligomeric species. Given the presence of RNA-bound hexamers and octamers observed in the ESI-IMS-MS spectrum (Figure 3), we interpreted the first shift as hexameric σNS and the second shift as octameric σNS. The protein-free RNA had an apparent FRET efficiency (EFRET(app)) of 0.73 ± 0.03, suggesting that the stem-loop is folded. The RNA hexameric shift had EFRET(app) of 0.48 ± 0.11, and this value decreased further for the octameric shift (EFRET(app) = 0.28 ± 0.05), indicating that octameric σNS has a greater capacity for destabilizing RNA structure than hexameric σNS (Figure 5D).

Given the RNA-binding footprint of σNS of ~20 nt (18,46), a third shift was observed at elevated σNS concentrations (>2 μM). This shift does not occur when σNS binds 20 nt RNA (Supplementary Figure S8), confirming that this shift is due to two σNS oligomers bound to the same 36 nt RNA. Similar protein saturation of 40mer RNA resulted in formation of a mixed population of higher-order species, as observed in SAXS (Supplementary Figure S9).

NSP2 and σNS differ in modes of RNA unfolding

Having examined the helix-destabilizing activity of σNS oligomers, we then investigated whether oligomer binding to a structured RNA is coupled to its unwinding activity. We used single-pair (sp)FRET to analyze the FRET states of discrete populations of protein-free and oligomer-bound RNAs using the dual-labeled RNA construct described above (Figure 5A). For stem-loop alone, a single high-FRET population is present (Figure 6A, gray histogram), in agreement with the MFE prediction that the RNA forms a stable hairpin with its 3′ and 5′ termini in proximity. In the presence of NSP2, two distinct FRET populations were observed—a high-FRET population similar to that of the stem-loop alone, and a low-FRET population (Figure 6A). Species-selective correlation analysis reveals that NSP2 is bound to RNA in both FRET populations (Figure 6B), indicating that NSP2 can bind to both folded and unfolded RNA stem-loops.

Similarly, we analyzed RNA stem-loop destabilization by σNS (Figure 6C). While the high-FRET population persists in the presence of σNS, there are also a range of lower FRET populations, notably an intermediate-FRET and a low-FRET population. The intermediate-FRET population is dynamic on a sub-millisecond time scale, revealing an ensemble of partially unwound, inter-converting σNS-bound RNA structures (Supplementary Figure S10). This
**Figure 5.** αNS oligomers have different helix-unwinding activities. (A) Structure of the dual-labeled RNA stem-loop with 5′-donor (‘D’) and 3′-acceptor (‘A’) fluorophores, used for helix-unwinding assays monitored by FRET. (B and C) EMSA of the dual-labeled stem-loop bound to αNS oligomers. Multiple shifts occur (blue, green and yellow asterisks) as αNS concentration increases. (D) Helix-unwinding activities of different αNS oligomers formed at increasing αNS concentration. Apparent in-gel FRET efficiencies ($E_{FRET(app)}$) of the unbound, hexamer-bound and octamer-bound RNA stem-loops were estimated for each band-shift shown in (B).

is in agreement with previous results suggesting that even at higher protein concentrations (>5 μM), αNS was unable to induce a single, fully unfolded RNA population (18). Species-selective correlation analysis indicates that αNS is bound to RNA in both the intermediate and low-FRET populations, but not the high-FRET (Figure 6D).

Ensemble FRET experiments conducted at a wider range of RNA and protein concentrations further demonstrate that NSP2 is more efficient at destabilizing RNA than αNS (Supplementary Figure S11). Collectively, these data suggest that NSP2 and αNS have different modes of RNA helix-destabilization. While NSP2 can bind to the folded stem-loop, its binding may not necessarily result in RNA unwinding. In contrast, αNS binding induces a range of unfolded and partially folded RNA conformations.

**NSP2 and αNS discriminate between RNA structures based on their relative stabilities**

As RNA unfolding activities of both proteins require ss-RNA binding, we investigated the relationship between RNA structure and binding affinity of NSP2 and αNS. We designed three fluorescently-labeled 20mer RNAs with different thermodynamic stabilities for use in binding measurements: an unstructured RNA, a metastable RNA ($ΔG = −3.8 \text{ kcal mol}^{-1}$), and a stable hairpin structure ($ΔG = −8.1 \text{ kcal mol}^{-1}$) (Figure 7A). All three RNAs bind NSP2 with near-identical affinities ($K_D = 20 ± 3.2 \text{ nM}$, Supplementary Table S2), indicating that NSP2 binds ssRNA independent of structure (Figure 7B). αNS binds the unstructured and metastable RNAs with similar affinities ($K_D = 37 ± 1.5 \text{ nM}$ and $24 ± 2.6 \text{ nM}$, respectively, Supplementary Table S2) but had lower affinity for the stable RNA ($K_D = 137 ± 1.9 \text{ nM}$) (Figure 7D).

To further investigate RNA structural preferences of either protein, we then examined the affinities of NSP2 and αNS for these RNAs in the presence of Mg$^{2+}$ ions, which stabilize RNA structures (47,48). Although the $K_D$ values of both NSP2 and αNS for unstructured RNA remains largely unchanged (1.4- and 1.5-fold increase, respectively), there is a 10- to 30-fold decrease in affinity of both NSP2 and αNS for the metastable and stable RNAs (Figure 7C and E; Supplementary Table S2). This suggests that although both
Figure 6. Helix-destabilizing activities of αNS and NSP2, examined by single-pair FRET (spFRET). (A) Histogram of spFRET efficiency of the dual-labeled RNA stem-loop (10 pM, shown in gray), and in the presence of 25 nM NSP2 (orange). (B) Species-selective correlation analysis of the high-FRET (green autocorrelation function, ACF) and low-FRET (red ACF) populations, and freely diffusing RNA (orange). A typical ACF of a freely diffusing stem-loop is shown in gray (RNA*). Note rightward shift in ACFs of protein-bound stem-loops due to slower diffusion. (C) Histogram of spFRET efficiency of the dual-labeled RNA stem-loop (see panel A), alone (gray) and in the presence of 25 nM αNS (blue). (D) Species-selective correlation analysis of the high-FRET (blue ACF), intermediate FRET (green ACF) and low-FRET (red ACF) populations, and freely diffusing folded RNA (high-FRET, orange). A typical ACF of a freely diffusing stem-loop is shown in gray (RNA*). Only intermediate and low-FRET species are bound to αNS.

Figure 7. Stability of RNA structure determines preferential binding by NSP2 and αNS. (A) Fluorescently-labeled unstructured (red), metastable (blue) and stable (green) 20-mer RNAs used for fluorescence anisotropy binding assays. (B and D) NSP2 binds unstructured and stable RNAs with similar affinities. In contrast, stable secondary structure impedes αNS binding. (C and E) Mg²⁺-dependent stabilization of RNA structure impairs binding of ssRNAs by both NSP2 and αNS. Note the apparent affinity of both proteins for unstructured 20-mer remains largely unchanged upon addition of 10 mM MgCl₂. Due to NSP2 aggregation at higher concentrations, protein titrations were only performed with [NSP2] up to 2 μM (C).
NSP2 and σNS exhibit preferential binding to unstructured RNAs. NSP2 can bind stable hairpins better than σNS. This reduction in affinity cannot be explained by Mg$^{2+}$-induced dissociation of protein oligomers (33), as binding affinities of either protein for unstructured RNA remain largely unaffected in the presence of Mg$^{2+}$.

Previous structural studies of NSP2 have demonstrated that sequence-independent ssRNA-binding occurs via a positively-charged groove (11,42). Although there is no such structural information for σNS, NSP2 and σNS both bind ssRNA with high affinity and without apparent sequence specificity, potentially via multiple electrostatic contacts. We therefore examined the dependency of binding affinity on ionic strength by measuring the dissociation constant of NSP2 and σNS for an unstructured 20mer ssRNA molecule at different ionic strength (Figure 8). We observed expected relationships (Eqs. 2-3) between binding affinity and ionic strength, indicating that NSP2 binding involves at most two salt bridges, whereas σNS binds RNA via 3–4 ion contacts (Figure 8C) (32). Under physiological ionic strength, NSP2 binds ssRNA with apparent free energy of binding, $\Delta G = -9.52$ kcal mol$^{-1}$, of which there is an estimated $\sim$17.5% electrostatic component (Supplementary Table S3). Under the same conditions, σNS binds RNA with $\Delta G = -9.50$ kcal mol$^{-1}$, of which there is a $\sim$30.4% electrostatic contribution (Supplementary Table S3). This indicates that despite their near-identical affinities for RNA, σNS has a greater electrostatic contribution to the overall free energy of binding relative to that of NSP2. Together, these results suggest that NSP2 and σNS interact with RNAs differently, allowing them to discriminate between ssRNAs based on their propensities to form stable secondary structures (Figure 9).

**DISCUSSION**

Inter-molecular RNA-RNA interactions have been postulated to underpin the selection and assembly of multi-segmented genomes in viruses comprising the *Reoviridae* family (5,17,49,50). Recent studies on RVs suggest that NSP2 promotes RNA-RNA interactions between its full-length genomic ssRNA segment precursors (17). Both NSP2 and σNS facilitate conformational rearrangements of ssRNAs *in vitro* (17,18) and mediate formation of inter-molecular RNA contacts between short, synthetic RNA fragments, including those derived from ARV genome and containing complementary sequences that may be involved in assembly of multiple RNAs (18). However, such interacting sequences have not yet been identified in mammalian or ARVs, and have only recently been mapped for a number of inter-segment contacts within the RV genome (17). Therefore we employed these RV RNAs as a model to directly compare the abilities of RV NSP2 and ARV σNS to mediate RNA–RNA interactions within the context of full-length RNA genomic precursors.

Here we have shown that despite the ostensibly similar RNA chaperone-like activities of NSP2 and σNS, only NSP2 is capable of promoting inter-segment interactions between RV RNAs. To gain insights into the mechanisms underpinning this selective, protein-mediated strand- annealing reaction, we performed a side-by-side comparison of the RNA binding, helix-destabilizing and strand-annealing activities of both proteins.

Previous studies conducted with ssRNA substrates suggest that both NSP2 and σNS bind ssRNA without apparent sequence specificity (11,12,18,33,42,46). Using RNA substrates with different secondary structure stabilities, here we have demonstrated that σNS displays reduced affinity toward highly stable hairpins, whereas NSP2 binds ssRNAs regardless of their propensity to form secondary structure. Further stabilization of RNA structure with Mg$^{2+}$ ions reveals that both proteins indeed preferentially bind unstructured RNAs. The observed change in the affinity is consistent with previously reported inhibition of the strand displacement activity of σNS in the presence of MgCl$_2$ (42). NSP2 was shown to dissociate into smaller oligomers in the presence of magnesium (33). However, NSP2 binding to unstructured 20-mer RNA in the presence of 10 mM MgCl$_2$ remains largely unaffected (Figure 7C and E), strongly suggesting that the observed inhibitory effects of Mg$^{2+}$ ions are due to RNA secondary structure stabilization. These structural preferences of NSP2 and σNS are consistent with the observed helix-destabilization upon protein binding to a stable RNA stem-loop. Incubation of the stem-loop with large molar excess of either protein results in the RNA helix disruption (Supplementary Figure S11). However, spFRET and fluorescence correlation spectroscopy (FCS) analyses reveal that NSP2 can efficiently bind both folded (high FRET) and unfolded (low FRET) RNA stem-loops. The initial NSP2 binding may not necessarily result in substantial RNA unwinding (Figure 6), which is ultimately achieved at higher protein concentrations (Supplementary Figure S11). In contrast, σNS binding results in gradual unfolding, producing a mixed population of partially unfolded RNA intermediates, failing to completely unfold RNA even at high molar excess, further indicating that both proteins have different modes of helix disruption (Figure 6).

Given that helix-destabilizing activity of both proteins is likely to be coupled to ssRNA binding, we examined how these proteins interact with unstructured RNAs. While NSP2 interacts with two RNAs as an octamer (33), σNS hexamer only binds a single RNA and undergoes a hexamer-to-octamer transition that appears to be a prerequisite for binding a second RNA molecule. Octamer formation is concomitant with the increased helix-destabilizing activity, potentially providing additional RNA-binding surface (51,52), hence increased capacity to compete with RNA secondary structure formation (Figure 9). Thus, failure of σNS to promote a specific inter-segment duplex formation can be attributed to its reduced capacity to interact with and disrupt stable intramolecular RNA structures.

**RNA secondary structure stability modulates chaperone activity**

Given the mechanistic differences between NSP2 and σNS, it appears that a major determinant of efficient protein-mediated inter-segment annealing is RNA structural stability. Intramolecular RNA structure can regulate binding by cognate RNA chaperone proteins, therefore affecting refolding and inter-molecular duplex annealing. Thus, RNA
secondary structure stability together with chaperone binding mode may serve to fine-tune the matchmaking activities of RNA chaperones that would otherwise interact with RNA without sequence preference (53,54). This may regulate specific RNA–RNA interactions required for a highly accurate assembly of a complete segmented RNA genome. Other RNA chaperones, including *Escherichia coli* StpA and Moloney murine leukemia virus NC, both of which bind preferentially to unstructured regions of RNA (55,56), exhibit similar relationships between chaperone activity and RNA structural stability (57,58). This suggests that fine-tuning of RNA remodeling by its stability may be a general feature of RNA chaperone activity.

RNA-driven oligomerization

Upon binding of two ssRNAs hexameric αNS assembles into octameric RNP complexes with defined stoichiometry. This mechanism of RNA-driven oligomerization is distinct from that of other viral RNA-binding proteins that assemble into large, non-discrete, higher-order oligomers in the presence of RNA (59–62). We propose that αNS hexamer assembles as a trimer of dimers (18), in dynamic equilibrium with a low octameric population (Figure 3). RNA binding results in equilibrium shift toward octameric species, which appear to be stabilized by RNP complex formation. This represents a novel mechanism of modulating RNA chaperone activity, whereby the oligomeric state of the protein defines its unwinding efficiency. Similar to ARV αNS, mammalian reovirus αNS and bluetongue virus NS2 have also been reported to exist in a range of oligomeric states (63–66). It is possible that different αNS oligomers may play distinct roles during viral replication.

Other functionally analogous proteins encoded by different members of the *Reoviridae* family form octamers, similar to NSP2 (notably P9-1 and Pns9 proteins) (67–69). Despite their similar toroidal architecture, NSP2 presents a continuous, basic RNA binding groove on the surface, while P9-1 binds ssRNA within a positively charged inner pore. Analysis of RNA binding by both NSP2 and αNS also reveals different number of salt bridges involved in protein–RNA interactions (Figure 8), with each protein predominantly interacting non-electrostatically. Such sequence-independent, non-electrostatic interactions with ribose and base moieties may explain the preference of both NSP2 and αNS for unfolded ssRNA over dsRNA substrates that are only accessible via the A-form backbone. Indeed, Raman difference analysis of αNS-bound RNA (Supplementary Figure S12) reveals decrease in the band inten-
distributed novel analytical tools and analyzed ESI-IMS-MS. In addition to this, the authors conducted experiments, analyzed data and jointly wrote the manuscript. A. Bo and R.T. managed the project. P.M. collected and analyzed Raman spectroscopy data. All authors contributed ideas, discussed the results and were involved in writing of the manuscript.

FUNDING

Wellcome Trust [103068/Z/13/Z to A.B.]; Biotechnology and Biological Sciences Research Council (BBSRC) White Rose DTP [BB/M011151/1 to J.P.K.B.]; European Regional Development Fund [CZ.02.1.01/0.0/0.0/15_003/0000441 to R.T.]; BBSRC [BB/P000037/1 to A.N.C., BB/E012558/1]; Deutsche Forschungsgemeinschaft (Project B3) [SFB1032 to D.C.L.]; Ludwig-Maximilians-Universität, München through the Center for NanoScience (CeNS) (to D.C.L.); FEBS (to A.B.); FEMS (to A.B.); Microbiology Society (to A.B.). Funding for open access charge: Wellcome Trust; BBSRC. Conflict of interest statement. None declared.

REFERENCES

1. Desselberger, U. (2014) Rotaviruses. Virus Res., 190, 75–96.
2. Mertens, P. (2004) The dsRNA viruses. Virus Res., 101, 1–13.
3. McDonald, S.M., Nelson, M.I., Turner, P.E. and Patton, J.T. (2016) Reassortment in segmented RNA viruses: mechanisms and outcomes. Nat. Rev. Microbiol., 14, 448–460.
4. McDonald, S.M. and Patton, J.T. (2011) Assortment and packaging of the segmented rotavirus genome. Trends Microbiol., 19, 136–144.
5. Sung,P.Y. and Roy, P. (2014) Sequential packaging of RNA genomic segments during the assembly of bluetongue virus. Nucleic Acids Res., 42, 13824–13838.
6. Lourenço,S. and Roy, P. (2011) In vitro reconstitution of Bluetongue virus infectious cores. Proc. Natl. Acad. Sci. U.S.A., 108, 13746–13751.
7. Anzola,J. V, Xu,Z.K., Asamizu,T. and Nuss,D.L. (1987) Segment-specific inverted repeats found adjacent to conserved terminal sequences in wound tumor virus genome and defective interfering RNAs. Proc. Natl. Acad. Sci. U.S.A., 84, 8301–8305.
8. Touris-Otero,F., Cortez-San Martin,M., Martínez-Costas,J. and Benavente,J. (2004) Avian reovirus morphogenesis occurs within viral factories and begins with the selective recruitment of σNS and λAA to μNS inclusions. J. Mol. Biol., 341, 361–374.
9. Netherton,C.L. and Wileman,T. (2011) Virus factories, double subcompartmentalization of the host translational machinery. EMBO, 30, 1–11.
10. Miller,C.L., Arnold,M.M., Broering,T.J., Hastings,C.E. and Nibert,M.L. (2010) Localization of mammalian orthoreovirus NS–RNP, SASDDU5 (σNS–RNP), SASDDU5 (σNS apoprotein).

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.
proteins to cytoplasmic factory-like structures via nonoverlapping regions of microNS. J. Virol., 84, 867–882.

17. Borodavka, A., Dykeman, E.C., Schirmpf, W. and Lamb, D.C. (2017) Protein-mediated RNA folding governs sequence-specific interactions between rotavirus genome segments. Elife, 6, 1–22.

18. Borodavka, A., Ault, J., Stockley, P.G. and Tuma, R. (2015) Evidence that avian reovirus NS is an RNA chaperone: implications for genome segment assortment. Nucleic Acids Res., 43, 7044–7057.

19. Borodavka, A., Singaram, S.W., Stockley, P.G., Gelbart, W.M., Ben-Shaul, A. and Tuma, R. (2016) Sizes of long RNA molecules are determined by the branching patterns of their secondary structures. Biophys. J., 111, 2077–2085.

20. Konarev, P.V., Volkov, V.V., Sokolova, A.V., Koch, M.H.J. and Svergun, D.I. (2003) PRIMUS: a Windows PC-based system for small-angle scattering data analysis. J. Appl. Crystallogr., 36, 1277–1282.

21. Konarev, P.V., Potoukhov, M.V., Volkov, V.V. and Svergun, D.I. (2006) ATSAS 2.1, a program package for small-angle scattering data analysis. J. Appl. Crystallogr., 39, 277–286.

22. Potoukhov, M.V., Franke, D., Shkumatov, A.V., Tria, G., Kikhney, A.G., Gajda, M., Gorba, C., Mertens, H.D.T., Konarev, P.V. and Svergun, D.I. (2012) New developments in the ATSAS program package for small-angle scattering data analysis. J. Appl. Crystallogr., 45, 342–350.

23. Franke, D. and Svergun, D.I. (2009) DAMMIF, a program for rapid ab-initio shape determination in small-angle scattering. J. Appl. Crystallogr., 42, 342–346.

24. Volkov, V.V. and Svergun, D.I. (2003) Uniqueness of ab initio shape determination in small-angle scattering. J. Appl. Crystallogr., 36, 860–864.

25. Morgner, N. and Robinson, C.V. (2012) Massigen: An assignment strategy for maximizing information from the mass spectra of heterogeneous protein assemblies. Anal. Chem., 84, 2939–2948.

26. Ruotolo, B.T., Bencsich, J.L.P., Sandercock, A.M., Hyung, S.-J. and Robinson, C.V. (2008) Ion mobility–mass spectrometry analysis of large protein complexes. Nat. Protoc., 3, 1139–1152.

27. Bush, M.F., Hall, Z., Giles, K., Hoyes, J., Robinson, C.V. and Ruotolo, B.T. (2010) Collision cross sections of proteins and their complexes: a calibration framework and database for gas-phase structural biology. Anal. Chem., 82, 9557–9565.

28. Smith, D., Knapman, T., Campuzano, I., Malham, R., Berryman, J., Radford, S.E. and Ashcroft, A. (2009) Deciphering drift time measurements from travelling wave ion mobility spectrometry-mass spectrometry studies. Eur. J. Mass Spectrom., 15, 113–130.

29. Ruotolo, B.T. and Robinson, C.V. (2006) Aspects of native proteins are retained in vacuum. Curr. Opin. Chem. Biol., 10, 402–408.

30. Marklund, E.G., Degiacomi, M.T., Robinson, C.V., Baldwin, A.J. and Bencsich, J.L.P. (2015) Collision cross sections for structural proteomics. Structure, 23, 791–799.

31. Davidovich, C., Zheng, L., Goodrich, K.J. and Cech, T.R. (2013) Promiscuous RNA binding by polycomb repressive complex 2. Nat. Struct. Mol. Biol., 20, 1250–1257.

32. Record, M.T., Lohman, T.M. and Haseth, P. de (1976) Ion effects on ligand-nucleic acid interactions. J. Mol. Biol., 107, 145–158.

33. Schuck, P., Taraporewala, Z., McPhie, P. and Patton, J.T. (2001) Rotavirus nonstructural protein NSP2 Self-assembles into octamers that undergo Ligand-induced conformational changes. J. Biol. Chem., 276, 9679–9687.

34. Kudryavtsev, V., Sikor, M., Kalinov, S., Mokranjac, D., Seidel, C.A.M. and Lamb, D.C. (2012) Combining MFD and PIE for accurate single-pair Förster resonance energy transfer measurements. Chemphyschem, 13, 1060–1078.

35. Voith von Voithenberg, L., Sánchez-Rico, C., Kang, H.-S., Madl, T., Zanier, K., Barth, A., Warner, L.R., Sattler, M. and Lamb, D.C. (2016) Recognition of the 3′ splice site RNA by the U2AF heterodimer involves a dynamic population shift. Proc. Natl. Acad. Sci. U.S.A., 113, E1769–E1775.

36. Nir, E., Michaela, X., Hamadani, K., Laurence, T.A., Neuhauser, D., Kovechov, Y. and Weiss, S. (2006) Shot-noise limited single-molecule FRET histograms: comparison between theory and experiments. J. Phys. Chem. B, 110, 22103–22124.

37. Tomov, T.E., Tsukano, R., Masoud, R., Libcr, M., Plavner, N. and Nir, E. (2012) Disentangling subpopulations in single-molecule FRET and ALEX experiments with photon distribution analysis. Biophys. J., 102, 1163–1173.

38. Eggeling, C., Fries, J.R., Brand, L., Gunther, R. and Seidel, C.A.M. (1998) Monitoring conformational dynamics of a single molecule by selective fluorescence spectroscopy. Proc. Natl. Acad. Sci. U.S.A., 95, 1556–1561.

39. Laurence, T.A., Kwon, Y., Yin, E., Hollars, C.W., Camarero, J.A. and Barsky, D. (2007) Correlation spectroscopy of minor fluorescent species: signal purification and distribution analysis. Biophys. J., 92, 2184–2198.

40. Schlimp, Waldemar (2018) PAM: A framework for integrated analysis of imaging, Single-Molecule, and ensemble fluorescence data. Biophys. J., 114, 518–518.
60. Reguera, J., Cusack, S. and Kolakofsky, D. (2014) Segmented negative strand RNA virus nucleoprotein structure. *Curr. Opin. Virol.*, 5, 7–15.
61. Durham, A.C., Finch, J.T. and Klug, A. (1971) States of aggregation of tobacco mosaic virus protein. *Nat. New Biol.*, 229, 37–42.
62. Butler, P.J.G. and Klug, A. (1971) Assembly of the particle of tobacco mosaic virus from RNA and disks of protein. *Nat. New Biol.*, 229, 47–50.
63. Mumtsidu, E., Makhov, A.M., Roessle, M., Bathke, A. and Tucker, P.A. (2007) Structural features of the Bluetongue virus NS2 protein. *J. Struct. Biol.*, 160, 157–167.
64. Butan, C. and Tucker, P. (2010) Insights into the role of the non-structural protein 2 (NS2) in Bluetongue virus morphogenesis. *Virus Res.*, 151, 109–117.
65. Butan, C., Van Der Zandt, H. and Tucker, P.A. (2004) Structure and assembly of the RNA binding domain of bluetongue virus non-structural protein 2. *J. Biol. Chem.*, 279, 37613–37621.
66. Gillian, A.L. and Nibert, M.L. (1998) Amino terminus of reovirus nonstructural protein sigma NS is important for ssRNA binding and nucleoprotein complex formation. *Virology*, 240, 1–11.
67. Akita, F., Higashiura, A., Shimizu, T., Pu, Y., Suzuki, M., Uehara-Ichiki, T., Sasaya, T., Kanamaru, S., Ariasaka, F., Tsukihara, T. et al. (2012) Crystallographic analysis reveals octamerization of viroplasm matrix protein P9-1 of Rice black streaked dwarf virus. *J. Virol.*, 86, 746–756.
68. Wu, J., Li, J., Mao, X., Wang, W., Cheng, Z., Zhou, Y., Zhou, X. and Tao, X. (2013) Viroplasm protein P9-1 of Rice black-streaked dwarf virus preferentially binds to single-stranded RNA in its octamer form, and the central interior structure formed by this octamer constitutes the major RNA binding site. *J. Virol.*, 87, 12885–12899.
69. Akita, F., Miyazaki, N., Hibino, H., Shimizu, T., Higashiura, A., Uehara-Ichiki, T., Sasaya, T., Tsukihara, T., Nakagawa, A., IwasaK, K. et al. (2011) Viroplasm matrix protein Pns9 from rice gall dwarf virus forms an octameric cylindrical structure. *J. Gen. Virol.*, 92, 2214–2221.