Importance of RNA length for in vitro encapsidation by the nucleoprotein of human respiratory syncytial virus

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Respiratory syncytial virus has a negative-sense single-stranded RNA genome constitutively encapsidated by the viral nucleoprotein N, forming a helical nucleocapsid which is the template for viral transcription and replication by the viral polymerase L. Recruitment of L onto the nucleocapsid depends on the viral phosphoprotein P, which is an essential L cofactor. A prerequisite for genome and antigenome encapsidation is the presence of the monomeric, RNA-free, neosynthesized N protein, named N⁰. Stabilization of N⁰ depends on the binding of the N-terminal residues of P to its surface, which prevents N oligomerization. However, the mechanism involved in the transition from N⁰-P to nucleocapsid assembly, and thus in the specificity of viral genome encapsidation, is still unknown. Furthermore, the specific role of N oligomerization and RNA in the morphogenesis of viral factories, where viral transcription and replication occur, have not been elucidated although the interaction between P and N complexed to RNA has been shown to be responsible for this process. Here, using a chimeric protein comprising N and the first 40 N-terminal residues of P, we succeeded in purifying a recombinant N⁰-like protein competent for RNA encapsidation in vitro. Our results showed the importance of RNA length for stable encapsidation and revealed that the nature of the 5' end of RNA does not explain the specificity of encapsidation. Finally, we showed that RNA encapsidation is crucial for the in vitro reconstitution of pseudo-viral factories. Together, our findings provide insight into respiratory syncytial virus viral genome encapsidation specificity.

The Mononegavirales order includes many human pathogenic viruses such as those responsible for rabies, measles, mumps, haemorrhagic fevers due to Ebola or Nipah viruses, and respiratory diseases induced by human respiratory syncytial virus (HRSV), metapneumovirus (HMPV), or parainfluenza viruses (1). All these viruses have a negative-sense single-stranded RNA genome that is constitutively encapsidated by the nucleoprotein N, forming the nucleocapsid (NC). This NC serves as a template for the viral polymerase L responsible for both viral transcription and replication (2, 3). For the vast majority of these viruses, transcription and replication are cytoplasmic, occurring in viral factories which are viro-induced organelles called inclusion bodies (IBs) (4–8).

Among Mononegavirales, HRSV is the prototype of the Pneumoviridae family and the Orthopneumovirus genus (9, 10). HRSV is the most common cause of acute lower respiratory infections in young children worldwide (11–13). It is also recognized as a major cause of severe respiratory infections in immunocompromised and elderly people (14–16). Currently, no vaccine is available, and the only specific treatment against HRSV is prophylactic, consisting of injection of humanized monoclonal antibodies directed against the fusion protein F (palivizumab, Sinagis) (17). However, the efficacy of this treatment is controversial and its high cost limits its prescription to at-risk children. The HRSV genome is approximatively 15.2 kb long and contains 10 genes encoding 11 proteins (2). Replication and transcription rely on 4 of these proteins: the RNA-dependent RNA polymerase L which exhibits all the enzymatic activities required for viral replication and transcription, its cofactor, the phosphoprotein P responsible for the recruitment of L on the NC, the nucleoprotein N involved in genome and antigenome encapsidation, and the transcription factor M2-1 that has been described as an “antiterminating” factor during transcription and interacts with P and viral mRNA.

During transcription, the polymerase L recognizes the leader sequence at the 3' end of the genome, and the polymerase is guided by the gene start (gs) and gene end (ge) sequences flanking each gene in the viral genome. The neosynthesized viral mRNAs are capped and methylated at the 5' end and polyadenylated at the 3' end by the polymerase. It is noteworthy that the synthesis of all viral mRNAs depends on the presence of the M2-1 protein, which is required for L processivity along the genome (18, 19). The mechanism of action of M2-1 is still poorly understood. However, it has recently been shown that RSV mRNAs concentrate with M2-1 in specific subcompartments of IBs called IBs-associated granules (8), suggesting that the M2-1–mRNA interaction may promote the release of mRNAs from the polymerase L.

On the contrary, replication of the viral genome only depends on L, P, and the NC. In that case, the polymerase
recognizes the leader or the trailer sequences at the 3′ end of the genome or antigenome respectively, and the RNA synthesis starts at the first nucleotide and the polymerase proceeds through the ge and gs sequences to the end of the genome (or antigenome) (20, 21). Genomes and antigenomes, which have a triphosphate 5′ end (22, 23), are directly encapsidated by the N protein during their synthesis, forming helical NC (24) which can both be incorporated in progeny virions (25, 26). Encapsulation of genome and antigenome thus prevents the formation of double strand RNAs but also their degradation as well as their detection by cellular sensors of the innate immune system. The structure of N expressed in *Escherichia coli* and purified as N-RNA rings composed of 10 protomers revealed that each N protomer interacts with 7 nucleotides (27). The RNA binds within a groove formed by the interface between the 2 globular domains of N (NNTD and NCTD) that are separated by a flexible hinge region. The N protein also possesses 2 N- and C-terminal extensions (N- and C-arms) that are involved in N oligomerization: the N-arm of the Ni protomer binds to the N01 protomer, whereas the C-arm of N0 binds to the top of the NCTD of the Ni+1 protomer (24, 27). RNA encapsidation therefore depends on both direct interaction with RNA and the ability of N to oligomerize, 2 mechanisms that are closely coupled. When expressed alone, the *Mononegavirales* N proteins all show a strong tendency to encapsidate cellular RNAs. This implies that viral replication depends on the ability to maintain a pool of RNA-free monomeric N, termed N0, available for the specific encapsidation of genomic and antigenomic RNAs. It is now well established that all these viruses share a common mechanism, with the neosynthesized N being maintained in the N0 form by P which acts as a molecular chaperone (28–36). Specifically for HRSV, and by analogy to the N0–P complex of the HMPV whose crystal structure has been solved (31), the binding of the 28 N-terminal residues of P to the NCTD surface overlaps with the binding sites of both the N- and C-arms of the Ni+1 and N01 protomers in the oligomeric form, preventing self-oligomerization of N (30, 37). Notably, N0 is characterized by a rotation of NNTD relative to NCTD compared to the oligomeric form and a stacking of the C-terminal arm of N into the positively charged RNA groove that blocks RNA binding (30). However, the mechanism involved in the switch from N0 to N-RNA, which regulates specific encapsidation of genomes and antigenomes remains poorly characterized.

In addition to their critical role in viral polymerase function, N and P have been shown to be the scaffold proteins responsible for IBs morphogenesis (38, 39). It has recently been shown that IBs are liquid organelles formed by liquid-liquid phase separation induced by N–P interaction (39, 40). Importantly, the P protein (241 residues long), which forms tetramers through its central oligomerization domain (residues 131–151) flanked by highly disordered N- and C-terminal domains (PNTD and PCTD respectively) that are involved in multiple protein–protein interactions, plays a central role in regulating polymerase activity as well as in IBs formation. The PNTD interacts with N0 (37), M2-1 (41), the cellular phosphatase PP1 (41), and the viral matrix protein M responsible for virion assembly (42), whereas PCTD is involved in the interaction with L (43, 44) and NC (45, 46) and is critical for IBs formation (39). Both the length of the PCTD and its interaction with N complexed to RNA are required for efficient IBs morphogenesis. However, the importance of N oligomerization and of RNAs in the morphogenesis of IBs remains to be characterized.

In this study, we investigated the specificity of RNA encapsidation by HRSV N. Using a fusion protein between the full-length N and the peptide derived from the first 40 N-terminal residues of P (P40) responsible for chaperone activity on N0, we managed to purify a recombinant monomeric and RNA-free N-P40 chimeric protein. The resulting N-P40 protein was competent for *in vitro* RNAs encapsidation, the length of RNAs being critical for stable oligomerization, and invariably leading to the formation of N-RNA rings under the conditions tested. Furthermore, using this N-P40 protein, we revealed that RNA encapsidation is critical for the morphogenesis of pseudo-IBs *in vitro.*

**Results**

**Production of a N0-like recombinant protein**

Recently, a strategy based on coexpression of PNTD with His-tagged N in *E. coli* allowed the purification of a N0–PNTD complex competent for encapsidation (47). Given the low affinity of P peptide for N (37), which could lead to the loss of peptide upon purification and thus to N aggregation, we chose here to use a strategy based on the expression of a fusion protein between His-tagged full-length N and the 40 N-terminal residues of P (P40) in *E. coli* (Fig. 1A). This approach was previously used to determine the crystal structure of the N0–P complexes of HMPV (31), measles virus (34), and parainfluenza virus 5 (32). After purification of this chimeric protein in the presence of 1 M NaCl, the sample was analyzed by size-exclusion chromatography, following the absorbance at 260 and 280 nm, and compared to the profile of purified WT N (Nwt) forming N-RNA rings (46). The elution profile of N-P40 showed 2 peaks (Fig. 1B). The major P1 peak with A260nm/A280nm ratio >1 and apparent mass of ~500 kDa (estimated from the Superdex 200 calibration profile), of slightly smaller size than N-RNA rings, that could correspond to N-RNA oligomers or aggregates. A minor P2 peak presented A260nm/A280nm ratio <1 and apparent mass of ~50 kDa, as expected for monomeric RNA-free N-P40 protein. The fractions of P1 or P2 peaks were pooled, and samples were analyzed by SDS-PAGE stained with Coomassie blue and by band shift assay on native agarose gel and compared to the sample corresponding to purified Nwt. As shown on Figure 1C (upper panel), as analyzed by SDS-PAGE, the presence of a single band with a molecular weight close to 45 kDa for Nwt sample and of 50 kDa for P1 and P2 peaks were observed, consistent with the expected mass of the Nwt and the N-P40 protein, respectively. On native agarose gel, the Nwt complexed to RNA and thus negatively charged clearly migrated within the gel. The N-P40 from the P1 peak migrated with a single band, close to the band observed for Nwt, suggesting that N-P40 could
oligomerize as rings (Fig. 1, lower panel). On the contrary, N-P40 from the P2 peak poorly migrated within the gel and in the opposite direction.

We then focused on the sample from P2 peak which was dialyzed to 300 mM NaCl and concentrated up to 1.5 mg/ml. Dynamic light scattering (DLS) analysis of the sample revealed the presence of a major protein peak near 6.7 nm in diameter that should correspond to monomers and a minor peak of oligomers near 17 nm in diameter (Fig. 1D). These data revealed that in our conditions, N-P40 tended to aggregate or oligomerize upon concentration and led us to work with a protein concentration of 1 mg/ml. We finally analyzed the secondary and tertiary structures of N-P40 by circular dichroism (CD), using as previously Nwt purified as N-RNA rings as a control. The far-UV spectra of both N-P40 and Nwt showed peaks at 208 and 222 nm, typical of secondary structures with predominant α-helical content (Fig. 1E). The peak intensity shift observed for N-P40 revealed a lower helical content than Nwt. This observation correlates with the insertion of linkers in the N-P40 constructs, which are expected to

Figure 1. Purification and characterization of recombinant N-P40 fusion protein. A, amino acid sequence of the N-P40 fusion protein. The N-terminal 6xHis tag is in green, the N sequence is in blue, and the sequence of P40 is in red. Amino acids in black correspond to additional residues and linkers. B, gel filtration profile of purified N-P40 (red) and Nwt (black). The curves corresponding to absorbance at 260 nm and 280 nm are presented as dash and solid lines, respectively. P1 and P2 indicate the 2 peaks detected for the gel filtration profile of N-P40. The asterisk indicates the peak corresponding to the C-terminal fragment of P used for Nwt purification (46). C, the fractions corresponding to the peak of purified Nwt and to the peaks P1 and P2 of the gel filtration profile of N-P40 were pooled, and the samples were analyzed by SDS-PAGE colored with Coomassie blue (upper panel) and by migration on native agarose gel (lower panel). D, dynamic light scattering (DLS) analysis of the N-P40 protein isolated from the gel filtration peak P2, showing a homogenous peak close to 6.7 nm, corresponding to N-P40 monomer and a minor peak of oligomers. E and F, far-UV (E) and near-UV CD (F) spectra of Nwt corresponding to N-RNA rings in black and N-P40 in gray.
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be unfolded. The near-UV CD allows the detection of signals from aromatic residues engaged in a rigid chiral environment, consistent with the presence of a tertiary structure. The detection of a broad signal around 280 nm for the N-P40 protein can be attributed to the signal from Tyr and Trp residues and validates that the protein has a tertiary structure (Fig. 1F). In comparison, the spectrum obtained for N wt revealed an intense positive signal between 260 and 280 nm, which can be attributed to the presence of RNA masking the signals from the aromatic residues of N.

Overall, these results show that N-P40 can be purified as a monomeric and RNA-free protein and that the fusion of P40 does not have a major impact on N folding. This latter observation correlates with previous results showing that binding of P40 to N does not induce major conformational changes in N (30).

N-P40 is competent for encapsidation of RNAs

We then assessed the capacity of the N-P40 protein to encapsidate RNA in vitro. As each N wt protomer has been shown to interact with seven nucleotides within purified N rings formed of 10 or 11 protomers (27), we first tested the ability of N-P40 to encapsidate a 70-nucleotide long (70-mer) RNA or DNA of similar sequence. The RNA sequence that we used was the antileader sequence (the 5′ end of the antileader sequence of HRSV, which is the initial RNA sequence synthesized by the polymerase using the genomic RNA as a template) (Table 1). Encapsidation of RNA by N-P40 was analyzed using a band shift assay on native polyacrylamide and agarose gels, using N wt rings as a control. As shown in Figure 2A, while only slight migration was observed for purified N-P40, a clear band shift was observed when N-P40 was incubated with RNA. However, the protein seemed to poorly interact with DNA. In parallel, the samples were observed by negative stain electron microscopy (ns-EM). As expected, images of N-P40 alone showed only very small species (Fig. 2C). While N-P40 incubated with 70-mer RNA formed ring-shaped N-RNA assemblies, only aggregates were observed for the sample of N-P40 incubated with DNA (Fig. 2C). These results suggest that, although N could interact with DNA, the affinity is not sufficient to allow the regular and constrained assembly of N on nucleotides, which, based on the crystal structure, implies twists on the ribose-phosphate chain (27). Using the same approach, we confirmed that N-P40 was also capable of encapsidating shorter 14-mer RNAs and forming N-RNA rings (Fig. 2, B and C). Furthermore, as the first nucleotides could initiate RNA encapsidation, we investigated the impact of substitutions of the 2′ first nucleotides on the capacity of N to encapsidate 14-mer RNAs (Table 1). However, we observed that N-P40 similarly encapsidated 14-mer RNAs of different sequences (Fig. 2B), suggesting that the first nucleotides are not critical for starting RNA encapsidation.

Our results thus show that, although fused to N, the P40 peptide is not sufficient to prevent RNA encapsidation in vitro. Furthermore, while N-P40 specifically encapsidates RNAs over DNA, no specificity in RNA length was observed under the conditions tested. It is noteworthy that, as previously observed by Gao et al. (47), in vitro encapsidation of RNA did not result in the formation of helical NCs but only in N-RNA rings.

Investigation of the minimal RNA length required for encapsidation

Using the same antileader sequence of HRSV, we further studied the minimal RNA length required for encapsidation, using 5- to 11-mer oligonucleotides (Table 1). As shown in Figure 3A, no band shift of N-P40 was detected on native polyacrylamide or agarose gels in the presence of 5- and 6-mer RNAs sequences

| RNAs length | 5′ end modification | RNA sequence* |
|-------------|---------------------|---------------|
| 70-mer      | OH                  | 5′-ACGCAGAAAAAUGCGUAACAAACACUUGCAUAACCCAAAAGAAUGGGCAGAAUAAGAAAUUGGAAGUAAGAC-3′ |
| 14-mer      |                     | 5′-ACGCGAAAAAUGGC-3′ |
| 14-mer (a)  |                     | 5′-GGGCGAAAAAUGGC-3′ |
| 14-mer (b)  |                     | 5′-GGGCGAAAAAUGGC-3′ |
| 14-mer (c)  |                     | 5′-GGGCGAAAAAUGGC-3′ |
| 11-mer      |                     | 5′-ACGCGAAAAA-3′ |
| 10-mer      |                     | 5′-ACGCGAAAAA-3′ |
| 9-mer       |                     | 5′-ACGCGAAAAA-3′ |
| 8-mer       |                     | 5′-ACGCGAAA-3′ |
| 7-mer       |                     | 5′-ACGCGA-3′ |
| 6-mer       |                     | 5′-ACGCGA-3′ |
| 5-mer       |                     | 5′-ACGCGA-3′ |
| 11-mer      | ppp                 | 5′-ACGCGAAAAAAA-3′ |
| 10-mer      |                     | 5′-ACGCGAAAAAAA-3′ |
| 9-mer       |                     | 5′-ACGCGAAAAAAA-3′ |
| 8-mer       |                     | 5′-ACGCGAAAAAAA-3′ |
| 7-mer       | 7′OHp               | 5′-ACGCGAAAAA-3′ |
| 5-mer       |                     | 5′-ACGCGAAA-3′ |
| 4-mer       |                     | 5′-ACGCGAA-3′ |

* All the sequences correspond to HRSV antileader sequence except those of the 14-mer b and c indicated in italics.

*b Corresponding 70-mer DNA was used in the study.
RNAs. Similar N-P40 band shifts were observed in the presence of 10- and 11-mer oligonucleotides compared with N-P40 incubated with 14-mer RNAs. Finally, intermediate band shift profiles were observed for N-P40 incubated in the presence of 7-, 8-, and 9-mer RNAs. Yet, according to ns-EM observations, incubation of N-P40 with 7-mer and 11-mer RNAs both led to the formation of N-RNA rings, similar to the ones observed with the 14-mer and 70-mer RNAs (Fig. 3B).

To clarify these results, we next investigated the stability of the N-P40–RNA complexes by incubating the samples overnight in the presence of RNase A followed by dialysis. Migration analysis of the samples on a native agarose gel showed no band shift for N-P40 incubated with 6- to 10-mer RNAs, suggesting RNA digestion, whereas a band shift was still observed for N-P40 in the presence of 11-mer and 14-mer RNAs (Fig. 3C), indicating that RNA was protected from digestion. Further analysis of the samples was performed by measuring the $A_{260nm}/A_{280nm}$ ratio and by investigating structural changes of N-P40 by DLS and near-UV CD. Before RNase A treatment, all samples displayed an $A_{260nm}/A_{280nm}$ ratio close to 1.2, similar to the value measured for Nwt (Table 2). After RNase A treatment, $A_{260nm}/A_{280nm}$ ratios were <1 for N-P40 samples incubated with 6- to 10-mer RNAs (Table 2), validating RNA digestion upon RNase A treatment. Of note, the $A_{260nm}/A_{280nm}$ ratio of these samples remained higher than that obtained for purified N-P40. For N-P40 incubated in the presence of 11- and 14-mer RNAs, the $A_{260nm}/A_{280nm}$ ratios were close to 1. These data suggested that, although the RNAs were partially digested by RNase A, N-P40 was still associated with RNAs, preventing complete RNA degradation. In parallel, determination of the hydrodynamic diameter (Dh) of the samples by DLS revealed that N-P40 incubated in the presence of 6- to 10-mer RNAs displayed a Dh between 6.7 and 9 nm after RNase treatment, close to the Dh obtained under the same conditions for N-P40 alone, whereas those of N-P40 incubated with 11- and 14-mer RNAs were 14.2 and 14.4 nm, respectively, similar to the Dh of Nwt (Table 2). Furthermore, the near-UV CD spectra showed that only the N-P40 sample incubated with 11-mer RNA exhibited a positive peak around 260 nm, whereas the spectra with 7- to 10-mer RNAs were similar to purified N-P40 alone or displayed only a weak signal around 260 nm (Fig. 3D). These data correlate with absorbance and Dh measurements, suggesting that RNase A treatment induced degradation of RNAs shorter than 11-nucleotides long and N rings disassembly.

Altogether, these results reveal that stable RNAs encapsidation by N-P40 depends on RNA length, with the minimal length to stabilize N-P40–RNA complexes being 11-mer RNAs.

**Impact of 5′ end modification of RNAs on encapsidation**

One of the differences between RNAs lies in post-transcriptional modifications, notably at the 5′ end. Genomic and antigenomic RNAs have been described as having a 5′ triphosphate (ppp) (22, 23), whereas viral mRNAs are capped and methylated by L during their synthesis (3). Therefore, we investigated whether posttranscriptional modifications of the 5′ end of RNAs could impact the ability of N to encapsidate RNAs. Again, N-P40 was incubated in the presence of synthetic 7- to 11-mer RNAs with either a 5′ ppp or a N7 and 2′-O methylated cap structure (7mGppp) (Table 1), followed by native agarose gel electrophoresis.
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Results showed that the presence of ppp or 7mGppp at the 5' end inhibited the ability of N to encapsidate 7- and 8-mer RNAs compared to 5'OH RNAs (Figs. 3A and 4A). However, N-P40 was still able to encapsidate 9-, 10-, and 11-mer RNAs, and no difference was observed between 5' end ppp or capped RNAs. Interestingly, independently of the length and the capping of the RNAs, N-P40–RNA rings could always be detected by ns-EM (Fig. 4B), suggesting that this method of observation may stabilize ring-shaped assemblies and highlights the importance of using multiple complementary techniques for the analysis of RNA encapsidation by N.

One more time, we investigated the impact of RNAs’ 5’ end modification on the stability of N-P40–11-mer RNA complexes upon RNase A treatment. Surprisingly, although band shifts were still observed (Fig. 4C), N-P40 samples incubated with 5’ppp and 5’7mGppp RNAs displayed A260nm/A280nm ratios of 0.9 and 1, respectively, and DLS analysis revealed the presence of different populations (not shown). Furthermore, while the near-UV CD spectrum of N-P40 incubated with 5’7mGppp RNAs was similar to the spectrum of N-P40 complexed with 5’OH RNA, the spectrum of N-P40 incubated with 5’ppp RNAs clearly differed from those of N-P40 alone or incubated with 5’OH RNAs (Fig. 4D).
These results revealed that modification of the 5’ end of RNAs only partially impairs encapsidation but is not a critical factor for the specificity of encapsidation, 5’ end modifications delay RNAs encapsidation maybe by creating steric hindrance.

**Impact of full-length P protein on the stability of RNAs encapsidation**

As our results were obtained in the presence of only a short peptide of P, we then assessed if full-length P protein could modulate the stability of RNAs encapsidation. By analysis on native agarose gel, we first showed that coincubation of P with N-P40 allowed to observe the formation of a complex, revealing that P was able to interact with N-P40, although this interaction appeared weak compared to the interaction between P and N wt (Fig. 5A). This result suggests that P could bind to N either by displacing P40 binding on N or through the binding of the C-terminal extremity of P on NNTD. In the presence of 14-mer RNA, a band of strong intensity, similar to the band observed in presence of Nwt, was observed, showing that P does not prevent RNA encapsidation by N-P40. We therefore studied the impact of P on encapsidation depending on RNA length. As previously, no band shift was detected in the presence of 5- and 6-mers. In the presence of 7- to 9-mers, a band of weak intensity migrating slightly faster and at the same level than the bands of strong intensity observed in the presence of 10-, 11-, and 14-mers was observed (Fig. 5B, upper panel). These data reveal that in the presence of RNA, the full-length P does not improve the stability of the monomeric N compared to the P40 peptide. Of note, upon treatment with RNase A, the band observed in the presence of 8- and 9-mers was not detected, and a clear decrease of intensity of the band observed in the presence of 10-mer RNA was observed (Fig. 5B, lower panel). Finally, in the presence of 5’ end ppp or 7mGppp RNAs, a band of strong intensity was observed only in the presence of 11-mer RNAs.

Altogether, these results reveal that the presence of P does not drastically impact RNA encapsidation or the stability of the oligomers.

**Role of RNA encapsidation on pseudo-IBs morphogenesis in vitro**

We have previously shown that in vitro coincubation of purified recombinant fluorescent N-RNA rings (mCherry-Nwt) with P can lead to the formation of liquid droplets that exhibit properties similar to cellular IBs formed during infection. We also identified the key properties of P required for pseudo-IBs morphogenesis, that is, oligomerization, PCTD length and flexibility, and interaction of the C-terminus of P with N (39). However, this study did not decipher the role of N oligomerization and of the presence of RNAs in this process. Furthermore, although our data suggested that the N/P ratio maybe important for optimal IBs formation, these results remained preliminary, and a systematic study of the impact of N/P ratio and concentration on the morphogenesis of IBs was missing. As phase separation is known to be facilitated by increased protein concentration (48–52), we here decided to limit the protein concentrations to avoid spontaneous pseudo-IBs formation. Using a fixed minimum concentration of 3.5 μM for mCherry-Nwt or P, the addition of the second partner at a 1:1 ratio allowed us to observe the formation of small droplets of 1 to 5 μm in diameter (Fig. 6). A fixed low P concentration limited the size and number of droplets that remained similar whatever the concentration of mCherry-N (Fig. 6A). On the other hand, in a fixed low N concentration condition, the number and size of droplets increased proportionally to the addition of P until reaching a plateau for a P/N ratio ≥4 (Fig. 6B). These results correlate with our previous observations (39) but also revealed that P concentration is a limiting factor for IBs formation.

To study the impact of N-P40 oligomerization in the presence of RNAs on pseudo-IBs morphogenesis, we then investigated whether pseudo-IBs could form when N-P40 was incubated with a 70-mer RNA in the presence of P-BFP. Since no droplets were observed with a 3 μM concentration of N-P40 in the presence of an excess of P-BFP (15 μM) and 70-mer RNA at 50 μM, we then screened for the optimal conditions allowing pseudo-IBs formation, with a minimum concentration of proteins. Pseudo-IBs of 10 to 15 μm in diameter were observed in the presence of 3.5 μM of P-BFP, 70-mer RNAs (50 μM), and a minimum of 4-fold excess of N-P40 (13.5 μM). Of note, pseudo-IBs of only 2 to 3 μm in diameter were observed when coincubating P-BFP with Nwt (RNA-N rings) in the same conditions (Fig. 7A). It is noteworthy that no droplets were detected when incubating P-BFP in the presence of either RNAs or N-P40 alone (Fig. 7A). We then assessed the impact of RNA length on pseudo-IBs morphogenesis by incubating N-P40 with RNAs of various lengths before the addition of P-BFP, under the same conditions. As shown in Figure 7B, the addition of 14-mer RNA resulted in the formation of droplets similar to those observed in the presence of 70-mer RNA. More interestingly, droplet size decreased when N-P40 was incubated with 11-mer and 8-mer RNAs, and only aggregates were observed in the presence of 6-mer RNA. Furthermore, whereas RNase A treatment of N-P40 incubated with 11-mer RNA still allowed to observe droplets, only aggregates were detected for the sample with 8-mer RNA (Fig. 7C). This result correlates with data showing the instability of N oligomerization in the presence of short RNAs. Finally, pseudo-IBs similar to those obtained in the presence of RNA encapsidation by the RSV nucleoprotein

**Table 2**

| Proteins and RNAs | A260nm/A280nm | Dh (nm) |
|-------------------|---------------|---------|
| Nwt               | 1.24          | 15.75   |
| N-P40             | 0.67          | 7.9     |
| N-P40 + 6-mer RNA | 0.79          | 8.9     |
| N-P40 + 7-mer RNA | 0.9           | 8.7     |
| N-P40 + 8-mer RNA | 0.88          | 6.7     |
| N-P40 + 9-mer RNA | 0.87          | 7.6     |
| N-P40 + 10-mer RNA| 0.94          | 9       |
| N-P40 + 11-mer RNA| 1.03          | 14.2    |
| N-P40 + 14-mer RNA| 1            | 14.4    |

* a Hydrodynamic diameter determined by DLS. Data representative of 2 independent experiments.

* b Purified as N-RNA rings.
Figure 4. Impact of RNA 5′ end modifications on the ability of N-P40 to encapsidate RNAs. A, analysis of N-P40 migration alone or incubated in the presence of 7- to 11-mer RNAs with 5′ ppp (left) or 5′7mGppp (right) by native agarose gel electrophoresis. Gels were stained with amido black. B, images of N-P40–RNA complexes formed upon incubation of N-P40 in the presence of 7- and 11-mer RNAs with 5′ ppp or 5′7mGppp, as observed by ns-EM. The scale bar represents 50 nm. Magnifications of selected areas (indicated by black squares) are presented at the right top of each panel. The scale bar represents 20 nm. C, comparison of the migration of N-P40 alone or incubated in the presence of 11-mer RNAs with 5′ OH, ppp, or 7mGppp after RNase A treatment by native agarose gel electrophoresis, gel stained with amido black. D, near-UV CD spectra of N-P40 alone or incubated in the presence of 11-mer RNAs with 5′ OH, ppp, or 7mGppp after RNase A treatment. ns-EM, negative stain electron microscopy.
Figure 5. Impact of full-length P on the stability of RNA encapsidation. A, migration on native agarose gel of P, Nwt, or N-P40 in the absence or presence of 14-mer RNA alone or coincubated in the absence or the presence of 14-mer RNA. The asterisk indicates the band corresponding to P–N-P40 complex migration. B and C, analysis by native agarose gel electrophoresis of P and N-P40 migration alone or incubated in the presence of 7- to 11-mer RNAs (B) before (upper panel) or after treatment (lower panel) by RNase A or with 9- to 11-mer RNAs displaying 5’ ppp or 7mGppp (C). The black and white arrows indicate the bands corresponding to the migration of P–N-P40 and P–N-P40–RNA complexes, respectively. The asterisk indicates the band corresponding to RNase A.

Figure 6. Impact of Nwt and P concentration and ratio on pseudo-IBs formation. Recombinant mCherry-Nwt (corresponding to mCherry-N-RNA rings) and P proteins were coincubated in the presence of 15% Ficoll and the formation of droplets was observed by fluorescence microscopy. A, P concentration was fixed to 3.5 μM, and mCherry-Nwt was added up to a ratio mCherry-Nwt/P of 5; B, mCherry-Nwt concentration was fixed to 3.5 μM, and P concentration increased up to a ratio P/mCherry-Nwt of 5. The scale bar represents 10 μm. IB, inclusion body.
11-mer RNA with 5’ OH were observed in the presence of 11-mer RNAs with 5’ ppp and 7mGppp (Fig. 7D).

Altogether, our data revealed that coincubation of P with monomeric N does not allow pseudo-IBs morphogenesis in vitro, and that the addition of short RNAs is not sufficient to induce phase separation. Overall, these results showed that N oligomerization in the presence of RNA is critical for the morphogenesis of pseudo-IBs.

Discussion

Like for the majority of viruses belonging to the Mononegavirales order, HRSV replication and transcription take place within cytoplasmic viral factories called IBs that concentrate the viral and cellular proteins required for these activities, as well as neosynthesized antigenomic and genomic RNAs and viral mRNAs. Replication leads to the synthesis and amplification of full-length negative-sense genomic and positive sense antigenomic RNAs, which are encapsidated by N in its RNA-free form N0, forming the so-called NCs, in contrast to viral mRNAs. This specificity of encapsidation of viral genomes and antigenomes by N remains unexplained. Given the strong tendency of N0 to oligomerize on RNAs, some particular mechanisms might regulate the specificity of viral RNA encapsidation and thus the transition from N0 to N-RNA. The aim of this work was to investigate whether specificity of encapsidation could be explained either by the sequence or the nature of the 5’ end of RNAs. Here, using a chimeric protein
composed of full-length N and P40, a peptide corresponding to
the 40 N-terminal residues of P, we showed that this construct
is competent for RNA encapsidation in vitro, even in the
presence of full-length P protein. This first observation shows
that P is not sufficient to prevent RNA encapsidation by N,
suggesting the existence of another factor that could regulate
the transition from monomeric N° N-RNA assembly in
infected cells, most likely by enhancing the stability of N°. In
agreement with a recent publication by Gao et al. (47), our
results also confirmed that RNA as short as 7-mer can be
encapsidated, and that N oligomerization on RNAs always
leads to the formation of N-RNA rings in vitro. We showed
that the minimum length for stable encapsidation was 11
nucleotides, revealing that RNA length is critical for the sta-
bility of N-RNA oligomers. However, we cannot exclude that
the presence of P40 fused to the C-terminus of N may partially
alter N oligomerization. In contrast to measles virus for which
the presence of P40 fused to the C-terminus of N may partially
modulate IBs (39). When P was mixed with a monomeric
protein (N-P40) and RNAs are too short to be
encapsidated, no phase separation was observed, indicating
that the presence of RNA-N oligomers is critical for pseudo-
IBs formation. Interestingly, the length of encapsidated
RNAs seems to modulate the size of pseudo-IBs. These ob-
servations correlate with the fact that RNAs have been shown
to play a major role in liquid-liquid phase separation mecha-
nisms (49, 50). Further characterization of IBs morphogenesis
in the presence of RSV helical NC should be of great interest,
as the size and high flexibility of these structures could
modulate IBs morphogenesis. A recent cryo-electron tomog-
raphy study of RSV-infected cells has revealed the presence of
N rings in filamentous RSV particles (57), showing for the first
time the heterogeneity of N assembly in the cell. The role of
these structures during viral cycle, that is, IBs morphogenesis,
control of the innate immune response to infection, and virion
assembly remains to be determined.

Experimental procedures

Plasmid constructs

The gene sequence of the nucleoprotein N was amplified
without stop codon by PCR using Pfu DNA polymerase
(Statagene) and cloned in the pET28a+ vector, at BamHI
and XhoI sites. The oligonucleotides used were as follows:
5'-GCCGCGGATCCATGCTTATTAGAAAAGTCAAGTTG-3' (BamHI)
and 5'-GAGGAGCTCGAGAAGCTTATCTTTTGAGTTCTCG-3' (XhoI). The sequence-coding
TEV cleavage site was then introduced at the BamHI site af-
after annealing of the oligonucleotides 5'-GATCCGAACCT
ATTATCCAGG-3' and 5'-GATCCCTGGAATATAGGT
TTCTCG-3'. Finally, the sequence coding for the 40 N-ter-
nimal residues of P was PCR amplified using the oligonucleo-
tides 5'-GAGGAGCTCGAGGATGGCGGTAGCGGTAG
CGTACGTAGAAAATGGTTGC TCTCGAATTTCC-3' (containing the sequence coding for 4 Gly/Ser, underlined)
and 5'-GAGGAGCTCGAGGATATAGGT
TTCTCG-3' (with a stop codon at the 3' end, underlined)
and inserted at the XhoI site. The viral sequences derived from
the human RSV strain ATCC VR-26 (Genbank accession
number:AY911262.1). The final plasmid allows the expression
of the fusion protein 6xHis-N-4xgs-P40 containing a TEV
cleavage site upstream of the N sequence. For expression and
purification of recombinant mCherry-N, N, and P-BFP pro-
teins, the previously described pET-mCherry-N, pET-N, and
pET-P-BFP plasmids were used (37, 39).

Expression and purification of the recombinant proteins

The E. coli BL21 (DE3) bacteria strain (Novagen)
were transformed by the pET-N-P40 plasmid. Cultures were
grown at 37 °C in 2xYT medium containing 50 µg/ml kanamycin.
After 8 h, an equal volume of 2xYT medium containing
50 µg/ml kanamycin was added to the cultures, and the
expression was induced by the addition of 80 µg/ml IPTG
overnight at 28 °C. Bacteria were then harvested by centrifu-
gation. Pellets were resuspended in lysis buffer (20 mM
Tris–HCl, pH 8, 500 mM NaCl, 0.1% Triton X-100, 10 mM
imidazole, 4 mM benzamidine, 1 mg/ml lysozyme, and com-
plete protease inhibitor cocktail (Roche)). After incubation on
ice for 30 min, the lysates were sonicated, benzonase (Novagen) (final concentration 5 U/ml) and RNase A (final concentration of 1 U/ml) were added to the lysate, followed by incubation for 30 min at room temperature, finally NaCl was added (up to a concentration of 1 M). After centrifugation at 10,000 g for 30 min at 4 °C, the lysates were incubated with chelating Sepharose Fast Flow beads (GE Healthcare) charged with Ni²⁺, for 1 h at 4 °C. Beads were washed with washing buffers (20 mM Tris–HCl, pH 8, 1 M NaCl, 4 mM benzamidine) with increasing imidazole concentrations (10, 50, and 100 mM), before elution of the protein using 600 mM imidazole. Purified proteins were then loaded on a Hi-Load 16/600 Superdex 200 column (GE Healthcare) and eluted in 20 mM Tris–HCl, pH 8.5, 1 M NaCl. Then, the purified protein was dialyzed against 10 mM Tris–HCl, pH 8.5, with decreasing NaCl concentrations (500 and 300 mM) buffers, for 4 h at each step, at 4 °C. Finally, the purified protein was concentrated using a centrifugal concentrator with a MWCO of 10 kDa (vivaspin turbo 4, Sartorius). WT mCherry-N, N, and P-BFP proteins were expressed and purified as previously described (39).

**Synthesis of RNA substrates**

RNA sequences were chemically synthesized on a solid support using an ABI 394 automated synthesizer with 2'-O-pivaloyloxymethyl 3'-O-phosphoramidite ribonucleosides and with 2'-O-methyl 3'-O-phosphoramidite adenosine to obtain 7mGpppA₉₉RNA (ChemGenes Corp.) (58). After RNA assembly, depending on the desired 5' end of the RNA, 3 different processes were applied. For 5' OH-RNA sequences, the solid support was directly subjected to a basic treatment (28% aqueous ammonia solution at r.t for 3 h) to recover the crude RNA material. In the other 2 cases, the 5'-hydroxyl group was phosphorylated and the resulting H-phosphonate derivative was oxidized and activated to a phosphorimidazolidate derivative to react with pyrophosphate or GDP, yielding ppp-RNA (59) or Gppp-RNA (60), respectively. After deprotection and release from the solid support by the same ammonia treatment as for 5'-OH RNA, all RNA substrates were purified by IEX-HPLC (>95% pure) and their identity was confirmed by MALDI-TOF spectrometry. N7-methylation of the cap structure (Gppp) was performed using human (guanine-N7)-MTase to obtain 7mGppp-RNA (60).

**RNA–N-P40 complexes formation**

RNA oligonucleotides (15 μM) and purified N-P40 protein (10 μM) in 20 mM Tris–HCl, pH 8.5, 300 mM NaCl buffer were coincubated at different P/N molecular ratio on glass slides, and the molecular-crowding agent Ficoll was

**Band shift on native polyacrylamide and agarose gels**

50% sucrose loading buffer was added to the samples before loading either on native 4% native polyacrylamide gel or on native 1% agarose gel. For polyacrylamide native gel analysis, migration was performed in 0.2 x TBE (pH 8.0 for 2 h at 200 V at 4 °C), and gels were stained with Coomassie blue. For agarose native gel analysis, migration was performed in 1 × Tris–Glycine buffer during 1 h 30 at 80 V before staining with amido black 10B.

**Dynamic light scattering**

Size measurement of purified N-P40 alone or incubated with RNAs was performed at 20 °C using a helium-neon laser wavelength of 633 nm and detection angle of 173° with a Zetasizer Nano (Malvern). Ten measurements were made, with an acquisition time of 10 s for each measurement. Hydrodynamic diameters were calculated using the Zetasizer software provided by the instrument manufacturer (https://www.malvernpanalytical.com/fr/support/product-support/software/Zetasizer-Nano-software-update-v3-30). The results were presented as size distribution (nm).

**CD spectroscopy**

CD experiments were performed on a J-810 spectropolarimeter (Jasco) in a thermostated cell holder at 20 °C. Spectra of N-P40 and RNA–N-P40 complexes were measured after dialysis against 10 mM NaP, 300 mM NaF, pH 8.5 buffer, and at concentrations of 10 μM. Far-UV spectra (190–260 nm) were recorded in a 0.5 mm path-length quartz cell using a bandwidth of 2 nm and an integration time of 1 s. Near-UV spectra (260–320 nm) were recorded in a 10 mm path-length quartz cell using a bandwidth of 2 nm and an integration time of 1 s. Each spectrum was the average of 3 scans, with a scan rate of 100 nm/min and 50 nm/min for far-UV and near-UV spectra, respectively. Correction by subtracting the signal from the buffer was made and the spectra were smoothed with the fast Fourier transform filter (Jasco Software), and data were treated as previously described (37).

**Negative stain electron microscopy observations of N-P40–RNA complexes**

Three microliters of sample were applied to the clean side of a carbon–mica interface and stained with 2% sodium silicotungstate. Micrographs were recorded on a Tecnai T12 microscope operated at 120 kV with a Gatan Orius 1000 camera. Images were recorded at a nominal magnification of 23000× resulting in a pixel size of 2.8 Å.

**In vitro assay of pseudo-IBs formation**

As previously described (39), mCherry-N and P proteins or P-BFP and WT N or N-P40 recombinant proteins alone or in the presence of RNAs in 20 mM Tris–HCl pH 8.5, 150 mM NaCl buffer were coincubated at different P/N molecular ratio on glass slides, and the molecular-crowding agent Ficoll was
added on the droplets of solution. Then, coverslips were laid on the droplets. For RNase treatment, N-P40 recombinant protein was preincubated in the presence of 8-mer and 11-mer RNAs, then incubated half an hour at room temperature in the presence of RNase A (PureLink, Invitrogen). Pseudo-IBs were then observed with a Nikon TE200 inverted microscope equipped with a Photometrics CoolSNAP ES2 camera. Images were processed using MetaVue software (Molecular Devices, https://www.nikonusa.com/fileuploads/pdfs/MetaVue.pdf) and ImageJ software (https://imagej.nih.gov/ij/index.html).

Data availability
All data are contained within the article.

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Conflict of interest—The authors declare that they have no conflicts of interest with the contents of this article.

Abbreviations—The abbreviations used are: DLS, dynamic light scattering; HMPV, human metapneumovirus; HRSV, human respiratory syncytial virus; IB, inclusion body; NC, nucleocapsid; ns-EM, negative stain electron microscopy.

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