1	terization of the interaction domains between the phosphoprotein and the nucleocapsid		
2	of human Metapneumovirus		
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4	Running title: Metapneumovirus N-P interaction		
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#### 20 ABSTRACT

21 Human metapneumovirus (HMPV) causes severe respiratory diseases in young children. The HMPV 22 RNA genome is encapsidated by the viral nucleoprotein, forming an RNA-N complex (N<sup>Nuc</sup>), which 23 serves as template for genome replication and mRNA transcription by the RNA-dependent RNA 24 polymerase (RdRp). The RdRp is formed by the association of the large polymerase subunit (L), which 25 has RNA polymerase, capping and methyltransferase activities, and the tetrameric phosphoprotein (P). 26 P plays a central role in the RdRp complex by binding to N<sup>Nuc</sup> and L, allowing the attachment of the L 27 polymerase to the nucleocapsid template. During infection these proteins concentrate in cytoplasmic 28 inclusion bodies (IBs) where viral RNA synthesis occurs. By analogy to the closely related pneumovirus 29 respiratory syncytial virus (RSV), it is likely that the formation of IBs depends on the interaction between 30 P and N<sup>Nuc</sup>. However, the HMPV P-N<sup>Nuc</sup> interaction still remains to characterize. Here, we finely 31 characterized the binding domains involved in HMPV P and N<sup>Nuc</sup> interaction by studying binding between 32 recombinant proteins, combined with the use of a functional assay of the polymerase complex activity 33 and the study of the recruitment of these proteins to IBs by immunofluorescence. We show that the last 34 6 C-terminal residues of HMPV P are necessary and sufficient for binding to N<sup>Nuc</sup>, that P binds the N-35 terminal domain of N (N<sub>NTD</sub>), and identified conserved N residues critical for the interaction. Our results 36 allowed to propose a structural model of the HMPV P-N<sup>Nuc</sup> interaction.

37

#### 38 IMPORTANCE

39 Like respiratory syncytial virus (RSV), human metapneumovirus (HMPV) is a leading cause of severe 40 respiratory infections in children but also affects human populations of all ages worldwide. Nowadays, 41 no vaccine or efficient antiviral treatments are available for these two pneumoviruses. A better 42 understanding of the molecular mechanisms involved in viral replication could help the design or 43 discovery of specific antiviral compounds. In this work we have investigated the interaction between two 44 major viral proteins involved in HMPV RNA synthesis, the N and P proteins. We finely characterized 45 their domains of interaction, in particular a pocket on the surface of the N protein that could be used as 46 a template for the design of compounds interfering with N-P complexes and viral replication.

### 48 INTRODUCTION

49 Pneumonia is the leading cause of death among children younger than 5 years worldwide, and severe 50 pneumonia is more frequently caused by viruses than bacteria (1). After the closely related respiratory 51 syncytial virus (RSV), human metapneumovirus (HMPV) is recognized to be one of the most important 52 cause of viral bronchiolitis and pneumonia in young children, causing 7 to 19% of all cases of acute 53 respiratory tract infections (1, 2). HMPV infects mainly newborn, children, elderly and 54 immunocompromised individuals worldwide. This virus was first reported in 2001 from Dutch children 55 with acute lower respiratory tract illness, and serological studies have revealed that virtually every child 56 has been exposed to HMPV by the age of 5 years (3). The clinical features of HMPV infection display 57 as mid-to-upper respiratory tract infection, and can be severe enough to cause life-threatening 58 bronchiolitis and pneumonia. As yet, there is no effective treatment or licensed vaccine for HMPV.

Together with RSV, HMPV belongs to the *Pneumoviridae* family in the order *Mononegavirales* (4). HMPV is an enveloped virus that forms pleomorphic or filamentous virions. The virus genome is composed of a negative-sense single-stranded RNA of approximately 13.3 kb in size which encodes eight genes in the following order: 3'-N-P-M-F-M2-SH-G-L-5' (5, 6). The M2 gene of HMPV contains two overlapping open reading frames (ORFs), encoding for M2-1 and M2-2 proteins which precise functions during HMPV replication remain unclear.

65 The HMPV genome is encapsidated by multiple copies of the nucleoprotein (N) forming helical 66 nucleocapsids (N<sup>Nuc</sup>). This N<sup>Nuc</sup> serves as template for genome replication and mRNA transcription by 67 the viral polymerase complex formed by the large polymerase subunit (L) and its main cofactor the 68 phosphoprotein (P). After virus binding to the cell surface and virus-cell membrane fusion, mediated by 69 surface glycoproteins (F and G), nucleocapsids are released into the cytoplasm. Replication and 70 transcription of the viral genome take place within viro-induced cytoplasmic inclusions named inclusion 71 bodies (IBs) (7). These structures can be observed upon expression of P and N proteins alone (8), and 72 it was recently shown for RSV that the interaction between P and N<sup>Nuc</sup> is critical for the formation of IBs 73 (9).

Among the components of the polymerase complex, P plays a pivotal role as a cofactor of the L polymerase but also as a molecular hub between viral and cellular partners. HMPV P, 294 amino acid residues in length, forms homo-tetramers. The atomic structure of the coiled-coil oligomerization domain (residues 171-194) was resolved by crystallography (10). Small angle X-ray scattering (SAXS) studies 78 indicated that the flanking N-terminal (residues 1-170) and C-terminal (residues 195-294) regions 79 (named PNT and PCT respectively) are intrinsically disordered, some of them, such as residues 195-80 237, having  $\alpha$ -helical propensity (10, 11). More recently, the structure of the LP complex was resolved 81 by cryo-EM (12). It revealed a tentacular arrangement of P, with each of the four protomers adopting a 82 distinct conformation, demonstrating a "folding-upon-partner-binding" mechanism. Depending on the 83 protomer, the L-binding region involved regions 171-236, 172-217, 170-231 or 169-264. On the other 84 hand, by binding to the nucleocapsid, P mediates the attachment of the L protein to the N<sup>Nuc</sup> template 85 for viral RNA synthesis.

86 By analogy to RSV (13), the C-terminal domain of HMPV P is also thought to be the N<sup>Nuc</sup> binding 87 domain, but this has not been shown yet. It is noteworthy that the C-terminal extremity of P supposed 88 to bind to N<sup>Nuc</sup> was not visible in the crystal structure of the LP complex, indicating that this region is 89 disordered in the absence of N<sup>Nuc</sup>. Furthermore, the encapsidation of neosynthesized genome or 90 antigenome necessitates a pool of monomeric, RNA-free N, termed N<sup>0</sup>, which is kept in an unassembled 91 state through an interaction with P which plays the role of molecular chaperone, until delivery to the sites 92 of viral RNA synthesis. The crystal structures of recombinant HMPV N protein (394 residues) expressed 93 in E. coli and purified either as an RNA-free monomeric N in complex with the N-terminal residues of P, 94 or as rings of 10 N protomers complexed to RNA were resolved (14). In both states, the structures show 95 that N presents two globular domains (N<sub>NTD</sub> and N<sub>CTD</sub>) separated by a flexible linker that forms the RNA 96 groove, and N- and C-arms. In the oligomeric state, the N- and C-arms play a key role in the interaction 97 between N protomers and oligomerization, the N-arm binding to the flank of the N<sub>i-1</sub> protomer and the 98 C-arm binding atop the  $N_{i+1}$  protomer (*i* corresponding to the middle subunit of three adjacent N 99 protomers). In the  $N^0$  state, the important conformational changes consist in packing of the C-arm of N 100 in the RNA groove, impairing RNA binding.

101 In this work, based on the structural data of HMPV P and N proteins and their strong homologies 102 with RSV N and P proteins, we finely characterized the binding domains involved in HMPV P-N<sup>Nuc</sup> 103 interaction. By combining biochemical and functional cellular assays, coupled with a rational mutational 104 approach, we identified residues of P and N critical for their interaction. Our data show that the last C-105 terminal residues of P bind to the N<sub>NTD</sub>. These results allowed to establish a structural model of the 106 interaction which could be used for the rational design of antivirals targeting the N<sup>Nuc</sup>-P interaction of 107 HMPV.

108

# 109 **RESULTS**

#### 110 The HMPV PCT domain allows to purify N-RNA rings expressed in bacteria

By analogy to RSV, it is thought that the N<sup>Nuc</sup> binding domain of HMPV P is located at its C-terminus. 111 112 For RSV, it was shown that co-expression of the C-terminal disordered region of P (PCT, residues 161-113 241) fused to GST together with N in bacteria allows the purification of complexes formed by GST-PCT 114 and ring shape N-RNA oligomers (13, 15). Similarly, when expressed alone in E. coli, HMPV N also 115 forms decameric rings containing RNA (14). In a first attempt to characterize the N<sup>Nuc</sup> binding site of 116 HMPV P, we thus decided to co-express recombinant HMPV GST-PCT (residues 200-294) and N 117 proteins (from HMPV CAN 97-83 strain) in E. coli. The purified complexes were analyzed by SDS-PAGE 118 stained with Coomassie blue. As shown on figure.1A, N was co-purified with GST-PCT to > 95% 119 homogeneity. The PCT was then separated from GST by thrombin cleavage and the solubilized complex 120 was analyzed by size exclusion chromatography, following optical density (OD) at 220, 260 and 280 nm 121 (Fig. 1A). The elution profile showed a major peak (P1) with a OD<sub>260nm</sub>/OD<sub>280nm</sub> ratio > 1 and an apparent 122 mass of ~ 500 kDa, in agreement with the expected size of N-RNA decamers (Fig. 1B). A second peak 123 of lower intensity (P2), with an apparent mass of ~ 10 kDa was detected only at 220 nm. This peak 124 should correspond to PCT which does not present aromatic residues and a higher mass weight than 125 predicted due to the elongated shape of this fragment. The fractions of P1 peak were pooled; analysis 126 by SDS-PAGE stained with Coomassie blue of the sample showed the presence of a unique band 127 corresponding to N (Fig. 1B). The separation of PCT and N upon gel filtration reveals the relatively low 128 affinity of monomeric PCT for N. We then further analyzed the sample collected from the P1 peak by 129 combining dynamic light scattering (DLS) and electron microscopy (EM) approaches. The profile of DLS 130 measurement confirmed the homogeneity of the sample, with a single peak at 18 nm (Fig 1C), in 131 agreement with the size of N-rings observed by EM (Fig 1D).

Altogether, these results show that the GST-PCT fusion protein is sufficient to interact withnucleocapsid-like N-RNA rings.

134

# 135 The last 6 residues of HMPV P constitute the minimum domain for N<sup>Nuc</sup> binding

We thus investigated the minimal domain of HMPV P involved in N<sup>Nuc</sup> binding. By analogy with RSV for
which the 9 C-terminal residues of P are sufficient to interact with N (13), a series of GST fused peptides

138 of 9 to 1 amino acid long derived from the C-terminal of P was generated. These constructs were co-139 expressed with N in E.coli, and their capacity to purify N was analyzed by SDS-PAGE and Coomassie 140 blue staining. As shown on figure. 2A, the minimal sequence required for N purification corresponds to 141 the 6 last C-terminal residues of P. In parallel, alanine scanning was performed with the GST-P[285-142 294] construct to characterize the residues of P involved in the interaction with N<sup>Nuc</sup>. Again, GST-P 143 constructs and N were co-expressed in bacteria, followed by purification by Glutathione-Sepharose 144 beads affinity, and co-purification of N was analyzed by SDS-PAGE and Coomassie blue staining. Only 145 the four I289A, Y290A, L292A and M294A substitutions abrogated the interaction with N (Fig. 2B). In 146 order to further investigate the potential role of residues Q291 and I293 of P in the interaction with N, 147 the impact of the double mutations Q291A/I293A inserted into GST-PCT on N binding was also tested. 148 These mutations did not affect N binding, confirming that these two residues are not directly involved in 149 the interaction (Fig. 2C). These results confirm that the last residues of P are directly involved in the 150 interaction with N. Of note, our results suggest that HMPV P-N interaction mainly involves hydrophobic 151 interaction. As both hydrophobic and acidic residues of RSV P were previously shown to be critical for 152 the interaction with N (Fig. 2D), these results suggest that the binding of HMPV P on N differs from RSV 153 and involves specific interactions. We thus tested the capacity of each PCT fragments to pulldown N 154 proteins. Figure 2E shows that HMPV PCT and RSV PCT did not allow to purify RSV N and HMPV N 155 respectively. This last result confirms that HMPV and RSV N and P proteins cannot cross-interact, and 156 that pneumoviruses P-N interactions are specific to each virus.

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#### 158 Search for the P binding site on the surface of HMPV N

159 Although HMPV and RSV P-N<sup>Nuc</sup> interactions have their own specificity and cannot cross-react, the 160 strong homologies between N proteins suggest that the PCT binding site on N protein could be partially conserved. For RSV, the P binding site on  $N^{\text{Nuc}}$  is located at the surface of the  $N_{\text{NTD}}.$  The residues of 161 162 RSV NNTD involved in the interaction with P, which are partially conserved between HMPV and RSV 163 (Fig. 3A), were shown to form a well-defined pocket constituted of hydrophobic and positively charged 164 residues (16, 17). Corresponding residues of HMPV N<sub>NTD</sub> also form a pocket at the surface of the domain 165 (Fig. 3B). In a first attempt to characterize the HMPV P binding domain on N, we thus tested the capacity 166 of P to interact with the N<sub>NTD</sub>. The N<sub>NTD</sub> was expressed in *E. coli* and purified using a 6xHis tag, showing 167 that this domain is soluble when expressed alone (Fig. 3C). In parallel, the GST-P or GST-PΔM294

168 (deletion of the last C-terminal residue of P) proteins were expressed in bacteria either alone and purified 169 using the GST tag, or co-expressed with N<sub>NTD</sub> followed by purification by 6xHis tag. Analysis of the 170 samples by SDS-PAGE showed that  $N_{NTD}$  allowed to co-purify the wild type P, and that deletion of the 171 last residue of P was sufficient to impair the interaction (Fig. 3C). We then substituted by alanine the 172 residues of the HMPV N<sub>NTD</sub> identified by analogy with RSV, i.e. residues L46, L47, E50, Y53, D128, 173 R132, M135, R151, P152, and S153 (Fig. 3B). Again, the capacity of GST-P to co-purify N mutant 174 proteins was tested. Whereas mutants L46A, L47A, and P152 could still interact with P, only a weak 175 band of N was detected for mutants E50A, M135A, R151A and S153A, and mutations Y53A, D128A, 176 R132A abrogated the interaction (Fig. 3D). Of note, it was previously shown that similar punctual 177 mutations did not impact RSV N folding and solubility, suggesting that these mutations on HMPV N only 178 affect the capacity of P to interact with N. These results thus reveal a critical role of hydrophobic 179 residues, but also of negatively and positively charged residues of NNTD in P binding. Overall, these data 180 confirm that the P binding site on N presents strong homologies between HMPV and RSV.

181

#### 182 Validation of P-N<sup>Nuc</sup> binding domains in eukaryotic cells

183 As P-N<sup>Nuc</sup> interaction is required for viral polymerase activity, we then studied the impact of selected 184 mutations of P (M294A) or N (R132A and R151A) on HMPV replication/transcription using a bicistronic 185 subgenomic minigenome, pGaussia/Firefly. This construct contains the Gaussia and Firefly luciferase 186 genes under the control of gene start and gene stop sequences, as well as Leader and Trailer 187 sequences of HMPV genome (Fig. S1). Briefly, BSRT7 cells were transfected with plasmids coding for 188 N (Nwt, NR132A or NR151A), P (Pwt or PM294A), L, and M2-1 proteins, the plasmid pGaussia/Firefly 189 minigenome, and pSV-β-Gal (used to normalize transfection efficiencies). In this system, expression of 190 luciferases depends on the HMPV polymerase complex activity which can thus be quantified by 191 luminescence measurement. As shown on figure 4A, mutations M294A of P or R132A of N abrogated 192 HMPV polymerase activity and R151A induced a strong decrease on its activity, with only 30% of activity 193 compared to the control condition, although the levels of expression of these proteins were similar to 194 those of wild type proteins (Fig. 4B). These results correlate with those of pulldown assays, the mutant 195 NR151A that still presented a weak interaction with P displaying a residual polymerase activity. 196 In HMPV-infected cells, P-N<sup>Nuc</sup> interaction induces the formation of inclusion bodies (8). For RSV, we

197 have shown previously that IBs correspond to viral factories where viral RNA synthesis occurs (18). We

198 then assessed the impact of these mutations on the capacity of N and P to form pseudo-IBs when co-199 expressed in cells. To facilitate N detection, N-GFP fusion protein was generated to allow its detection 200 by epifluorescence. Based on previous studies on RSV P-N<sup>Nuc</sup> interaction, the GFP was fused to the C-201 terminus of N and this construct was co-expressed with wild-type N at a ratio of 1:1. Cells were thus co-202 transfected with plasmids encoding N-GFP, N (both WT or mutants R132A or R151A), and P (WT or 203 mutant M294A). The P protein was detected by immunolabelling using rabbit anti-P antibody, and nuclei 204 were stained with Hoechst 33342. When expressed together in BSRT7 cells, WT N-GFP/N and P were 205 found to co-localize into cytoplasmic inclusions similar to IBs observed upon HMPV infection (7, 8) (Fig. 206 4C). On the contrary, when expressing  $P_{M294A}$  in the presence of N<sub>WT</sub> or  $P_{WT}$  in the presence of N<sub>R132A</sub>, 207 P presented a diffuse cytoplasmic distribution and N protein was mostly detected in cytoplasmic 208 aggregates (Fig. 4C). These observations suggest that in the absence of N-P interaction, overexpressed 209 N presents a strong tendency to aggregate. Finally, co-expression of the mutant N<sub>R151A</sub> with P still 210 allowed to observe IBs, although less numerous and smaller than those observed in the presence of 211 WT proteins. This last result correlates with previous results showing that mutation R151A of N 212 drastically reduced but did not abrogate N-P interaction and polymerase activity. Altogether, these 213 results confirm that the residues of P and N previously identified as critical for N<sup>Nuc</sup>-P interaction *in vitro* 214 are also critical for the formation of IBs and polymerase activity in cells.

215

#### 216 Molecular modelling of PCT-N<sub>NTD</sub> interaction

217 To gain insight into the possible binding mode of P with N<sub>NTD</sub>, we built models of the last 6 residues of 218 P (peptide I289-YQLI-M294 named P6) bound to N, by analogy with the PCT-N<sub>NTD</sub> complex of RSV. For 219 RSV, only the last 2 residues of P (D240-F241) were resolved in the PCT-NNTD X-ray structures (17), 220 and are bound to a hydrophobic pocket of N similar to the one we characterized at the surface of HMPV 221 N. Using comparative modeling and refinement with HADDOCK (19), we obtained three clusters of 222 HMPV P6/NNTD complex structures representing possible binding modes of P6 (Table S1). The first two 223 clusters in terms of HADDOCK-score were the most populated and displayed similar Root-mean-square 224 deviation (RMSD) among their members (considering only the last three residues of P, i.e. residues 225 L292-I293-M294, called P3). In those two clusters, the C-terminal carboxyl group of M294 is involved in 226 an ionic interaction with the guanidinium of either R131 or R152 side-chains of N. Additionally, only L292 227 and M294 of P3 consistently display hydrophobic contacts with N. This result is consistent with our 228 mutation data showing that I293 of P is not required for binding to N, contrary to L292 and M294. The 229 10 best-scoring structures from cluster 1 and 2 are shown in Fig. S2, superimposed with the RSV 230 P2/NNTD structure (residues D240-F241 of P). Owing to the best HADDOCK-score and non-bonded 231 energies of cluster 2, we considered this cluster as most representative for the potential binding mode 232 of HMPV P3 to N<sub>NTD</sub>. Interestingly, more than half of the structures from this cluster display a hydrogen-233 bond between the backbone oxygen of I293 of P3 with the side-chain of R151 of N, while the side-chain 234 of R132 of N forms a salt-bridge with the C-terminal carboxyl group of M294 of P. Thus, the interactions 235 determined by molecular simulation correlate with the experimental data showing the importance of 236 R132 and R151 of N and of M294 of P for PCT-N interaction.

237 The interactability propensity of each modeled N<sub>NTD</sub> conformation was further profiled with our in-house 238 tool InDeep in order to select a subset of N<sub>NTD</sub> models being more likely to interact with the P3 peptide. 239 On this subset of N<sub>NTD</sub> conformations, the hydrophobic channel of InDeep was also used to locate 240 regions where hydrophobic moieties of P3 are expected to interact at the N<sub>NTD</sub> surface. Three 241 hydrophobic patches are detected on the N<sub>NTD</sub> surface, that present a good match with hydrophobic 242 moieties of P3 structures belonging to cluster 2 of HADDOCK: i) one occupied by L292 of P3; ii) another 243 occupied by M294 of P3; and iii) the last one located in an unoccupied region of the N pocket (Fig.5A). 244 Based on experimental and modelling information, the structure shown in Figure 5 represents a 245 consistent model of P3/N<sub>NTD</sub> complex. The N<sub>NTD</sub> conformation of this structure is predicted to be 246 favorable to protein-protein interaction by InDeep, and the P3 conformation, belonging to cluster 2 of 247 HADDOCK, binds to N<sub>NTD</sub> with a combined effect of electrostatic/polar interactions and hydrophobic 248 contacts in agreement with the mutational experiments. In addition, a procedure of virtual alanine 249 scanning using the same InDeep approach confirmed some of the mutagenesis results, such as the 250 importance of R132, M135, and R151 of N for the interaction with P. It also suggests that residue L139 251 of N may be key for the N<sub>NTD</sub> patch for partners' binding.

These analyses reveal that despite the different physico-chemical nature of residues involved in the interaction, PCT binding at the N surface is structurally similar between HMPV and RSV.

254

#### 255 **DISCUSSION**

Beyond its interest in better understanding the molecular mechanisms of virus replication, the study of
 protein-protein interactions required for Pneumoviruses' polymerase complex activity also presents

258 interest for the development of specific antiviral compounds. The activity of this complex depends on 259 multiple highly specific protein-protein interactions that have no cellular counterparts. Among those, 260 N<sup>Nuc</sup>-P interaction which is critical for viral polymerase activity constitutes a singular potential target for 261 antivirals. We have previously described RSV N<sup>Nuc</sup>-P interaction at the molecular level, showing that the 262 C-terminus of P binds to a well-defined pocket at the surface of N (9, 13, 16). We also made the proof 263 of concept that small chemical compounds can target this pocket of N and inhibit viral replication in cells 264 (17). More recently, Sá et al. showed that flavonoids derived molecules could also specifically bind to N 265 pocket and block its interaction with P (20). Given the strong homologies between RSV and HMPV N 266 and P proteins, it could be expected that a similar strategy could be used to block HMPV replication. It 267 is noteworthy that ideally, the discovery of compounds that could inhibit replication of these two closely 268 related viruses at the same time would be much more advantageous.

269 In the present study, we investigated the domains involved in HMPV P-N<sup>Nuc</sup> interaction using pulldown 270 assays between recombinant proteins (either truncated or mutated proteins) expressed in E. coli. As for 271 RSV, the HMPV N protein was purified as nanorings formed by N and bacterial RNA (Fig 1). Our results 272 revealed that mechanisms of binding between P and N<sup>Nuc</sup> are similar between RSV and HMPV, the C-273 terminus of P interacting with a pocket located on the surface of the N N-terminal globular domain N<sub>NTD</sub>. 274 However, it appeared that P and N amino acid residues involved in these interactions have different 275 contributions/roles between RSV and HMPV, and that the P-N<sup>Nuc</sup> interaction is specific of each virus. 276 More specifically, we showed that the minimal domain of HMPV P required for N binding is constituted 277 by the last 6 C-terminal residues of P which are mostly hydrophobic residues (IYQLIM), and determined 278 that only Q291 and I293 are not critical for the interaction (Fig. 2). This P fragment is thus shorter than 279 for RSV for which N binding minimal domain involves the 9 C-terminal residues of P and is composed 280 of both hydrophobic and acidic residues (13, 16). We also identified 7 residues of N involved in P binding, 281 i.e. residues E50, Y53, D128, R132, M135A, R151A and S153A, which form a well-defined pocket at 282 the NNTD surface (Fig 3). Of note, the pivotal role of residues R132 and R151 of N for polymerase activity 283 and IBs formation sustained in vitro binding assays (Fig. 4). Of most interest, residues R132 and R150 284 of RSV N were also shown to play a key role in P-N<sup>Nuc</sup> interaction, the aliphatic part of the R132 side 285 chain being involved in the stacking of the aromatic ring of the residue F241 of P. In order to better 286 understand the binding mode of HMPV P to N, molecular modelling was thus performed and the best 287 structural models support that R132 and R151 of N and of M294 of P are critical for PCT-N interaction

 $\begin{array}{ll} \text{(Fig. 5). Nevertheless, molecular modelling was restricted to the binding of the last 3 C-terminal residues} \\ \text{of P, and did not allow to predict the interaction of all the 6 critical residues of P on N. As also observed} \\ \text{in the crystal structures of RSV PCT/N_{NTD} complexes (17), the last C-terminal residue of P seems to} \\ \text{drive the binding to a well-defined N_{NTD} pocket and this primary binding could allow transient contacts} \\ \text{with upstream P residues outside of the N pocket.} \\ \end{array}$ 

293 In conclusion, our data confirm the strong structural homologies between HMPV and RSV P-N 294 complexes but also highlight some singularities. These data suggest that the HMPV P binding pocket 295 at the N<sub>NTD</sub> surface could represent a new target for the rational design of antivirals.

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- 297

# 298 MATERIALS AND METHODS

## 299 Plasmid constructions

The N and P sequences used for bacterial expression were derived from HMPV CAN97-83 strain. The sequence of N or N<sub>NTD</sub> (residues 30-255 of N) were cloned into pET-28a(+) vector (Novagen) at Ncol/Xhol or BamHI/Xhol restriction sites respectively, to allow bacterial expression of full-length N or N<sub>NTD</sub> domain with a C-terminal or N-terminal 6xHis tag. The sequence of P or PCT (residues 200-294 of P), were cloned into pGEX-4T3 vector, at BamHI and Smal sites to express in bacteria GST-P and GST-PCT fusion proteins. For the construction GST-P[285-294], complementary antiparallel oligonucleotides were hybridized and inserted at the BamHI/Smal sites in the pGEX-4T-3 vector.

307 A HMPV minigenome plasmid containing Gaussia/Firefly luciferases was designed and synthesized by 308 Genscript, cloned in pUC57 vector, and containing the trailer, leader, gene start (GS) and gene end 309 (GE) sequences derived from HMPV CAN97-83 strain (see Fig S1). The first ORF of this 310 pGaussia/Firefly minigenome codes for the Firefly luciferase and the second one codes for the Gaussia 311 luciferase. The plasmids pP, pN, pL, and pM2-1 corresponding to the sequences of NL99-1 HMPV strain 312 cloned into pCite vector. For expression of N-GFP fusion protein, the mGFP gene was amplified by PCR 313 and cloned in frame at the 3' end of N at the EcoRI site of pN, using In-fusion HD cloning kit (Takara 314 Bio). Point mutations were introduced in the P and N sequences by site-directed mutagenesis using the 315 Q5 site-directed mutagenesis Kit (New England Biolabs). Sequence analysis was carried out to check 316 the integrity of all constructs. All the oligonucleotides sequences are available on request.

#### 318 Antibodies

Antisera used in this study included polyclonal rabbit antisera raised against recombinant HMPV N and
P expressed in bacteria. A mouse monoclonal anti-β-tubulin (Sigma) and secondary antibodies
directed against mouse and rabbit Ig G coupled to HRP (P.A.R.I.S) were also used for immunoblotting.
Secondary antibody directed against rabbit Ig G coupled to Alexafluor-594 (Invitrogen) was used for
immunofluorescence experiments.

324

#### 325 Cell culture and transfections

BHK-21 cells (clone BSRT7/5) constitutively expressing the T7 RNA polymerase (21) were grown in Dulbeco Modified Essential Medium (Lonza) supplemented with 10% fetal calf serum (FCS), 2 mM glutamine, and antibiotics. The cells were grown at 37°C in 5% CO2, and transfected using Lipofectamine 2000 (Invitrogen) as described by the manufacturer.

330

## 331 Minigenome replication assay

332 BSRT7/5 cells at 90% confluence in 48-well dishes were transfected using Lipofectamine 2000 333 (Invitrogen) with a plasmid mixture containing 0.125 µg of pGaussia/Firefly minigenome, 0.125 µg of 334 pN, 0.125  $\mu$ g of pP (WT and mutants), 0.06  $\mu$ g of pL, 0.03  $\mu$ g of pM2-1, as well as 0.03  $\mu$ g of pSV- $\beta$ -Gal 335 (Promega) to normalize transfection efficiencies. Cells were harvested at 24 h post-transfection and 336 lysed in Firefly lysis buffer (30 mM Tris [pH 7.9], 10 mM MaCl2, 1 mM dithiothreitol [DTT], 1% [vol/vol] 337 Triton X-100, and 15% [vol/vol] glycerol). The Firefly luciferase activity was determined for each cell 338 lysate with an Infinite 200 Pro (Tecan, Männedorf, Switzerland) and normalized based on β-339 galactosidase (β-Gal) expression. Transfections were done in triplicate, and each independent 340 transfection experiment was performed three times. For proteins expression analysis, cells were lysed 341 in Laemmli buffer and analyzed by Western blotting (WB) using anti-N, anti-P, and anti-tubulin antibodies 342 according to standard protocols.

343

### 344 Fluorescence microscopy

345 Immunofluorescence microscopy was performed with cells grown on coverslips and previously 346 transfected with pN-GFP, pN (both WT or mutants R132A and R151A, at a ratio 1:1), and pP (WT or 347 mutant M294A). At 24 h post transfection, cells were fixed with 4% paraformaldehyde (PFA) for 25 min, 348 made permeable, and blocked for 30 min with PBS containing 0.1% Triton X-100 and 3% bovine serum 349 albumin (BSA). Cells were then successively incubated for 1 h at room temperature with primary and 350 secondary antibody mixtures diluted in PBS containing 0.3% BSA. For nucleus labeling, cells were 351 exposed to Hoechst 33342 stain (Invitrogen) during incubation with secondary antibodies. Coverslips 352 were mounted with ProLong Gold antifade reagent (Invitrogen) and observed with an inverted 353 fluorescence microscope (Zeiss Axiovision). Images were processed with ZEN software (Zeiss) and 354 ImageJ software.

355

# 356 Expression and purification of recombinant proteins

357 E. coli BL21 bacteria (DE3) (Novagen) transformed with pGEX-P plasmids (WT or mutant) or pET-N<sub>NTD</sub> 358 were grown at 37°C for 8 hours in 100 ml of Luria Bertani (LB) medium containing 100µg/ml ampicillin 359 or 50µg/ml kanamycin, respectively. Bacteria transformed with pET-N-derived plasmids together with 360 pGEX-P derived plasmids were grown in LB medium containing ampicillin (100 µg/ml) and kanamycin 361 (50 µg/ml). The same volume of LB was then added and protein expression was induced by adding 362 80µg/ml isopropyl-ß-D-thio-galactoside (IPTG) to the medium. The bacteria were incubated for 15 hours 363 at 28°C and then harvested by centrifugation. For purification using the GST-tag, bacteria were re-364 suspended in lysis buffer (50 mM Tris-HCl pH 7.8, 60 mM NaCl, 1 mM EDTA, 2 mM DTT, 0.2% Triton 365 X-100, 1 mg/ml lysozyme) supplemented with complete protease inhibitor cocktail (Roche), incubated 366 for 1 hour on ice, sonicated, and centrifuged at 4°C for 30 min at 10,000g. Glutathione-Sepharose 4B 367 beads (GE Healthcare) were added to clarified supernatants and incubated at 4°C for 3 hours. Beads 368 were then washed two times in lysis buffer and three times in PBS 1X, then stored at 4°C in an equal 369 volume of PBS. To isolate GST-free P fragments or N rings, beads containing bound proteins were 370 incubated with thrombin (Novagen) overnight at 20°C. Purified recombinant N proteins were loaded onto 371 a Superdex 200 16/30 column (GE Healthcare) and eluted in 20 mM Tris/HCl pH 8.5, 150 mM NaCl.

For purification of N<sub>NTD</sub>-6xHis fusion protein purification, bacterial pellets were re-suspended in lysis buffer (20 mM Tris-HCl pH8, 500 mM NaCl, 0.1% TritonX-100, 10 mM imidazole, 1 mg/ml lysozyme) supplemented with complete protease inhibitor cocktail (Roche). After sonication and centrifugation, lysates were incubated 30 min with chelating Sepharose Fast Flow beads charged with Ni<sup>2+</sup> (GE Healthcare). Beads were then successively washed in the washing buffer (20 mM Tris-HCl, pH 8, 500 mM NaCl) containing increasing concentration of imidazole (25, 50, and 100 mM), and proteins were

eluted in the same buffer with 500 mM imidazole. For SDS-PAGE analysis, samples were prepared in
Laemmli buffer, denatured 5 min at 95 °C, separated on 10% plolyacrylamide gel and detected by
Coomassie brilliant blue.

381

#### 382 Dynamic light scattering (DLS)

Size measurement of purified N oligomers 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 ( $D_H$ ) were calculated using the DLS software provided by the instrument manufacturer. The results were presented as size distribution (nm).

388

# 389 Negative stain electron microscopy observations of recombinant nucleoproteins

Three microliters of sample were applied to the clean side of carbon on a carbon-mica interface and stained with 2% sodium silicotungstate. Micrographs were recorded on a FEI Tecnai T12 microscope operated at 120 kV with a Gatan Orius 1000 camera. Images were recorded at a nominal magnification of 23 000 × resulting in a pixel size of 2.8 Å.

394

## 395 Molecular modelling

396 In the X-ray structures of HMPV N<sup>Nuc</sup> (PDB 5fvc) or N<sup>0</sup>-P (PDB 5fvd), the N<sub>NTD</sub> residues L110 to M113, 397 corresponding to a long beta-hairpin close to the PCT binding site, are not modeled. We used Modeller 398 (22) to first construct a complete model HMPV N<sub>NTD</sub> (residues 32 to 251) using as templates the X-ray 399 structures of HMPV N<sup>Nuc</sup> (residues 32 to 109 and 114 to 251) and RSV NNTD (PDB 4uc6) for the missing 400 beta-hairpin (residues 110 to 113). Next, using Modeller again, we generated 100 models of HMPV NNTD 401 complexed with the last six residues of P (peptide I289-YQLI-M294 called P6) using the RSV NNTD 402 structure in complex with the last two residues of RSV P (PDB 4uc9), i.e. D240-F241 (called P2), as 403 template for HMPV P6 by aligning the last 2 residues of HMPV-P and RSV-P. To compensate for 404 possible alternate conformations of exposed residues in apo and holo-form of N<sub>NTD</sub>, no template was 405 used for R132 and R151 side-chains. Next, P6-NNTD models where M294 side-chain of P is pointing 406 towards the N<sub>NTD</sub> hydrophobic pocket (by analogy with the binding of F241 of P on N RSV) were 407 selected, to obtain an ensemble 41 P6-N<sub>NTD</sub> models. Finally, the models were submitted to a water

408 refinement procedure using the refinement interface of the HADDOCK2.2 server (23). Each input P6-409 N<sub>NTD</sub> complex model was refined 20 times to obtain 820 refined models. The best 400 models with the 410 lowest HADDOCK-score were clustered based on similarity of the L292-I293-M294 residues of P, called 411 P3 (I289-Y290-Q291 were not considered for clustering since they appear very floppy in the refined 412 models and had no structural templates during homology modeling). Models were first superposed on 413 their NNTD backbone atoms and the RMSD was computed on the backbone atoms of L292-I293-M294 414 of P. Clustering was performed using the HADDOCK tool cluster\_struc (24) with a 2Å cut-off and a 415 minimal cluster size of 10.

416 In addition, the 820 P6-N<sub>NTD</sub> conformations generated by HADDOCK were profiled using InDeep 417 software in order to select the most suitable N<sub>NTD</sub> conformations to bind P3. InDeep is a deep-learning 418 model based on a FCN (Fully Convolutional Network) trained from known structures of protein 419 complexes capable of predicting protein-protein interaction interfaces. Each N<sub>NTD</sub> pocket conformation 420 is placed within a 3D grid composed of 1  $Å^3$  voxels. The model computes a probability value for each 421 voxel representing its interaction propensity. The predictions were performed on the putative P6 binding 422 site on NNTD surface. An "interactability-score" of the pocket is computed by taking the mean of the 150 423 best voxels probabilities. This value represents the binding propensity of a given N<sub>NTD</sub> pocket 424 conformation. Then, on the 200-top conformations with the highest "interactability-score" the 425 hydrophobic channel of InDeep was applied in order to locate the most probable hydrophobic patches 426 on the N<sub>NTD</sub> pocket where P3 hydrophobic side chains could interact. In addition, once a set most 427 promising conformations was identified, a virtual alanine scanning was performed using the same tool 428 but by mutating one by one the residues surrounding the patch of interaction of P3. The results are 429 provided as a set of correlations, one for each residue, highlighting the impact of the simulated mutation 430 on the interactability of the N<sub>NTD</sub>.

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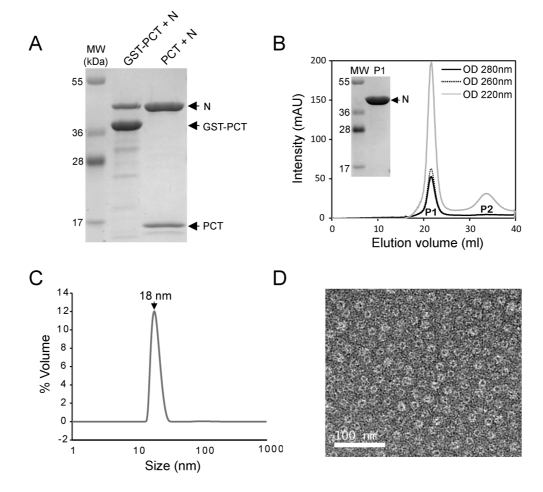
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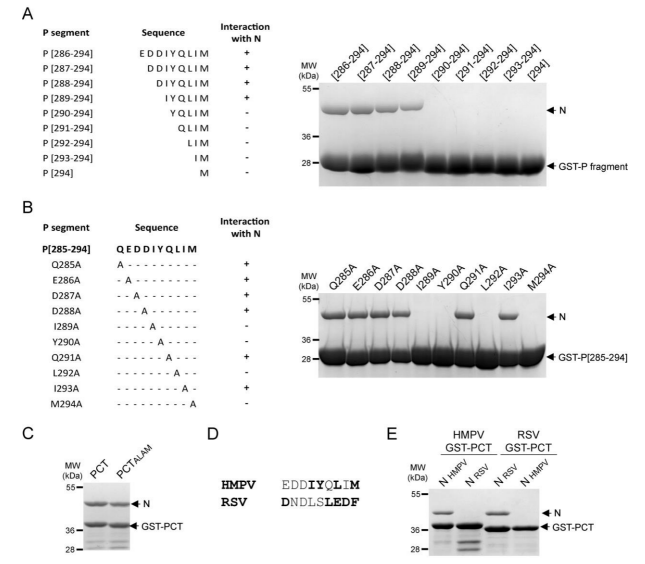
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- 441
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- 443 H.D., M.G. and J.-F.E. designed experiments. H.D., M.B. performed molecular, and cellular assays.
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- d45 observations. C.-A.R. performed gel filtration and *in vitro* study. M.G. wrote the paper with contributions
- from all authors, and J.-F.E. and M.G. edited the manuscript. All authors commented on the manuscript.
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# 450 Figure legends



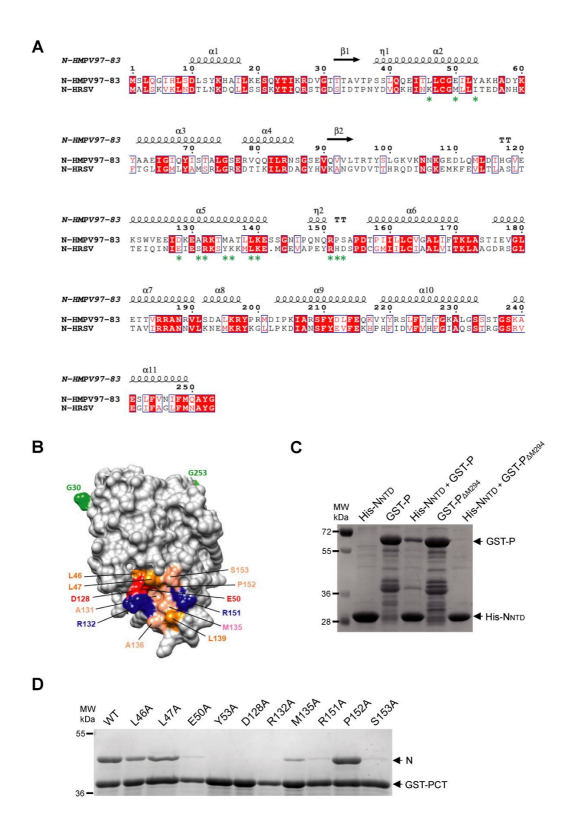
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452 Figure 1: Purification of HMPV N-RNA rings using GST-PCT. A/ GST-PCT and N proteins were co-453 expressed in E. coli, followed by purification using the GST tag. The product of purification was analyzed 454 by SDS-PAGE and Coomassie blue staining (left lane). The sample was then incubated in the presence 455 of thrombin in order to cleave GST (remaining on beads) and isolate PCT-N in the supernatant (right 456 lane), B/ Gel filtration profile of purified PCT-N complex. The curves corresponding to OD spectra at 220 457 nm, 260 nm, and 280 nm are presented. P1 and P2 indicate the two peaks detected. The fractions 458 corresponding to P1 were pooled and the sample was analyzed by SDS-PAGE colored with Coomassie 459 blue, C/ Dynamic light scattering (DLS) analysis of the purified N protein, showing a homogenous peak 460 at 18 nm, corresponding to N oligomers. D/ Image of purified HMPV N-RNA rings as observed by 461 negative stain electron microscopy. Bar, 100 nm.



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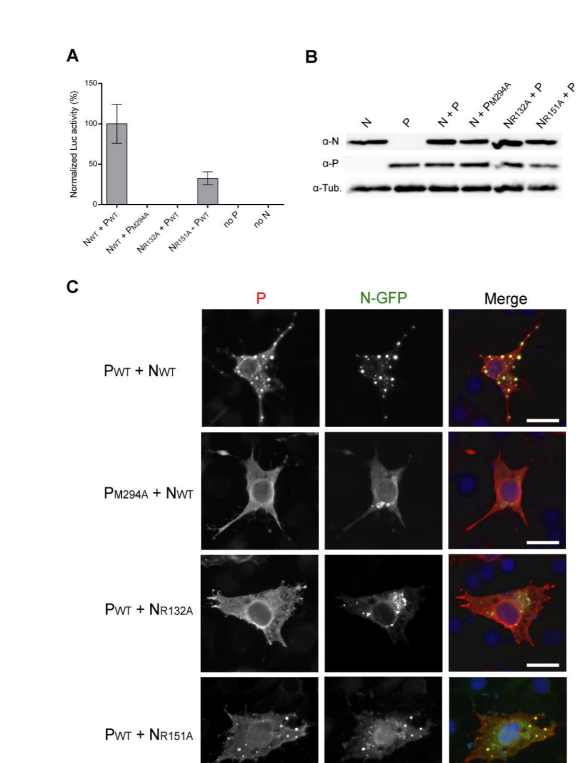
464 Figure 2: Identification of the residues of P involved in N binding. A/ GST-P fragments and B/ GST-465 P[285-294] (WT or mutants) (sequences indicated on the left) were co-expressed with N in bacteria, 466 followed by purification using the GST tag. The products of purification were analyzed by SDS-PAGE 467 and Coomassie blue staining (right). C/ GST-PCT or GST-PCTALAM corresponding to the double substitution of residues Q291 and I293 residues of P by alanine were co-expressed with N. The products 468 469 of purification by GST were analyzed by SDS-PAGE and Coomassie blue staining. D/ Sequence 470 alignment of the 9 last C-terminal residues of HMPV and RSV P proteins. The residues critical for the 471 interaction with the N protein are indicated in bold. E/ HMPV and RSV GST-PCT (residues P(200-294) 472 for HMPV and P(161-241) for RSV) constructs were co-expressed with either HMPV or RSV N proteins. 473 Copurification of N proteins by GST-PCT fragment was analyzed by SDS-PAGE and Coomassie blue 474 staining.



476

Figure 3: Search for the PCT binding site on HMPV N. A/ Amino acid sequence alignments between
the N-terminal domains of N (residues 1-252/253) proteins of HMPV (strain CAN 97-83) and human
RSV (HRSV, strain Long VR-26). Invariant residues are highlighted in white font on a red background.
The secondary structure elements observed in the crystal structure of HMPV N protein (11) are indicated
above the sequence. Green asterisks below the sequence indicate the residues constituting the P
binding pocket of RSV N. Uniprot accession codes: NCAP\_HRSVA (Human RSV); A1DZS3\_9MONO
(Human MPV). Sequences were aligned with Clustal W and treated with ESPript 3. B/ Surface

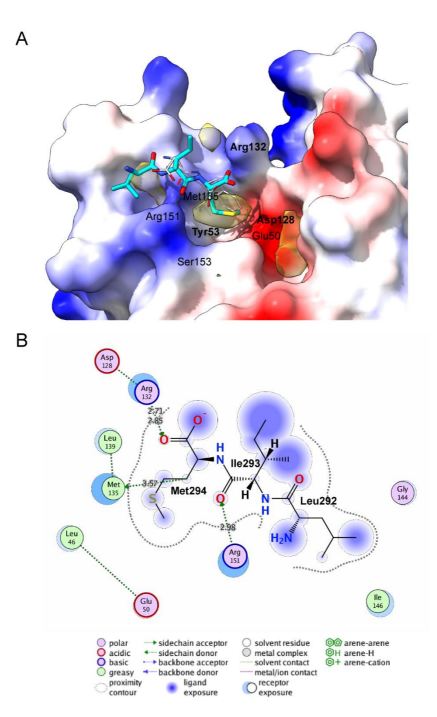
representation of HMPV N<sub>NTD</sub> (from Gly30 to Gly 253, indicated in green) showing the potential P binding pocket, with acidic amino acids colored in red, basic residues in blue, and hydrophobic residues in orange, C/ GST-P or GSP-P<sub> $\Delta$ M294</sub> and N<sub>NTD</sub>-6xHis proteins were co-expressed in *E. coli*, followed by purification using the 6xHis tag. GST-P or GSP-P<sub> $\Delta$ M294</sub> alone were purified using the GST-tag. The products of purification were analyzed by SDS-PAGE and Coomassie blue staining, D/ GST-PCT and mutant N proteins were co-expressed in *E. coli*, followed by purification using the GST tag. The product of purification was analyzed by SDS-PAGE and Coomassie blue staining.



#### 493

Figure 4: The residues involved in P-N<sup>Nuc</sup> interaction are critical for the polymerase activity and the
formation of inclusion bodies. A/ HMPV polymerase activity in the presence of P and N mutants.
BSRT7/5 cells were transfected with plasmids encoding the P (WT or M294A mutant), N (WT, or R132A
and R151A mutants), L, and M2-1 proteins, the pGaussia/Firefly minigenome, together with pCMV-βGal
for transfection standardization. Viral RNA synthesis was quantified by measuring the Firefly luciferase
activity after cell lysis 24 h after transfection. Each luciferase minigenome activity value was normalized
based on β-galactosidase expression and is the average of three independent experiments performed

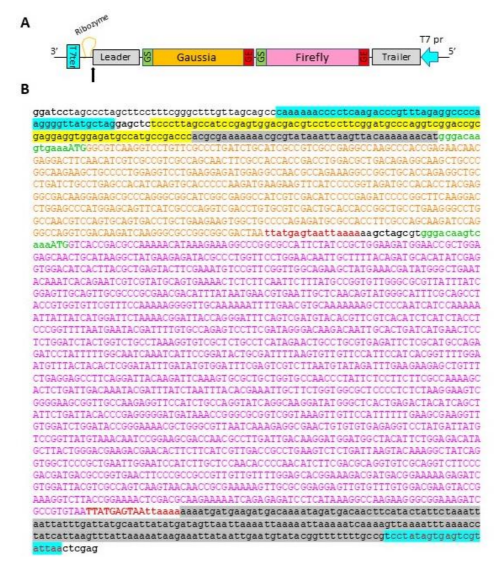
in triplicate. Error bars represent standard deviations calculated based on three independent
experiments made in triplicate. B/ Western blot showing the expression of N protein variants in BSRT7/5
cells. C/ Impact of P and N mutations on the formation of HMPV cytoplasmic IBs. N-GFP and P (or
mutants) proteins were coexpressed in BSRT7/5 cells; cells were then fixed 24 h post-transfection,
labeled with anti-P antibody and the distribution of viral proteins was observed by fluorescence
microscopy. Nuclei were stained with Hoechst 33342. Scale bars, 10 µm.



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**Figure 5:** Model of HMPV P3/N<sub>NTD</sub> complex from HADOCK refinement. A/ The N<sub>NTD</sub> protein is colored by electrostatic surface and the P3 peptide (residues L292-I293-M294 of P) is shown as sticks. The residues of N<sub>NTD</sub> critical for the interaction with P are indicated, with residues for which mutation to alanine abrogate P binding are labeled in bold. Yellow Surface shows InDeep hydrophobic channel prediction where hydrophobic contacts are expected to occur at the N<sub>NTD</sub> surface. B/ Interaction diagram between P3 peptide (Leu292-IIe293-Met294) and N<sub>NTD</sub>. Legend indicated below the diagram.

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519 Supplementary Figure 1: Minigenome construct. A/ Schematic representation of the elements 520 constituting the minigenome sequence. The sequences of the leader and trailer (grey), gene start (GS, 521 green) and gene end (GE, red) are derived from HMPV stain CAN98-87. The full sequence is framed 522 by T7 promoter (T7pr, blue arrow) and terminator (T7ter, blue) sequences. The black arrow indicates 523 the cleavage site induced by the presence of the Ribozyme (yellow). B/ Nucleotide sequence of the 524 HMPV minigenome. The elements are highlighted according to the color code used in A/. Sequences 525 of Firefly and Gaussia luciferase are indicated by uppercase letters. The enzymatic restriction sites are 526 indicated in bold. 527

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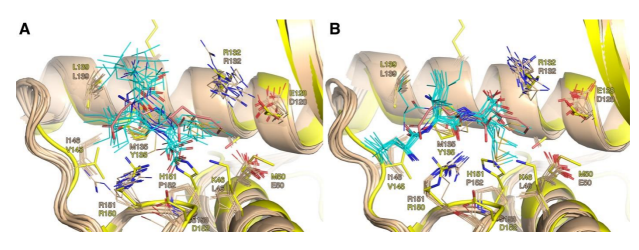
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#### Supplementary Table 1. Cluster statistics of HADDOCK refined models obtained for HMPV P6/NNTD

#### complex

	Cluster 1	Cluster 2	Cluster 3
HADDOCK score (all)	-47.3 +/- 4.3	-52.5 +/- 7.9	-45.5 +/- 3.4
HADDOCK score (top10 <sup>a</sup> )	-57.8 +/- 2.8	-69.3 +/- 4.0	-49.9 +/- 2.8
Cluster size	198	134	36
Cluster RMSD (Å)	0.95 +/- 0.28	0.99 +/- 0.29	1.12 +/- 0.56
RMSD from the overall lowest-energy structure Å	1.69 +/- 0.18	0.99 +/- 0.29	1.47 +/- 0.24
Van der Waals energy	-15.8 +/- 3.2	-19.3 +/- 3.4	-15.8 +/- 3.0
Electrostatic energy	-83.5 +/- 26.1	-87.1 +/- 20.0	-77.1 +/- 12.3
Buried Surface Area (Ų)	620.4 +/- 101.1	673.7 +/- 102.8	606.1 +/- 104.4
lonic/polar interactions frequencies			
(N)R132(P)M294 salt-bridge	2.0 %	76.9 %	2.8 %
(N)R151(P)M294 salt-bridge	75.3 %	21.6 %	75.0 %
(N)R132-Nη1/2(P)I293-O H-bond	0 %	-	0 %
(N)R151-Nη1/2(P)I293-O H-bond	-	57.5 %	-
# Hydrophobic inter-molecular contacts <sup>b</sup>			
(P)L292	1.0 +/- 1.2	1.0 +/- 1.4	0.0 +/- 0.2
(P)I293	0.3 +/- 0.8	0.3 +/- 0.6	0.4+/- 0.8
(P)M294	3.6 +/- 2.3	4.2 +/- 2.2	3.3+/- 1.7

<sup>a</sup> 10 lowest-energy structure of the cluster ; <sup>b</sup> average number of contacts between Carbon or Sulfur atoms within 3.9 Å



Supplementary Figure 2. 10 best-scoring structures of clusters 1 (A) and 2 (B) from HADDOCK refinement of HMPV P3 (residues L292-I293-M294 of P)/N<sub>NTD</sub> complex superimposed with RSV P2/N<sub>NTD</sub>. N<sub>NTD</sub> is colored in beige (HMPV) or yellow (RSV), and HMPV P3 peptide in cyan (HMPV) or RSV P2 peptide (corresponding to residues D240-F241 of P) in pink. Side-chains of residues in contact with the P peptide in HADDOCK models are also shown, along with their RSV equivalent.

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