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## **3D structure of three jumbo phage heads**

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## 29 Running title: 3D structure of three jumbo phage heads

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32

#### 34ABSTRACT

35Jumbo phages are bacteriophages that carry more than 200 kbp of DNA. In this study we 36 characterized two jumbo phages ( $\Phi$ RSL2 and  $\Phi$ XacN1) and one semi-jumbo phage ( $\Phi$ RP13) 37at the structural level by cryo-electron microscopy. Focusing on their capsids, three-38 dimensional structures of the heads at resolutions ranging from 16 Å to 9 Å were calculated. 39Based on these structures we determined the geometrical basis on which the icosahedral 40 capsids of these phages are constructed, which includes the accessory and decorative proteins 41that complement them. A triangulation number novel to *Myoviridae* (**PRSL2**; T=21) was 42 discovered as well as two others which are more common for jumbo phages ( $T=27$  and 43T=28). Based on one of the structures we also provide evidence that accessory or decorative 44 proteins are not a prerequisite for maintaining the structural integrity of very large capsids.

45

#### **Introduction** 47

48Bacteriophages are viruses that infect bacteria. They are extremely numerous and the number 49 of known bacteriophages has increased at a rate of approximately 100 per year for decades 50[1]. An increasing number of studies have suggested that bacteriophages are an attractive 51 option for alternatives to antibiotics [2]. Bacteriophages can be polyhedral, filamentous, or 52 pleomorphic and may have either a long, short, contractile, or flexible tail. Some are even tail-53less. They can contain either single- or double-stranded DNA or RNA. Caudal bacteriophages 54 represent the vast majority of known bacterial viruses and have been classified in different 55 families according to their tail morphology which include Siphoviridae (long flexible tail), *Myoviridae* (long contractile tail), and *Podoviridiae* (short tail). Bacteriophages carrying more 56 57than 200 kbp of DNA are commonly known as "jumbo phages"  $[3, 4]$ .

58

59The capsids of phages have icosahedral symmetry and are constructed from a basic brick 60 which, until now, has always been based on the canonical structure of the HK97 major capsid 61 protein. The number of currently known jumbo phages is around 100 [5] with the dimension 62of these phages varying from 100 to 160 nm in diameter for the head and a triangulation 63number between 19 and 52 [6]. Only a few of the known jumbo phages have been 64characterized structurally, including  $\Phi$ kZ [7],  $\Phi$ RSL1 [8],  $\Phi$ M12 [9] and  $\Phi$ N3,  $\Phi$ Pau,  $65\Phi$ PBS1,  $\Phi$ 121Q, and  $\Phi$ G [6]. The highest resolution of their three-dimensional structures has 66been limited to 9 Å. Particularly within the jumbo phage family and more generally in the 67bacteriophage world, viruses use different types of accessory or decorating proteins. Because 68of the large size of their genome, they also exhibit large heads with high triangulation 69numbers (usually higher than 20). The larger the genome is, the larger the capsid has to be. 70This is a general rule in the virus world with a correlation between genome length and capsid 71 size<sup>[10]</sup>. However there are deviation to the rule: for the characterized jumbo phages, the 72average density of packed DNA has been measured experimentally to be between 0.39 and 730.55 bp/nm3 which represent a variation of  $40\%$  [6].

74

Towards the aim of contributing to a better understanding of the structural characteristics of 75 76 jumbo phages including DNA packing, we examined the structures of two large jumbo phages  $77(\text{QRSL2}$  with a medium-sized genome of 224 kbp [11] and  $\Phi$ XacN1with a large genome of 78385 kbp [12]) and one smaller semi-jumbo phage (ΦRP13) carrying also a smaller genome of 79 about 180 kbp (phages with a genome smaller than 200 kbp but close to 200 kbp are defined 80as semi-jumbo phages). Here we present three new three-dimensional structures of jumbo 81 phage heads: two of them representing new examples of known triangulation numbers ( $T=27$ 82and T=28) and one new type of geometry (T=21) never described for a *Caudovirales*. Two of 83phage heads are decorated with different proteins (on the outside and also on the inside of the 84 capsid) but  $\Phi$ XacN1 appears to be naked, with the capsid built only by one type of protein 85(with the exception of the vertices). The DNA packing density is in the common range of 0.39 86and 0.55 bp/nm3 for  $\Phi$ XacN1 and  $\Phi$ RP13 but much lower for  $\Phi$ RSL2 (0.29). All together 87this is showing that there is no rule that can be applied for phages neither in the capsid 88 composition nor in the genome-capsid size correlation.

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90

#### **Materials and Methods** 91

#### 92**Bacteriophage production and purification**

93Ralstonia phages PRSL2 [11] and PRP13 [13] were isolated from Japan and Thailand, 94 respectively. They were propagated with *Ralstonia solanacearum* MAFF 106603 as the host. 95Host bacterial cells were cultured in CPG medium containing  $0.1\%$  (w/v) casamino acids, 961.0% (w/v) peptone, and 0.5% (w/v) glucose [14] at 28°C with shaking at 200-300 rpm. When 97the cultures reached an  $OD_{600}$  of 0.05, each bacteriophage was added at a multiplicity of 98infection (MOI) of 0.1. After culturing for a further 12-24 h, the cells were removed by 99 centrifugation at 5,000 x g for 15 min at  $4^{\circ}$ C in a R12A2 rotor in a Hitachi himac CR21E 100 centrifuge. The supernatant was membrane-filtered (0.45-um pore; Steradisc, Kurabo Co. 101Ltd., Osaka, Japan), and the pellet was dissolved in SM buffer (50 mM Tris-HCl at pH 7.5, 102100 mM NaCl, 10 mM MgSO<sub>4</sub>, and 0.01% gelatin) after centrifugation at 15,000  $\times$  *g* for 1h at 1034°C. For further purification, the phage suspension was layered on a 20-60% sucrose gradient 104 and centrifuged with a P28S rotor in a Hitachi CP100 $\beta$  ultracentrifuge at 40,000  $\times$  *g* for 1 h. 105The purified phages were stored at  $4^{\circ}$ C. Xanthomonas phage  $\Phi$ XacN1 [12] was isolated in 106Japan and propagated with *Xanthomonas citri* MAFF 301080 as the host. *Xanthomonas* cells 107were cultured in NB medium (Difco, BBLBD, Cockkeysville, MD, USA) at 28°C with 108shaking at 220 rpm. When cultures reached an OD<sub>600</sub> of 0.03,  $\Phi$ XacN1was added at a MOI of 1090.1. After culturing for a further 12-24 h, the cells were removed by centrifugation at 5,000 x 110g for 15 min at  $4^{\circ}$ C and the supernatant was membrane-filtered as above.  $\Phi$ XacN1 was 111 pelleted by centrifugation at  $15,000 \times g$  for 1h at 4°C and dissolved in SM buffer as above. 112For further purification, the phage suspension was layered on a 20-60% sucrose gradient and 113 centrifuged at  $40,000 \times g$  for 1 h as above.

115

116 Negative staining electron microscopy

117Negative-stain grids were prepared using the mica-carbon flotation technique [15]. Briefly, 118 samples were adsorbed on the clean side of a carbon film previously evaporated on mica and 119then stained using  $2\%$  (w/v) Ammonium Molybdate pH 7.5 for 30 s. The sample/carbon 120 ensemble is then transferred to a grid and air-dried. Images were acquired under low dose 121 conditions (<30 e<sup>-/</sup>Å<sup>2</sup>) on a Tecnai 12 FEI electron microscope operated at 120 kV using a 122Gatan ORIUS SC1000 camera (Gatan, Inc., Pleasanton, CA).

123

#### 124Cryo-EM

3.5 μl of concentrated sample were applied to glow discharged (25 mA, 40 s) R3.5/1 125 126 quantifoil copper grids (Quantifoil Micro Tools). The excess of solution was blotted using a 127Vitrobot (20 $\degree$ C, 100% humidity, 2-s blotting time, and blot force 1) and subsequently flash-128 frozen in liquid ethane.

129The grids were transferred to a Tecnai F30 Polara electron microscope working at 300 kV. 130 Movies (40 frames of 0.1 s and a dose of 1 electron/ $A^2$  per frame) were recorded manually on 131a K2 summit direct electron detector using the low dose module in the GMS3 software 132(Gatan) software at a nominal magnification of  $\times$ 12,000 in super resolution mode (1.64 Å per 133 pixel at the sample level for  $\Phi$ RSL2) and x20,000 in counting mode (1.94 Å per pixel at the 134 sample level for  $\Phi$ XacN1 and  $\Phi$ RP13).

135

## 136Image Analyses

137For the ΦRSL2 dataset, the re-alignment of the frames has been performed automatically

138 using the Latitude S software. For the two other datasets, Motioncor2 has been used excluding 139 frames 1 and 2 [16]. CTF parameters were determined using GCTF [17].

140

141ΦRSL2

142The images have been binned four times (final pixel size of  $6.57 \text{ Å}$ ). The initial 3D model of 143full ΦRSL2 capsids has been calculated with the RIco software [18]. Thereafter all image 144 analyses and capsid reconstructions have been performed using the Relion software [19] 145 imposing icosahedral symmetry. The final reconstruction includes 250 particles out of 499 146 for a resolution of 16 Å (FSC determined using the gold-standard method implemented in 147 Relion [19] at 0.143 threshold; Supplementary Figure 1).

148  $\Phi$ XacN1 and  $\Phi$ RP13

149The images have been binned two times (final pixel size of 3.88 Å). All the image analysis 150 including generation of an initial model have been performed using the Relion software [19]. 151 The final reconstruction of  $\Phi$ XacN1 and  $\Phi$ RP13 respectively includes 1149 and 669 particles 152for a resolution of 9.1 Å and 9.4 Å (FSC determined using the gold-standard method 153 implemented in Relion [19] at 0.143 threshold; Supplementary Figure 1).

154Reducing the binning to 2 for  $\Phi$ RSL2 and no binning for  $\Phi$ XacN1 and  $\Phi$ RP13 did not 155 improve the resolution of the corresponding map.

156 Figures were generated using Chimera [20]. All the statistics are summarized in Table 1. 157

158Fitting of the bacteriophage HK97 MCP into the EM map

159The X-rays structure of HK97 MCP (pdb 2FT1) was fitted into the  $\Phi$ XacN1 and  $\Phi$ RP13 map 160 using Chimera [20]. Briefly the entire structure (7 monomers) was first roughly placed by 161 hand in the EM map and in a second step only one monomer was used. For this second step, 162the long alpha helix of HK97 monomer, which is easily recognizable was used as a landmark. 163 Final refinement of the fitting was performed using the Chimera function "fit in map".

#### 164

#### 165Results

166Two of the bacteriophages described here ( $\Phi$ RSL2 [11] and  $\Phi$ RP13 [13]) were isolated from 167the phytopathogen Ralstonia solanacearum. The third one (**ØXacN1** [12]) also a jumbo 168phage, was isolated from the phytopathogen *Xanthomonas citri*. The three phages belong to 169the *Caudovirales* order and exhibit the typical *Myoviridae* morphology with a contractile tail 170 and an isometric head (Figure 1). Negative staining images of  $\Phi$ RSL2 (Figure 1A) clearly 171show that the tail is decorated by fibres at two different levels (arrows). For  $\Phi Xa cN1$ , an 172 annular structure of unknown function is present around the tail (Figure 1B, arrow). ΦRP13 173 differs from the other two phages because it exhibits a double-layered baseplate like the 174Twort-like phage  $\Phi$ 812 [21]. The [tail length]: [capsid diameter] ratios are quite different 175 among the different viruses (Table 2). For this study we mainly focused our structural 176 analyses on the virus head.

177

#### 178 **PRSL2**

179Cryo-electron microscopy images show that the sample was a mixture between 180bacteriophages with a head full of DNA and others which have released their DNA (appearing 181as light shades in the images). Only capsids full of DNA were selected to perform icosahedral 182 image analysis. The resulting structure shows a capsid having a diameter of 139 nm from

183 vertex to vertex (5-fold axis) and 128 nm along the 2-fold axis. The triangulation number, 184 which determines the number of protein copies forming the capsid, is  $T=27$  and was deduced 185from the hexagonal lattice present on the surface. It was calculated using the formula  $186T=h^2+hk+k^2$  where h and k are the number of local symmetry axes to be crossed to go from 187one 5-fold axis to the next [22]. For  $\Phi$ RSL2, the observed numbers were h=3 and k=3 (Figure  $1882A$ ).

#### 189

190The resolution obtained for  $\Phi$ RSL2 was limited to 16 Å due to a limited number of particles 191(250 particles; Figure 2A and Table 1). Each facet of the capsid is flat and composed of 13 192hexamers. These hexamers as well as the pentamers are most probably made of the same 193 protein: the major capsid protein (MCP) which, for this virus, is encoded by ORF117 194(predicted size of 82,440 Da but observed size of 70 kDa according to SDS-PAGE and LC-MS/MS analyses as described before [11]). This is the largest known MCP. An icosahedral 195 196 capsid with a triangulation number of  $T=27$  is assembled from  $27x60$  asymmetric units (or 19720x13 hexamers plus 12 pentamers). In the case of a caudal bacteriophage, a pentamer must 198be removed because one of the vertices is occupied by the portal. The total number of MCPs 199 in this capsid is therefore 1615 since it appears at this resolution that the 11 vertices that do 200 not bind the tail are composed of the same protein.

#### 201

202A hollow tube can be found at the centre of each hexamer with a diameter of 40 Å and a 2031ength of 70 Å. The 260 cylindrical structures project outward from the capsid and appear to 204 cross the capsid and slightly extend out from the inner face of the hexamer (Figure 3C). The 205 stoichiometry of this protein is difficult to assess at this resolution as it lacks recognisable 206 features. The outside of the capsid is further decorated with another cylindrical protein (70 Å 207in length and 30 Å in diameter) that is bound to the periphery of the hexamers and lying 208 parallel to the capsid surface. It associates in 810 dimers and forms bridges between 209 neighbouring hexamers/pentamers (Figure 3B). The structure and organisation of the dimer is 210 reminiscent from that observed in the  $\Phi$ KZ [7] and  $\Phi$ PBS1 capsids [6] which have the same 211triangulation number. It is interesting to note the presence of extra densities on the inside of 212the capsid, at the level of the 5-fold axis (Figure 3A, black arrows). This kind of structure is 213 different from the one observed in  $\Phi$ RSL1 which exhibits a much more complex structure 214 made of a trimer and a dimer [8]. Isosurface visualisation of these densities from the inside of 215the capsid (Figure 3D) shows that they are directly connected to the capsid at the 5-fold axis 216level (central globular structures) but also to the base of the cylindrical structures present at

217the centres of each hexamer. The nature of these densities is unknown, but they may 218 correspond to proteins or DNA/protein complexes that link the DNA to the capsid and allow 219the DNA to be organised.

#### 220

221 Focusing on the inside of the capsid, one can note the absence of DNA organised in 222 concentric layers which is probably not due to the lack of resolution as it becomes visible at 223about a 20 Å resolution. It is possible to distinguish a hexagonal mono domain organisation of 224DNA with a cylindrical rod spanning the interior of the head and oriented along the tail axis 225(dotted rectangle at bottom of Figure 1A; Supplementary Figure 2C). When  $\Phi$ RSL2 was 226 exposed to a very high dose of electrons ( $>100$  e $\angle$ A<sup>2</sup>), an "inner body" similar to the one 227 observed for  $\Phi KZ$  [23] and  $\Phi$ 121 $Q$  [6] can be visualized. However, this bubblegram is 228 slightly different from that observed with  $\Phi KZ$  as it has an arch at one end of the cylinder that 229 is close to the tail. This arch is reminiscent of the structures visible under the 5-fold axis in 230Figure 3D.

231

#### *XacN1* 232

2331149 particles have been used out of 1775 to obtain a 9 Å resolution three-dimensional map 234of the  $\Phi$ XacN1 head. Only particles loaded with DNA were analysed. The diameter of the 235final reconstructed full capsid is 139.5 nm from vertex to vertex and 116 nm along the 2-fold 236 axis (Figures 1B and 2B). The triangulation number of this head is  $T=28$  (h=4, k=2). The 237 major capsid protein of  $\Phi$ XacN1 is 463 aa in length and about 49 kDa in mass as described 238before [12]. Even with the medium resolution of this reconstruction, it was possible, due to the 239 presence of a long "spinal" alpha helix, to unambiguously fit the X-ray structure of HK97 240[24] into the cryo-electron microscopy map (Supplementary Figure 3A). The handedness of 241 the structure is therefore most likely to be *dextro* (T=28,d). There are only two other known 242capsids from the *Caudavirales* order where three-dimensional reconstructions show such 243 symmetry:  $\Phi$ 121Q [6] and PhAPEC6 [25].

244

245 The structure of the external and internal faces of the capsid is very smooth compared to other 246phages (Figures 2B, 3F, and 3G). This is especially true if compared with  $\Phi$ 121Q which 247 harbours two types of decoration proteins at the periphery and on the middle of the hexamer. 248The only protruding components are located at the 5-fold axis with the presence of a turret-249like structure (dimensions of 38 Å in height and 60 Å in diameter; Figure 3F). This kind of 250 extension is quite common in the bacteriophage world  $[8, 9, 26]$ . Inspection of the central

251 slice of the three-dimensional reconstruction clearly shows that there are at least six 252 concentric layers of DNA separated by 23.6 Å (Figure 3E). When irradiated at a high electron 253 dose, the  $\Phi$ XacN1 head did not exhibit any bubblegram-type structure (data not shown). 254

## *RP13* 255

256Image analysis of the DRP13 head started with 1311 particles. The best 669 particles yielded 257a three-dimensional reconstruction image at 9 Å resolution. This capsid shows a triangulation 258 number of  $T=21$ , which was the smallest triangulation (and dimensions) of the three jumbo 259phages analysed here. The diameter of the particle from vertex to vertex is only 114 nm and 260between two opposed two-fold axes it is 97 nm. The obtained resolution enabled fitting of the 261HK97 X-ray structure into the  $\Phi$ RP13 capsid reconstruction leading to the assumption that the 262capsid handedness is T=21 *laevo* (supplementary Figure 3B). The **PRP13** phage is the first 263HK97-related phage to exhibit this kind of triangulation number. Only lipidic phages like 264 $\Phi$ PM2 [27], FLiP (*Flavobacterium*-infecting, lipid-containing phage [28]), or P23-77 [29] 265have shown an organization with similar geometry but with a pseudo *T*=21 *dextro* 266triangulation number. These types of phages do not use the canonical HK97 hexameric 267 structure to build up their capsid but rather incorporate an adenovirus-like trimeric structure. 268Based on the three-dimensional structure it is clear that the vertices of the capsid are built by 269the same protein as the facet (Figures 3I and 3J, pentamer and hexamer). The capsid is 270 therefore composed of 1255 copies of the major capsid protein  $(20x10)$  hexamers per facet 271 plus 11 pentamers).

272

273 Decoration proteins can be found on the top of the major capsid protein in a position crossing 274 $local$  two-fold axes somewhat similar to what was observed in  $\Phi$ RSL2. The shape of the 275 dimeric decoration proteins surrounding the hexamer is more globular in case of  $\Phi$ RSL2 276 compared to that of  $\Phi$ RP13. The  $\Phi$ RP13 decoration protein also appears to be hollow. On the 277 inside of the particle, in the middle of each hexamer, one can also find a globular extra density 278(Figure 3J, right, arrow). On the DNA level, it is possible to distinguish at least five 279 concentric layers of DNA separated by 26.3 Å. Like  $\Phi$ XacN1, an irradiation-sensitive inner 280body was not detected for this virus (data not shown).

281

282

## 283Discussion

284We determined the icosahedral capsid structures of three jumbo phages. Two of these phages 285 are representatives of known triangulation number groups (T=27 and T=28), whereas the third 286 one has a triangulation symmetry number that was not previously known to exist for caudal 287bacteriophages (T=21,1). Surprisingly,  $\Phi Xa$ cN1 has a smaller size compared to  $\Phi RSL2$  even 288though it has a higher triangulation number, but this may be due to the difference in size of 289 their respective MCPs (46 Da vs  $70 \text{ kDa}$ ). The distance between the centre of two adjacent 290hexamers is slightly higher for  $\Phi$ RSL2 compared to  $\Phi$ XacN1 and  $\Phi$ RP13 (120 Å vs 113 Å, 291 respectively). Different decoration proteins have been visualized on the outer portion of  $292\Phi$ RSL2 and within its inner area. In contrast,  $\Phi$ XacN1 has the most basic capsid of the three 293phages studied here as no decoration proteins were observed on the exterior of the capsid. 294This proves that these accessory proteins are not essential to ensure the solidity of 295bacteriophage capsids, even when faced with the enormous internal pressures required to 296 compact up to 400 kbp of DNA. For  $\Phi$ RSL2, because its MCP is much larger compared to the 297 other two phages, one cannot be completely sure if the dimer present at the periphery of the 298 hexamer is an extra protein or part of the MCP itself. However, it is likely that this dimer 299 consists of accessory proteins since the same kind of dimer is present in  $\Phi KZ$  and  $\Phi PBS1$  and 300 because the MCPs are much smaller in these phages. A new kind of phage decoration protein 301 that forms a hollow tube was also found in the  $\Phi$ RSL2 capsid. One would need to study this 302at higher resolution to determine what role this protein may have.

303

 $304\Phi RSL2$  carries only 224 kbp of DNA but it has the largest capsid.  $\Phi RSL2$  is also the only 305phage in this study that has an inner body that does not exhibit concentric organisation of 306DNA layers within its three-dimensional structure. According to the structure we determined, 307 large protein extensions are present on the inner part of the capsid and the DNA appears to be 308 connected to the lower part of the hollow tube present at the centre of each hexamer. All of 309this together suggests that there are different types of DNA organisation in the jumbo phage 310 world: one organised around an inner body with DNA connected to the capsid through 311 dedicated structures near the 5-fold axes of the capsid; and another one as classical toroidal 312structures.

313Two out of the three phages exhibits a "classical" DNA packing density of 0.49 and 0.39 314 whereas the third one has as very low one:  $\Phi$ RSL2 (0.29). This shows that the exceptions still 315 exist and also shows the value of continuing extensive structural studies on viruses.

317The ΦRP13 capsid protects 180 kbp of DNA: less than most jumbo phages but more than 318 classical bacteriophages.  $\Phi$ N3 DNA is 207 kbp for T=19,l and with a capsid diameter of 120 319nm (vertex to vertex) compared to 114 nm for  $\Phi$ RP13 [6]. Because the capsid dimensions and 320the DNA size are in the same range for  $\Phi$ N3 and  $\Phi$ RP13, and even if  $\Phi$ RP13 does not 321 technically meet the jumbo phage criteria ( $> 200$  kbp), one can say that  $\Phi$ RP13 belongs to a 322new "semi-jumbo" category.

323

324

## 325Conclusion

326We found a new triangulation number symmetry to add to the bacteriophage morphology 327 catalogue as well as a new kind of decoration protein. Many triangulation numbers are 328 observed in nature.  $\Phi$ RP13 exhibits a higher T number than  $\Phi$ N3 but with DNA size that is 329too small for jumbo phage classification. We propose that PRP13 belongs to a new group of 330semi-jumbo phages. It seems that the distinction between classical myophages and jumbo 331 phages is not clearly defined and that there is a continuum in the DNA sizes carried by *Myoviridae*. The diversity of accessory/decoration proteins is enormous. One can imagine that 332 333 during evolution each phage has dipped into a common well to create its own combination of 334 decorative/accessory proteins. Extensive study of these accessory proteins would help to 335 discover novel protein properties (e.g. recognition of ligand).

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## 339 **Data availability**

340Cryo-EM maps have been deposited in the Electron Microscopy Data Bank:



351T.K. produced and purified the three bacteriophages. O.C. isolated  $\Phi$ RP13 in Thailand. G.S 352and E.N. prepared cryo-EM grids. E.N. and G.S. collected cryo-EM data on a FEI Polara EM. 353E.N. performed cryo-EM image processing and cryo-EM 3D reconstructions with the help of 354GE and LFE. The manuscript was written by G.S., E.N. and T.Y. with input from all authors. 355G.S. was responsible for the conception and direction of the work, analysing and interpreting 356 data and revising the final drafts of the manuscript. All co-authors have reviewed and 357 approved of the manuscript before submission and agree to be accountable for all aspects of 358the work. This manuscript has been submitted solely to this journal and is not published, in 359press, or submitted elsewhere.

360

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#### 371

### **Conflicts of interest**  372

373The authors declare that there are no conflicts of interest. 374

375

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## **Figures legend:** 380

381

**Figure 1: Electron microscopy of bacteriophages RSL2, XacN1 and RP13.** 382 383

384A - **QRSL2** 

385Top: Negative staining image of  $\Phi$ RSL2. The black arrows indicate decoration of the phage 386tail with fibrillary structures.

387Bottom: Cryo electron microscopy image of the jumbo phage. The inner electron-dense body 388is highlighted by a rectangle or an arrow. The white arrow indicates some free DNA released 389from the bacteriophage.

390

 $391B - \Phi XacN1$ 

392Top : Negative staining image of  $\Phi Xa cN1$ . The arrows highlight an annular density 393 decorating the phage tail.

394Bottom : Cryo electron microscopy image of the jumbo phage. The arrow points to the same 395 structure as the one highlighted in negative staining.

396

397C – **PRP13** 

 $398$ Top : Negative staining image of  $\Phi$ RP13. The double arrow highlights the presence of a 399 double layered baseplate in the virus. The inset show the bacteriophage in a contracted state 400 with the inner tube of the tail sticking out.

401Bottom: Cryo-EM image of ΦRP13. The double arrow points to the double baseplate.

402

403The scale bar represents 100 nm.

404

405

**Figure 2: Three-dimensional reconstruction of the three jumbo phages obtained from** 406 **cryo-EM images.** 407

408

409Three-dimensional reconstruction of the three jumbo phages represented as isosurface on the 410top of the figure. One facet is highlighted by a black triangle. The structures are color coded 411 according to the radius of the particle as indicated in D.

412

A diagram showing the organization of the asymmetric units in one of the facet of the 413 414 icosahedron is drawn for each virus. The organization of the hexamers in this facet makes it 415 possible to determine the triangulation number that characterizes each of the bacteriophages. 416The different decoration proteins are also shown. The scale bars represent 20 nm.

417

418A –  $\Phi$ RSL2. The bacteriophage head has a triangulation number T=27 (h=3; k=3; T=h<sup>2</sup> + hk  $419 + k^2$ ).

#### 420

421B -  $\Phi$ XacN1. The bacteriophage head has a triangulation number T=28, *dextro* (h=4; k= 2;  $422T=h^2 + hk + k^2$ ).

423

424C -  $\Phi$ RP13. The bacteriophage head has a triangulation number T=21, *laevo* (h=4; k= 1;  $425T=h^2 + hk + k^2$ ).

426

 $427D - Color code used in A-C to color the capsids according to their radius (in nm).$ 

428

#### **Figure 3 : Central section and decoration protein** 429

430

#### **RSL2** 431

 $432A$  – Half of the central section of the  $\Phi$ RSL2 density map. The protein density is in white. 433Black arrows are highlighting densities on the inner part of the capsid. The white asterisk is 434 highlighting the cylindrical spike and the white arrow the dimer around the hexamers. The 435 scale bar represents 20 nm.

436

437B - Schematic view (left) and enlarged isosurface view (center) of the 5-fold axis showing 438that the vertex is surrounded by dimers. The vertex is also prominent compared to the rest of 439the capsid.

440

441C - Schematic view (left) and enlarged isosurface view (center) of one pseudo 6-fold axis 442 showing that the hexamers are surrounded by dimers and that there is a hollow cylindrical 443 density sticking out from its center. On the right, isosurface view of a hexamer seen from the 444 inside of the capsid. The arrow is pointing to the inner part of the spike present in the center 445 of the hexamer.

446

447D - Detailed view of the inside of the capsid along the 5-fold axis. The blue densities are 448 corresponding to the extra densities indicated in A by the arrows. They represent probably 449 nucleoprotein complexes involved in the DNA organization.

450

#### **XacN1** 451

452E - Half of the central section of the PhXacn1 density map showing that the capsid is roughly 453 smooth. Turret-like structures are only seen at the 5-fold axes, and some extra densities are 454also visible under the 5-fold axes (arrow). It is possible to distinguish at least 6 concentric 455 layers of DNA. The protein/DNA densities are in white. The scale bar represents 20 nm. 456

457F - Schematic view (left) and enlarged isosurface view (center) of the 5-fold axis showing the 458 vertex composition. The right panel is a side view of the vertex showing the extra protein 459 forming the turret-like structure.

460

461G - Schematic view (left) and enlarged isosurface view (center) of one pseudo 6-fold axis 462 showing that the capsid is quite smooth and probably only composed by the major capsid 463protein.

464

#### **RP13** 465

466H - Half of the central section of the  $\Phi$ RP13 density map showing that the capsid is decorated 467by dimers (black arrow). No extra-densities are visible neither on the outside nor on the inside 468of the 5-fold axes. It is possible to distinguish at least 5 concentric layers of DNA. The 469protein/DNA densities are in white. The scale bar represents 20 nm.

470

471I - Schematic view (left) and enlarged isosurface view (center) of the 5-fold axis showing that 472the vertex is surrounded by dimers. The protein forming the vertex has, at this resolution, the 473 same shape as the one forming the hexamers (the major capsid protein) (see J).

474

475J - Schematic view (left) and enlarged isosurface view (center) of one pseudo 6-fold axis 476 showing that the hexamer formed by the MCPs is surrounded by dimers. The right panel is 477 showing an isosurface view of a hexamer seen from the inside of the capsid. The arrow is 478 pointing to an extra density which is probably a non-identified protein.

479

480

2915

**Table 1 : Electron microscopy statistics for the image analysis of the three bacteriophages.**

# **Table 2 : Dimensions of the different parts of the three bacteriophages (capsid, tail and**

485DNA). The inner volume of the capsid has been measured using UCSF Chimera as described 486in [6].

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Figure 3 Neumann et al.





Supplemental Figure 2 Neumann et al.



Supplemental Figure 3 Neumann et al.



Table 1 Neumann et al.



**Table 2 Neumann et al.**