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Bacteriophage P22 Ejects All of its Internal Proteins Before its Genome

Yan Jin¹, Sophia M. Sdao², John A. Dover², Natalia B. Porcek^{2,3}, Charles M. Knobler¹, William M. Gelbart^{1,4,5}, and Kristin N. Parent²

¹Department of Chemistry and Biochemistry, University of California, Los Angeles, Los Angeles CA, 90095; USA

²Michigan State University, Department of Biochemistry and Molecular Biology, East Lansing MI, 48824; USA

³Michigan State University, Department of Microbiology and Molecular Genetics, East Lansing MI, 48824; USA

⁴Molecular Biology Institute, University of California, Los Angeles, Los Angeles CA, 90095; USA

⁵California NanoSystems Institute, University of California, Los Angeles, Los Angeles CA, 90095; USA

Abstract

Double-stranded DNA bacteriophages are highly pressurized, providing a force driving ejection of a significant fraction of the genome from its capsid. In P22-like *Podoviridae*, internal proteins ("E proteins") are packaged into the capsid along with the genome, and without them the virus is not infectious. However, little is known about how and when these proteins come out of the virus. We employed an *in vitro* osmotic suppression system with high-molecular-weight polyethylene glycol to study P22 E protein release. While slow ejection of the DNA can be triggered by lipopolysaccharide (LPS), the rate is significantly enhanced by the membrane protein OmpA from *Salmonella. In contrast*, E proteins are not ejected unless both OmpA and LPS are present and their ejection when OmpA is present is largely complete before any genome is ejected, suggesting that E proteins play a key role in the early stage of transferring P22 DNA into the host.

Keywords

bacteriophage infectivity; genome ejection; internal protein ejection; receptors; osmotic suppression

Correspondence to: William M. Gelbart; Kristin N. Parent.

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Introduction

The detailed mechanisms by which bacteriophages deliver their genomes into their bacterial hosts are not fully understood. In general, for tailed double-stranded (ds) DNA phages, proteins located in the phage tail first contact the surface of the bacterium and the phage diffuses along the surface until it finds a specific receptor. Upon binding to the receptor, the phage tail undergoes a series of conformational changes that result in release of the DNA and in its translocation from the capsid into the cytoplasm of the host (Poranen *et al.*, 2002; Bhardwaj *et al.*, 2014). In order to overcome the defense barriers of the bacteria – e.g., the outer and inner membranes along with the periplasmic space in between – bacteriophages use different strategies based on their tail morphology. Compared to the long-tailed *Myoviridae* (with contractile tails) and *Siphoviridae* (non-contractile tails) (Leiman *et al.*, 2012; Davidson *et al.*, 2012), *Podoviridae*, a family of bacteriophages with tails shorter than the width of the periplasm, cannot directly use their tails to penetrate both membranes. While studies of the particular podoviruses phi29 and T7 (González-Huici *et al.*, 2006; Molineux, 2001) have shed much light on their mechanism of infection; to date no generalized mechanism of infection for phages has been identified.

P22 is a member of *Podoviridae* that infects *Salmonella enterica*. It is a dsDNA phage with a 43.5 kbp genome that is packaged via a headful mechanism (Casjens *et al.*, 1988) into an icosahedral procapsid formed from assembly of the coat, scaffolding, and portal proteins (King *et al.*, 1976). There are three internal proteins (called "pilot, "ejection", or "E" proteins) packaged inside P22 procapsids, all incorporated by the scaffolding protein in the early stages of assembly: gp16, gp20 and gp7 ("gp" = gene product), each with 10-20 copies (Israel, 1977). A short tail "machine" is then connected at a unique five-fold vertex to complete the P22 virion. Several of these structural proteins have been identified in recent cryoEM investigations (Tang *et al.*, 2011; Lander *et al.*, 2009). Although density in these reconstructions had been ascribed to the E proteins (Chang *et al.*, 2006; Lander *et al.*, 2006), a crystal structure of the complete portal, as well as the most recent asymmetric reconstructions (Olia *et al.*, 2011; Tang *et al.*, 2011) show this to be incorrect. Accordingly, the location of the three E proteins within the mature virion remains unknown.

Along with their DNA, most phages eject proteins that have been packaged into the capsid, and this is generally essential for infection (Molineux *et al.*, 2013). Although the exact roles that each of these E proteins plays in infection have not been determined, some information is available. For example, cells infected with gp16-deficient phage (a mutant lacking a functional gp16 E protein) continue to divide normally and do not replicate the P22 genome, but co-infections show that gp16 can work *in trans* to complement a gp16-deficient phages do not induce the superinfection-exclusion response and gp16 is not part of the replication machinery (Israel *et al.*, 1972). Gp7 and gp20 often co-purify with gp16, and gp7 ejection does not occur when gp16 is absent (Israel, 1977). Indirect evidence supports a membrane-breaching role because purified gp16 disrupts dye-loaded lipid vesicles (Perez *et al.*, 2009). It has been proposed that the E proteins protect the DNA in the periplasm during infection (Israel, 1977), but their function and location post-infection has yet to be determined

experimentally. Taken together, these facts suggest that the E proteins, gp7, gp16, and gp20, may be linked to the efficiency and dynamics of DNA ejection across host membranes.

If the receptor is known and can be solubilized, phage ejection can be triggered *in vitro*, in which case the extent of DNA ejection can be controlled by the presence of an osmolyte in the surrounding buffer solution (Tzlil et al., 2003; Evilevitch et al., 2003; Castelnuovo and Evilevitch, 2007; Bauer et al., 2013). More explicitly, each concentration of high-molecularweight polyethylene glycol (PEG8000) in the host ("external") solution corresponds to a certain amount of water being drawn out of the phage capsid (from which PEG is excluded). As a result, the water inside is under tension, thereby producing a force resisting the ejection of DNA. Such osmotic suppression studies done with dsDNA phages like lambda (Evilevitch et al., 2003) and T5 (Leforestier et al., 2008), both members of Siphoviridae, have shown that the virus capsid is highly pressurized as a result of the electrostatic selfrepulsion and bending of the densely-packed, negatively-charged, semi-rigid DNA (Riemer and Bloomfield, 1978; Tzlil et al., 2003). In these in vitro experiments, in which pressure is the only driving force for DNA ejection, the length of DNA ejected can be tuned by the osmotic pressure difference between the outside and inside of the capsid (Evilevitch et al., 2003). In the present study we use osmotic suppression to examine for the first time the ejection of both the DNA and the E proteins from P22, which sheds light on ejection mechanisms in Podoviridae.

Results/Discussion

For P22, the O-antigen portion of lipopolysaccharide (LPS) located on the surface of the host *Salmonella enterica* serovar Typhimurium works as a primary receptor for infection (Iwashita and Kanegasaki, 1973). Further, it has been demonstrated (Andres *et al.*, 2010) that LPS can trigger a slow ejection of DNA from P22 *in vitro*, which makes it possible to study the process with the osmotic suppression technique.

Osmotic suppression experiments show release of DNA from P22 is inhibited at a pressure of 16.8 atm

Using LPS to trigger ejection, we determined the fraction of the DNA remaining unejected in the presence of an osmotic pressure by separating the capsids from the ejected DNA, which was degraded by DNase, and then recovering and analyzing the DNA remaining in the capsids as in the experiment of Evilevitch *et al.* (2005). Fig. 1A is a cartoon showing the ejection behavior of P22 in the presence of LPS under different osmotic pressures. Ejected DNA is digested into nucleotides, but the DNA that is kept inside the capsids by the PEG pressure is protected; this DNA is extracted (see Materials and Methods) and its length analyzed by gel electrophoresis. As a control, we have demonstrated that the isolated P22 genome can be degraded by DNAse in the presence of PEG8000, as is consistent with previously published data (Andres *et al.*, 2010).

The results are indicated by the open circles in Fig. 1B. A representative agarose gel showing unejected DNA from the capsid under different osmotic pressure conditions is included in the supplemental materials (Fig. S1); the length of DNA remaining in the capsid is seen to increase with increasing PEG concentration until ejection is completely

suppressed at 16.8 atm. Because of the jump from 10.8 to 16.8 atm in our osmotic pressure measurements, it is possible that complete suppression occurs anywhere between these two pressures. But it is clear from the variation of ejection fraction with pressure for the set of measured points that 16.8 atm is an upper bound (and very close) to the value at which the suppression is complete. This is significant because, as we shall see in the following sections, ejection of the E proteins is largely *complete* at 16.8 atm in the presence of OmpA. For all samples, more than 80% of the capsids were triggered open as was confirmed by plating experiments in which the number of remaining plaque-forming units (PFUs) was counted after treatment with receptor, as previously described (Parent *et al.*, 2014). Note that partially-ejected capsids, and even capsids triggered open, but for which DNA ejection was

completely suppressed (those bound by LPS in high [PEG]), lose their infectivity, i.e., are unable to form a plaque. As an additional control, we incubated P22 in increasing [PEG] in the absence of LPS and analyzed the samples by agarose gels, confirming that PEG alone does not trigger capsid opening (data not shown).

E protein ejection does not occur when P22 is treated with LPS alone

Using the osmotic-suppression method described above to control the extent of ejection for a series of phage samples, we centrifuged ³⁵S-labeled samples (see Materials and Methods) to separate capsids from ejected protein and DNA. We then analyzed by SDS-PAGE, autoradiography and densitometry the protein content in the pellet, i.e., those proteins still associated with the capsid/receptor macromolecular complex. Fig. 2 shows the results for pellet fractions corresponding to different osmotic pressures, with LPS used to trigger ejection. For all the pressures reported in Fig. 1B, we assayed a portion of each reaction and counted PFUs before and after treatment with LPS to confirm that more than 80% of virions were triggered for ejection (Fig. S2).

If any (or all) of the E proteins had been ejected, they would be present in the supernatant and consequently the bands representing those proteins would become less intense or even disappear, depending on the fraction ejected. As is evident, each lane has the same viral protein pattern with similar band intensities, independent of the presence of LPS and the amount of PEG. The same result was found in 10 independent trials. Similar results were also found when we performed a trypsin digestion in parallel samples to digest any protein that was ejected and hence no longer protected by the capsid (data not shown). Therefore, we can exclude the possibility that protein was ejected but pelleted along with the capsid. Trypsin digestion was also performed with purified E proteins, and they were completely and rapidly degraded (Carol Teschke, personal communication), indicating that these proteins are highly sensitive to protease when they are outside of the capsid. We can thus conclude that LPS alone did not trigger E protein ejection *in vitro*.

P22 ejection efficiency increases *in vitro* in the presence of outer membrane protein A (OmpA)

Studies by Seckler and colleagues have shown that while LPS from *Salmonella* can trigger P22 ejection *in vitro*, the kinetics are very slow (~ 5 hours) compared to the phage life cycle (~1 hour) (Andres *et al.*, 2010; 2012). Because of the close relationship between Sf6 and P22 (Casjens *et al.*, 2011), and the fact that both the outer membrane protein OmpA and

LPS are required for Sf6 ejection *in vitro* (Parent *et al.*, 2014), we examined if OmpA was able to enhance the rate of ejection in P22. As seen in Fig. 3A, as early as 10 min after incubation with both LPS and OmpA, around 75% of the virions in the sample have lost infectivity versus only a 30% infectivity loss upon incubation with LPS alone. Samples incubated with OmpA alone showed essentially no loss of infectivity, indicating that OmpA does not trigger ejection without LPS (Fig. S2).

We also performed the same experiment with two P22 hybrids (Leavitt *et al.*, 2013) where the head of the phage is comprised of P22 proteins but the tail needle tip is from another phage type. These hybrids are constructs where the tips were fused downstream of the helical core of the P22 tail needle (includes P22 residues 1-140). For one such hybrid, containing an Sf6 tail needle knob (Sf6 residues 132-282) fused to the P22 helical core, we obtained results (Fig. 3B) similar to P22. But for a mutant with a T4 "foldon" tail needle (Fig. 3C), neither LPS with OmpA nor LPS alone was able to trigger loss of infectivity. The "foldon" is a 25-residue trimer found at the C terminus of T4 fibritin (Tao *et al.*, 1997) and all of the foldon residues are added to the 140 amino acid P22 helical core. It appears then that for LPS and OmpA to function as receptors, the tail needle structure must be closely similar to that of P22, consistent with the earlier suggestion that the P22 tail needle is part of the trigger that determines DNA ejection (Leavitt, *et al.*, 2013).

P22 E proteins are ejected *in vitro* in the presence of both LPS and an outer membrane protein

Since OmpA enhanced P22 ejection kinetics and overall efficiency (Fig. 3A), we asked if P22 capsids would release the E proteins *in vitro* when both receptors were present. The pressure driving DNA ejection was determined again, but this time in the presence of both LPS and OmpA (Fig. 1B—filled squares). Within the precision of the measurements, the data obtained for fast ejection, i.e., with the two receptors, are identical to those for slow ejection, i.e., for LPS alone.

However, when both LPS and OmpA were present, the E protein behavior was different. Fig. 4 is an autoradiograph showing pellets from samples with both receptors at osmotic pressures from 0 to 4 atm. Comparing lanes with receptors (lane 1-4) and the lane without receptors (lane 5), we can clearly see that bands representing gp16 and gp20 disappear in the presence of the two receptors, independent of the osmotic pressure. Similar results were found in more than 10 independent trials, but in most of them the E protein bands for samples of virus with both receptors – while many times weaker than the band for pure virus – were still visible and measurable. From these 10 trials we obtain (see discussion below) an average intensity of the E protein band associated with the pelleted phage in the presence of both receptors, for each of increasing osmotic pressures. A test with trypsin digestion, but without separation by centrifugation, gave consistent results for the autoradiograph – confirming that the E proteins were rapidly digested when the capsids were triggered "open" with both LPS and OmpA, indicating their release into the solution (data not shown). As above, all samples were analyzed by measuring PFU, and a large fraction of the phages (>90%) were triggered for ejection (Fig. S2).

The percentages of ejection of all three E proteins against osmotic pressures ranging from 0 to 38 atm were calculated as follows: For each autoradiograph, there is one lane involving receptor-free virus at a concentration identical to that of the other samples in the gel. The intensity of each E protein band for each sample is determined relative to the coat protein band from the same sample (see Materials and Methods). We also compared E protein intensity to that of portal and tailspike protein bands in the same lanes. In all instances, the ratios were consistent (data not shown). These results are presented in Fig. 5: 60 - 70% of each of the three E proteins is ejected from the capsid, independent of osmotic pressures ranging from 0 to 16.8 atm, whereas no E protein can be detected at higher pressures (Fig. S3).

In comparison with the incomplete (\approx 70%) ejection of E proteins that we have observed, *in vivo* studies of P22 E protein ejection performed in the early 1970s showed that more than 90% of gp16 and gp20, as well as 70-90% of gp7 were ejected from the capsid during infection. In our system LPS and OmpA are present as free molecules in solution while *in vivo* the LPS and OmpA are both in the outer membrane of the bacterium and closely situated. In contrast, in our *in vitro* solution studies it is possible that virus particles interact with LPS without simultaneously being in contact with OmpA, thereby triggering DNA ejection only and leaving E proteins inside the capsid. An alternative explanation is that perhaps OmpA is not as functional when purified in detergent micelles as it is *in vivo*. However, our data show very clearly that virtually no E protein release occurs with LPS alone, whereas near-physiological amounts of protein ejection occurs when OmpA is also present.

If we compare the DNA partial ejection data (Fig. 1B) with the E protein ejection data (Fig. 5), we see that while at 16.8 atm DNA ejection is completely suppressed, most of the E proteins are still ejected. From this we conclude that in the presence of LPS and OmpA all three E proteins are ejected prior to DNA ejection. It is only at an osmotic pressure 2 atm higher that the ejection of the E proteins is inhibited. However, we were not able to determine the *order* of E protein ejection. It is possible that all the proteins bind together and are ejected together, or that they are ejected in a random sequence that is not detectable in our bulk population measurements. Even if an order exists, it is also likely that, because all the proteins are ejected before the highly stressed DNA is ejected, their order cannot be resolved using the osmotic suppression method because the contribution of each protein to the overall pressure is too small.

The force driving ejection of the protein from P22 arises from the confined DNA and is expected to be similar to that associated with the ejection of DNA from λ phage, which like P22 has a T=7 capsid and is about the same size. Grayson *et al.* (2006) carried out osmotic suppression measurements on λ for the 48.5 kb wild-type genome and a 37.7 kb mutant and found that the ejection was completely inhibited at pressures of 20-25 and 10-15 atm, respectively. The pressure required to inhibit the E proteins that we have observed for P22 originates from 43.5kb-length of DNA (longer than the 41.7 kb genome because of head-full packaging) confined to a volume slightly smaller than the capsid interior because of the presence of the E proteins. The fraction of the internal volume occupied by the E proteins can be estimated from their number and molecular masses, using a typical protein density of

1.4g/cm³. Taking the internal volume to be 1.1×10^5 nm³ (Patterson *et al.*, 2012), one estimates that the E proteins occupy about 2% of the volume. This decrease in volume available to the DNA is quite small. Nevertheless because of the exponential dependence of osmotic pressure of highly condensed DNA (Rau *et al.*, 1984 and Tzlil *et al.*, 2003) it would be expected to increase the ejection pressure by about 10%, comparable to the difference we observe between the pressures (16.8atm and 19.2atm, respectively) that inhibit the ejection of the DNA and proteins. Finally we note that the turgor pressure in both the periplasm and cytoplasm of *Salmonella* is about 3.5 atm (Stock *et al.*, 1977), so that pressure-driven ejection from P22 can be expected to transfer only about 60% of the DNA from the phage to the host (see Fig. 1B); another mechanism is necessary to account for the remaining 40%, and this remains to be clarified.

Conclusions

Productive virus infection requires accurate recognition of the host cell to avoid wasted genome release. This process is largely controlled by specific interactions between the virus and the receptors from the host. It has been shown previously (Andres *et al.*, 2010; 2012) that LPS from the host *Salmonella* can trigger slow DNA ejection of P22 *in vitro*. In our present work, we have demonstrated for the first time that, together with LPS, purified OmpA can: (1) dramatically increase the rate and the efficiency of DNA ejection of P22 *in vitro*; and (2) facilitate the *in vitro* ejection of P22's internal (E) proteins. OmpA is one of the major outer membrane proteins in bacteria, with ~100,000 copies in each cell (Koebnik *et al.*, 2000). Our results suggest that it can serve as a potential secondary receptor during P22 infection. This finding is similar to the case of Sf6 (another *Podovirus* closely related to P22) for which LPS from the host *Shigella* cannot trigger DNA release by itself. Only when a secondary receptor like OmpA or OmpC is also present can the virus trigger DNA release (Parent *et al.*, 2014). It remains to be determined how LPS alone or LPS and OmpA interact with the portal complex. Perhaps interaction with both receptors present induces a conformational change that promotes effective ejection of both DNA and E proteins.

Another open question, especially for short-tailed *Podoviridae*, is the following: after specific binding to the outer membrane, how is viral DNA translocated across both membranes of the host to initiate the infection? Compared to some phages whose long tails are in principle long enough to span the space between the outer surface and the cytoplasm of their host, *Podoviridae* may require a more complex mechanism to deliver their genome. One of the best studied *Podoviridae*, T7, likely utilizes its internal proteins to further extend its tail across the membranes to facilitate genome delivery as seen in cryo tomograms of T7 infecting mini cells (Garcia, 1995; Hu *et al.*, 2013; Molineux *et al.*, 2013). A recent example of a non-tailed phage, phiX174, has shown that this icosahedral ssDNA phage can use an internal protein to build a structure required for DNA translocation. These results suggest that even in the absence of a tail machine, a protein tube is still formed by internal phage proteins (Sun *et al.*, 2014). So far there is no such structural evidence for P22. Our finding that in *in vitro* experiments with OmpA that all E proteins are ejected *before* the DNA, and the fact that all of them are essential for infectivity, along with the finding that gp16 can work *in trans* (Hoffman *et al.*, 1975; Israel; 1977), suggest that the these proteins may play a

key role *in vivo* in transferring P22 DNA through the periplasm and inner membrane by either protecting the DNA from periplasmic nuclease digestion or disrupting the membrane.

We have shown that the ejection of the proteins precedes that of the DNA but we have been unable to determine if they are ejected in a specific order. The question might be resolved if there were information about the location of the proteins in the capsid, or about their specific functions after ejection. One possibility for elucidating more about these mechanisms is to determine by cryo-EM high-resolution structures during ejection, i.e., distinguishing "before" and "after" conformations. More generally, it will be interesting to compare and contrast these issues for short- vs long- tailed phages and to unravel the roles of primary and secondary receptors in the genome ejection and delivery processes.

Materials and Methods

Plaque assay for determining the virus ejection rate

For all experiments, the P22 strain used was a clear plaque mutant that is an obligate lytic strain (Casjens *et al.*, 1987) and was purified from *Salmonella enterica* serovar Typhimurium DB7136 (leuA414am, hisC525am, su⁰) LT2 cell lysates. *S. enterica* serovar Typhimurium strain DB7136 (Winston *et al.*, 1979) was used for all phage plaque assays. Hybrid P22 with either an Sf6 tail needle knob or a T4 foldon tail were induced from prophage, and were a gift from Sherwood Casjens (University of Utah). LPS was purified using a kit (Bulldog Bio), and OmpA was purified as described (Parent *et al.*, 2014; Porcek and Parent, 2015). Each type of virus with final concentration of 10^9 PFU/ml was incubated with LPS (0.25 mg/ml), or OmpA (0.2 mg/ml), or both LPS and OmpA, all in the presence of detergent Triton X-100 [1.06% (w/v)]. All plating experiments were performed in phage dilution buffer (10 mM Tris, 10 mM MgCl₂, pH = 7.6). At each time point (0, 10, 20, 30, 60 min post mixing), an aliquot was removed for titering. The percentage of infectious virus remaining was determined using the PFU from each sample divided by that for a sample containing no receptor.

Genome ejection assay

P22 labeled with radioactive ³⁵S was purified as described in (Parent *et al.*, 2004). The virus sample (10^{10} PFU/ml) was first treated with DNase I (New England Biolabs) to remove any free DNA. Samples of P22 and LPS in phage dilution buffer were mixed with PEG 8000 (at concentrations of PEG corresponding to osmotic pressures of 0, 1.78, 6.2, 10.8 or 16.8 atm, respectively) (Stanley and Strey, 2003) and were incubated overnight at 37°C to ensure maximum genome ejection. The ratios between virus and the receptors were the same as those employed in the plaque assays. For samples containing both LPS and OmpA as receptors, the same concentrations of P22, LPS, OmpA, Triton X-100 and PEG used in the plaque assays were mixed and incubated at 37°C for 1 hr. In both cases, DNase I (5 units) was added to the sample after ejection was triggered, and the mixture was kept at 37°C for 4 hr to digest the DNA ejected from the capsid. The procedure for determining the amount of the genome remaining in the capsids after partial ejection is essentially that described by Evilevitch *et al.* (2005). Before recovering the unejected DNA from the capsid, 1mM EDTA was added and the sample heated for 10 min at 75°C to denature the DNase I. After addition

of an equal volume of lysis buffer (25mM EDTA, 200mM Tris, 250mM NaCl, 1%SDS, pH 7.5) and 1 unit of Proteinase K, the samples were incubated at 65°C for 1 hr to disrupt the capsids, allowing the DNA inside to be released. Phenol / chloroform extraction was carried out twice to separate protein from DNA, followed by ethanol precipitation to concentrate the DNA. The DNA was then resuspended in TE buffer and analyzed by gel electrophoresis. In order to resolve the relatively long DNA, a low-percentage agarose gel (0.3%) was used and the running condition was 6 hr at 3V/cm; SYBR gold was used to stain the gel in the last step (Fig. S1).

SDS-PAGE for determining E protein ejection

We used samples prepared as described above for the plaque assays. Samples with LPS as the only receptor were incubated overnight to ensure maximum ejection. DNase I was then added followed by another 2 hr incubation at 37° C to completely digest the ejected DNA. An aliquot of each sample was used to determine the efficiency of genome ejection. The remainder of the sample was centrifuged at $27,000 \times g$ for 1.5 hr to separate the virus capsids from possible ejected proteins and free nucleotides. Resuspended pellets containing capsids were TCA-precipitated and loaded into a 10% SDS-PAGE gel. The gel was then fixed, dried and exposed to HyBlot CL autoradiography Xray film (Denville). Gel densitometry on the developed film was performed with a BIORAD Gel Doc XR+ documentation system.

The ejection fraction (*f*) for each E protein is calculated from the intensity (*I*) of that band, corrected by the receptor efficiency (α) found from the plaque assay, divided by that of pure virus sample (I_0). More explicitly:

$$f = \frac{I_0 - I}{aI_0}$$

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations

LPS	lipopolysaccharide
PFU	plaque forming units
PEG	polyethylene glycol
dsDNA	double-stranded DNA

ssDNA single-stranded DNA

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Highlights

- P22 *in vitro* genome ejection was monitored by osmotic suppression measurements
- Ejection proteins do not exit the capsid when triggered with LPS only
- Both LPS and an outer membrane protein were needed for Ejection protein release
- Ejection proteins, gp7, gp16, and gp20 are ejected prior to DNA release



Figure 1. The extent of genome ejection is controlled by osmotic pressure

A. cartoon of P22 *in vitro* ejection. The purified phage is incubated in PEG8000 and DNase. Receptor is then added, triggering ejection of progressively less DNA as the osmotic pressure increases; ejected DNA is digested by the DNase, and the protected DNA that remains in the capsid is extracted and its length analyzed by gel electrophoresis. B. Measured DNA ejection percentage from P22 at various osmotic pressures under different receptor conditions: P22+LPS (○); P22+LPS+OmpA (■). Ejection is triggered by addition of receptor (LPS, or LPS and OmpA) in the presence of PEG and DNase I; the DNase is

inactivated; and the DNA remaining in the capsids is extracted, run on an agarose gel, and the unejected length calculated from an accompanying DNA ladder. % DNA ejection is relative to the full-length DNA. The solid curve is drawn to aid the eye.



Figure 2. LPS alone is insufficient to eject E proteins

SDS-PAGE visualized by autoradiography showing the protein content in the pellet after triggering ejection by LPS under different external osmotic pressures. The left-most lane is the control which does not contain any receptor. Boxes highlight the three E proteins.



Figure 3. Loss of infectivity is enhanced by OmpA

In vitro genome ejection of (A) wild type P22, and of two P22 hybrids, one with an Sf6 tail (B) and the other with a T4 "foldon" tail (C). The "fraction of infectious virions remaining" was calculated at each time point as the number of PFU remaining after incubation with LPS, OmpA or LPS and OmpA, divided by the number of PFU when incubated with buffer at t = 0 min.



gp1, portal protein

gp 9, tailspike protein gp16, E protein

gp 5, coat protein gp 20, E protein

Figure 4. E proteins are ejected in the presence of both LPS and OmpA

SDS-PAGE gel visualized by autoradiography showing the protein content in the pellet after ejection was triggered by LPS and OmpA at different external osmotic pressures. The right-most lane is the control which does not contain any receptor. This is a representative film that most clearly shows the release of gp 16 and gp 20.



Figure 5. E proteins are ejected in the presence of LPS + OmpA, independent of osmotic pressures up to 16.8 atm

In vitro E protein ejection fractions at different osmotic pressures were calculated from the intensity of E protein band in each sample divided by the intensity of that band in the absence of receptors. Each value is an average of at least five experiments and the error bars represent one standard deviation. The arrow is to emphasize the pressure at which it was first observed that, after triggered ejection by LPS and OmpA, all of the DNA remained inside the capsid but the majority of the E proteins ejected.