Penetration of Membrane-Containing Double-Stranded-DNA Bacteriophage PM2 into Pseudoalteromonas Hosts

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The icosahedral bacteriophage PM2 has a circular double-stranded DNA (dsDNA) genome and an internal lipid membrane. It is the only representative of the Corticoviridae family. How the circular supercoiled genome residing inside the viral membrane is translocated into the gram-negative marine Pseudoalteromonas host has been an intriguing question. Here we demonstrate that after binding of the virus to an abundant cell surface receptor, the protein coat is most probably dissociated. During the infection process, the host cell outer membrane becomes transiently permeable to lipophilic gramicidin D molecules proposing fusion with the viral membrane. One of the components of the internal viral lipid core particle is the integral membrane protein P7, with maturalytic activity that apparently aids the process of peptidoglycan penetration. Entry of the virion also causes a limited depolarization of the cytoplasmic membrane. These phenomena differ considerably from those observed in the entry process of bacteriophage PRD1, a dsDNA virus, which uses its internal membrane to make a cell-envelope-penetrating tubular structure.

The key step in the life cycle of most viruses is the successful interaction with an appropriate host via a receptor on the cell surface. Several obstacles must be overcome for successful entry of a virus into the host cell. In the case of gram-negative bacteria, the envelope contains three layers: two membranes and a rigid peptidoglycan layer. In addition, nucleases in the periplasmic space form a challenge to bacteriophage entry. Binding to a specific outer membrane (OM) receptor has been well described for several phages (44, 45), but the mechanism of DNA entry into the bacterial cell is still a poorly understood process.

Bacteriophages use diverse strategies for genome entry (for reviews, see references 30, 59, 60, and 77). Tailed double-stranded-DNA (dsDNA) bacteriophages of Escherichia coli, such as T4, T5, T7, and lambda, have stable icosahedral capsids with a proteinaceous tail associated with the portal vertex. This unique vertex is used for genome packaging and delivery into the host bacterium. Receptor binding by the tail leads to structural rearrangements, and then the linear DNA is injected with the aid of the tail. The empty capsid is left on the cell surface. Mechanisms of penetration vary among tailed phages: the genomes of T5 and T7 are transported across the envelope through a protein-rich channel partially formed by phage proteins (35, 59, 67, 68), whereas the channel for the transfer of the lambda genome is formed by host-derived proteins (11, 79). Translocation of the T7 genome occurs simultaneously with its transcription by the host and T7 polymerases (36, 94), while phage T4 induces fusion between the OM and cytoplasmic membrane (CM) (88, 89). However, in the case of bacterial viruses that contain a membrane and no tail, the lipid component is used for cell entry. For example, the enveloped dsDNA phage φ6 fuses its external membrane with the host OM (5, 76) and the dsDNA phage PRD1 uses its internal membrane to form a tubular structure that penetrates the cell envelope (4, 38, 61).

Transport of viral DNA across the host cell membranes is a highly efficient energy-dependent process. It has been demonstrated that some phages utilize the proton motive force (Δp), consisting of membrane voltage (ΔΨ) and pH difference across the CM (ΔpH) for DNA delivery. The injection of phage T4 DNA into the host is strictly dependent on the metabolic state of the cell (50). A minimal CM voltage of ~90 mV is needed for the proper delivery of the genome (56). However, other tailed phages such as T5 and lambda can transfer their genomes across the envelope of deenergized cells (57, 79). Changes in the CM permeability and the depolarization of the membrane indicate that the genomes of phages T4, T5, and lambda are injected through a transient channel formed in the CM (14, 15, 41). Surprisingly, during the penetration of phage PRD1 DNA, the host CM is not depolarized but the membrane voltage increases (27).

Measurements of ion gradients across the cell envelope during bacteriophage infection have been used to study DNA entry processes (27, 59, 60). The OM of gram-negative bacteria forms a relatively permeable barrier allowing small hydrophilic metabolites and inorganic ions to pass through porins (72). However, the highly charged lipopolysaccharide (LPS) and proteins in the outer leaflet of the OM make a permeability barrier to lipophilic compounds (73). The bacterial CM is a highly selective barrier. It is permeable to lipophilic ions like tetraphenylphosphonium (TPP+) and ionophoric antibiotics like gramicidin D (GD), while inorganic ions like K+ do not cross the CM. The efflux of intracellular potassium is used as an indicator of increased CM permeability (for details, see references 27 and 59). Measuring the distribution of the lipophilic membrane voltage indicator TPP+ between cells and
Medium or binding of phenyldecaborane (PCB–) to the cellular membranes has been used to study the energy state of the CM and the permeability of the OM. The ion fluxes across the cell envelope can be observed by using selective electrodes that monitor the concentrations of indicator ions in the media.

Bacteriophage PM2 (31) is a lipid-containing dsDNA virus that infects two marine pseudoalteromonads, *Pseudoalteromonas espejana* BAL-31 and the virus were isolated from coastal seawater in Chile (32). Another host, ER72M2, was isolated from the East River, New York City, N.Y.. Based on its ribosomal 16S sequence, ER72M2 also belongs to the pseudoalteromonads (9). Later, the genus *Alteromonas* was divided into *Pseudoalteromonas* and *Alteromonas* (37).

PM2 is the only known isolate of the *Corticoviridae* family (1, 7). The virus (~65 nm in diameter) consists of an icosahedral capsid surrounding a protein-rich membrane vesicle, the lipid core (LC), which encloses the circular dsDNA genome of 10,079 bp (43, 48, 54, 62). The highly supercoiled phage genome replicates in association with the host CM, using the rolling-circle mechanism initiated by the virally encoded replication initiation protein P12 (18, 62). Among its 21 putative genes, ten (I to X) have been shown to encode structural components, proteins P1 to P10 (48, 53, 54, 62). All structural proteins, except P9, are encoded by the late operon, which is regulated by two phage-encoded transcription factors, P13 and P14 (63). For transcription, PM2 uses the host RNA polymerase (95). Virus-derived endolysin activity induced during PM2 infection has been described (90).

The mass of the virion is distributed among protein (72%), lipid (14%), and nucleic acid (14%) (22, 23, 24). Cryo-electron microscopy-based three-dimensional image analysis of native PM2 showed that trimers of the major capsid protein P2 are organized on an icosahedral pseudo-T = 21 lattice (48). There are pentameric spike complexes composed of protein P1 at the fivefold vertices. The proximal N-terminal domains of P1 are connected to the P2 capsid, and the distal C-terminal domains are exposed to the medium (54). The PM2 virion is a metastable structure stabilized by Ca2+ ions. In the absence of calcium, the coat dissociates revealing the spherical hydrophobic LC particle (54). LC-associated proteins (P3 to P10) occupy about one-third of the PM2 membrane, and the rest are lipids derived from the host CM during virus maturation (33, 43). The capsid and the underlying LC are linked by 60 transmembrane anchors near the twofold symmetry axes (48).

The process of entry of the viral genome into its host in general, and into a gram-negative cell in particular, is poorly understood. The internal membrane component in the PM2 virion suggests its involvement in this process. Currently, there is no information describing the early stages in the PM2 life cycle. In this investigation, we characterized the adsorption parameters and measured cellular energy changes during PM2 entry. We also identified a virion protein with murolytic activity that potentially is involved in locally digesting the peptidoglycan during virus entry.

### MATERIALS AND METHODS

**Bacterial strains, virus, and plasmids.** Bacterial strains and plasmids used in this study are listed in Tables 1 and 2, respectively. Phage PM2 (31) and *Pseudoalteromonas* cells were cultured in SB broth (53) or in a marine defined rich medium (63, 70) at 28°C. *E. coli* strains were propagated in Luria-Bertani broth (LB) at 37°C. When HB101 was used in adsorption tests, LB medium was supplemented with 10 mM CaCl2 to maintain phage viability.

**Transfection of *Pseudoalteromonas* cells by phage DNA.** PM2 DNA was isolated as described by Mannisto et al. (62) and transferred into *Pseudoalteromonas#

### TABLE 1. Bacteria used in this study

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*a Plasmid pDMI has a p15A replicon encodes resistance to kanamycin as the selective marker, and contains the gene for the lac repressor.

### TABLE 2. Plasmids used in this study

<table>
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*a PM2 nucleotide coordinates are according to Mannisto et al. (62) (GenBank no. AF155037).
cells by transfection according to van der Schans et al. (91). Transfected spheroplasts were grown for 2 h followed by plating on a lawn of PM2-sensitive host cells.

**Isolation of phase-resistant mutants.** For the isolation of spontaneous PM2-resistant ER72M2 cells, confluent plates were prepared. After overnight growth at 28°C, PM2-resistant plaques were picked, and five subsequent single-colony purifications were carried out. The sensitivity of the resulting strains to phage was tested, and the total protein compositions were analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) to detect possible contaminating species.

**Growth and purification of the phage.** Radioactive labeling of the PM2 virion was carried out in marine defined rich medium (see above) not including methionine. PM2 was labeled either with l-[^35]Smethionine (11 μCi/ml; Amersham, St-Jovite) 10 min after infection or with [3P] (11 μCi/ml; Amersham BF1003) 30 min prior to infection of a BAL-31 culture (7.5 × 10^7 CFU/ml), using a multiplicity of infection (MOI) of 10. The virus particles from the lysate were concentrated with polyethylene glycol 6000 and purified by rate zonal centrifugation (53, 54). A similar amount of virus to ER72M2 cells was also analyzed by rate zonal centrifugation. After a 15-min incubation the cells were layered on a linear 5 to 17% sucrose gradient (SB buffer; Sorvall TH641 rotor; 24,000 rpm, 60 min, 15°C). The two supernatant fractions were combined (soluble fraction), and the pellet (insoluble fraction) was resuspended in deionized water. After overnight storage at 4°C, the fractions were used in the adsorption test as described above. In previous studies an increase of adsorption of these fractions was observed with exponentially growing ER72M2 cells and incubated for 15 min at 28°C. The adsorption was determined after a 10-min adsorption as described above.

**Adsorption in the presence of isolated cell fractions.** Twenty milliliters of exponentially growing ER72M2, BAL-31, and HB101 cells (2–2 × 10^8 CFU/ml) was collected and resuspended in 2 ml of 10 mM Tris-HCI, pH 8.0, containing 1 mM EDTA. Cells were disrupted by sonication on ice (four 5-s pulses), and intact cells were removed by centrifugation (Heraeus BioBiofe; 10,000 rpm, 30 s, 4°C). The broken cells were retained by centrifugation (Heraeus BioBiofe; 13,000 rpm, 10 min, 4°C), and the resulting pellet was washed with the above-mentioned buffer and centrifuged again. Two supernatant fractions were combined and centrifuged. The supernatant fractions were combined (soluble fraction), and the pellet (insoluble fraction) was resuspended in deionized water. After overnight storage at 4°C, the fractions were used in the adsorption test as described above. In previous studies an increase of adsorption of these fractions was observed with exponentially growing ER72M2 cells and incubated for 15 min at 28°C. The adsorption was determined after a 10-min adsorption as described above.

**Adsorption to protease-treated cells.** ER72M2 and HB101 cells were grown to a density of ~5 × 10^8 CFU/ml. HB101 cultures were transferred to 28°C, and tetracycline was added to both cultures at a final concentration of 100 μg/ml to prevent protein synthesis. After 30 min, pronase, trypsin, and proteinase K were added to reach final concentrations of 2 mg/ml, 100 μg/ml, and 400 μg/ml, respectively, followed by incubation at 28°C for 10, 30, or 60 min. The cells were washed twice to remove the proteases and resuspended in SB medium containing 1 μg/ml of tetracycline. Treated cells were incubated for 15 min at 28°C before usage in the adsorption test as described above (10-min adsorption time). Cell densities and viabilities were determined by plating the cells on SB or LB plates. All proteases were purchased from Boehringer Mannheim.

**Electron microscopy.** Bacteria were grown to a cell density of ~7 × 10^8 CFU/ml and infected with fresh virus stock to obtain an MOI of 160. Samples were taken at 2, 5, and 10 min after infection, and the cells were fixed with 3% (vol/vol) glutaraldehyde for 20 min at 22°C. Fixed cells were collected by centrifugation, washed twice, and prepared for transmission electron microscopy as previously described (3). The measurements were taken using a JEOL 1200 EX electron microscope operating at 60 kV (Electron Microscopy Unit, Institute of Biotechnology, University of Helsinki).

**Measurements of ion fluxes and determination of membrane voltage.** Ion flux measurements were performed as described by Daugelavicius et al. (27). Briefly, cells were grown to a density of 7 × 10^8 CFU/ml, centrifuged, concentrated 200-fold, and dissolved in 50 mM Tris-HCI, pH 7.4, containing 50 mM NaCl. After a 15-min incubation with protease, an aliquot of cells was collected and resuspended in deionized water at 4°C for 4 h. The water and phenol phases were separated by centrifugation. The resulting phenol layer was extracted with another 5 ml of water. The two water phases were combined and dialyzed against deionized water at 4°C for five nights. After dialysis, the mixture (~10 ml) was concentrated threefold under vacuum, stored overnight at 4°C, and used in the adsorption test as described above. Increasing amounts of the isolated LPS fraction were mixed with a constant number of exponentially growing ER72M2 cells and incubated for 15 min at 28°C with aeration. After this, the cells were infected with PM2 (MOI ~0.4). The amount of nonadsorbed phages was determined after a 10-min adsorption as described above.

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bated as described for the ion flux measurements. Fifty microliters of the cell suspension was withdrawn and mixed with 750 μl of 100 mM Tris-acetate (pH 7.75), 2 mM EDTA buffer, and 200 μl of ATP Monitoring Reagent (BioOrbit). The amount of light produced was measured by using a model 1250 Luminometer (LKB-Wallace). The total ATP content of the cells was measured by adding 200 μl of ATP Releasing Reagent (BioOrbit) to the bacterial suspension. The amount of ATP present in the incubation buffer was calculated from a calibration curve. The intracellular ATP concentration of the cells was calculated from the external concentrations, assuming that 1 × 1010 cells correspond to an intracellular water volume of 5 μl (see above).

Expression and purification of PM2 proteins P1 and P2. The E. coli DH5α(pDMM)(pRM206) strain was used for expression of protein P1 (see Tables 1 and 2). Standard molecular cloning techniques were used (82). The cells were grown overnight in LB supplemented with 150 μg of ampicillin/ml and 25 μg of kanamycin/ml. The culture was diluted ~30-fold in the same medium and grown to a density of 2 × 10^8 CFU/ml at 28°C. Expression was induced by adding isopropyl-β-D-thiogalactopyranoside to a final concentration of 1 mM, and the culture was transferred to 18°C. After 16 to 18 h, the cells were collected (Sorvall SLA3000 rotor; 5,000 rpm, 10 min, 5°C), resuspended in 20 mM Tris-HCl (pH 8.0) buffer containing 5% (vol/vol) glycerol (1/100 of the original volume), and stored at −80°C. The cells were disrupted by two passages through a precooled French pressure cell (~105 MPa, 35-ml cell), and the cell debris was removed by centrifugation (Sorvall SS34 rotor; 12,000 rpm, 20 min, 5°C). The supernatant was cleared by centrifugation (Beckman Ti 50 rotor; 40,000 rpm, 1 h and 40 min, 5°C). Protein P1 was precipitated from the supernatant by 35% (vol/vol) (NH₄)₂SO₄, resuspended in 20 mM Tris-HCl, pH 8.0, containing 150 mM NaCl, and purified by gel filtration (Superdex 200 16/60 [Amersham Pharmacia Biotech] equilibrated with 20 mM Tris-HCl [pH 8.0]–150 mM NaCl). The peak fractions containing protein P1 were pooled and used for the adsorption inhibition tests. It was estimated from SDS-PAGE that this fraction contained ~80% P1.

For the expression of protein P2, the E. coli HMS174(DE3)(pRM607) strain was used (see Tables 1 and 2). The cells were grown in LB supplemented with 25 μg of kanamycin/ml and stored and disrupted as described above for protein P1 except that the pH of the resuspension buffer was 7.5. Protein P2 was precipitated with 42.5% (vol/vol) (NH₄)₂SO₄, resuspended in 20 mM Tris-HCl, pH 7.5, containing 150 mM NaCl, and purified by size exclusion chromatography (Superdex 200 26/60 [Amersham Pharmacia Biotech] equilibrated with 20 mM Tris-HCl [pH 7.5]–150 mM NaCl). The fractions containing P2 were pooled, dialyzed against 20 mM Tris-HCl (pH 7.5) containing 30 mM NaCl (Spectra/Por membrane; cutoff, 12,104 kDa), and loaded onto 1-ml Hitrap Q Sepharose columns (Amersham Pharmacia Biotech). Flowthrough containing protein P2 was collected, stored at 4°C, and used in the adsorption inhibition tests. With this technique, P2 was purified to near homogeneity.

The multimericity of recombinant proteins was determined by gel filtration as described above and by sedimentation in a linear 10 to 40% (wt/vol) sucrose gradient in 20 mM Tris-HCl, pH 7.5, containing 150 mM NaCl buffer (Sorvall T84 rotor; 42,000 rpm, 15°C). Protein P1 was 23.75 kDa (90.6 kDa) trimers isolated from PM2 virus particles (54), protein P3 (120 KDa) and P5 (85 kDa) trimers isolated from PRD1 virus particles (4), bovine serum albumin (68 kDa; Sigma), and lysozyme (14 kDa; Boehringer Mannheim) were used as molecular mass markers. Gradient fractions were analyzed by SDS-PAGE.

Adsorption tests and ion flux measurements in the presence of protein P1 or P2. A constant number of ER72M2 cells (grown to ~7 × 10⁸ CFU/ml) was mixed with increasing amounts of purified P1 in equal volumes of SB broth (taking into account that the P1 specimen contained ~20% impurities). After 15 min of incubation at 28°C with shaking, the cells were infected with the fresh virus agar stock (MOI, ~0.3). After 10 min, adsorption was stopped and the number of nonadsorbed phages was determined as described for the phage adsorption test.

For the measurements of ion fluxes, ER72M2 cells in 50 mM Tris-HCl, pH 8.0, containing 450 mM NaCl and 10 mM CaCl₂, were combined with a mixture of P1 and P2 proteins isolated from phage particles and purified recombinant P1 (see above) or purified recombinant P2 (see above). The cells were incubated for 5 min with protein prior to the addition of phages. The measurements were carried out as described above.

Lytic activity assays by zymogram analysis. The production of 2X purified PM2 virions and isolated lipid cores from freeze-thawed phage particles for zymogram assays were carried out as described previously (54). For the expression of PM2 proteins P3, P4, P5, P6, P7, and P8, the corresponding coding sequences were amplified by PCR and inserted into expression vectors (see Table 2). Strains DH5α(pDMM)(pRM203), HMS174(DE3)(pRM313), HMS174(DE3) (pRM304), HMS174(DE3)(pRM315), HMS174(DE3)(pRM316), and HMS174(DE3)(pRM325) were used for the expression of proteins P3, P4, P5, P6, P7, and P8, respectively. DH5α(pDMM)(pDS12) and HMS174(DE3)(pJJ2) were used as negative controls. Cells carrying different plasmids were grown in LB medium supplemented with appropriate selective antibiotics (ampicillin, 150 μg/ml; kanamycin, 25 μg/ml), and protein expression was carried out as described for protein P1. After expression, cells were collected and concentrated tenfold by resuspension in 20 mM Tris-HCl (pH 7.2) buffer. The cells were disrupted by sonication, and the soluble and insoluble fractions were separated (Heraeus Biofree; 10,000 rpm, 4 min, 4°C). Bacteriophage PRD1 particles used as a control in zymogram assays were purified as previously described (8). All samples were stored at −20°C until used in zymogram analysis.

The peptidoglycan succus of E. coli DH5α was isolated as described by Hoeve and Beveridge (47) by boiling 500 ml of stationary-phase cells in 4% SDS followed by removal of the peptidoglycan-associated proteins with 2 M NaCl (10). The isolated peptidoglycan preparation was resuspended in 5 ml of water and stored at −20°C until used. Zymogram analysis was performed as previously described by Bernardsky et al. (10). SDS–16% polyacrylamide gels (101 by 82 by 0.5 mm) were cast as described by Olkkonen and Bambord (75) except that the peptidoglycan preparation homogenized by sonication was added at a final concentration of 7.5% (vol/vol) in the separation gel. After electrophoresis, gels first were soaked in deionized water and then in renaturation buffer (25 mM KPO₄ [pH 7.2], 0.1% [vol/vol] Triton X-100) for 30 min. Following this, zymogram gels were transferred into fresh renaturation buffer and the incubation was continued for 40 to 60 h. After renaturation, gels were rinsed with water, stained with 0.1% methylene blue in 0.01% KOH for 1 h at 28°C, and destained with deionized water.

Analytical methods. Protein concentration was measured by using the Coomassie blue method with bovine serum albumin as a standard (17). Proteins were separated by SDS-PAGE using either 14% polyacrylamide–tricine–SDS gels (53, 85) or 16% polyacrylamide–SDS gels (75). For amino-terminal amino acid sequencing, the proteins were transferred from the gel to a polyvinylidene difluoride membrane (Millipore) and stained with Coomassie blue (PhastGel Blue R; Pharmacia Biotech). Sequencing was performed by using a Procise 494A protein sequencer (Perkin-Elmer/Applied Biosystems, Foster City, Calif.) at the Protein Chemistry Laboratory, Institute of Biotechnology, University of Helsinki.

RESULTS

Large amounts of phage particles bind to the host cells, but no empty capsids are seen on the cell surface. When a number of Pseudalderomonas species were screened for PM2 sensitivity using the plaque assay, only one new isolate, ER72M2, in addition to BAL-31 cells, was able to plate PM2, indicating that the receptor is not a common Pseudalderomonas surface structure (53). Also note that Pseudalderomonas sp. strain A28 was unable to plate PM2 (see below). The adsorption of the phage to different cells was studied under vigorous aeration in normal growth medium resembling seawater. The adsorption was strictly aeration dependent. If the cells were not aerated, no virus binding was detected. The plating efficiency of PM2 on BAL-31 and ER72M2 is the same, but the binding of virus to the cell surface differed considerably (Fig. 1A and B). The adsorption rate constants calculated for the two hosts were 1.4 × 10⁻¹⁰ ml/min and 2.2 × 10⁻¹⁰ ml/min for ER72M2 and BAL-31, respectively (Fig. 1A). Virus adsorption to the sensitive strains was compared to that to PM2-resistant control cells (Pseudalderomonas sp. strain A28 or E. coli HB101). In addition, 50 independent spontaneous PM2-resistant ER72M2 cell lines were isolated (see Materials and Methods). The electrochemical characteristics of the resistant mutants were the same as those of the sensitive ER72M2 strain, indicating that the envelopes of the resistant cells were not altered. All the tested cell lines showed no binding of the virus above the background level in the adsorption test (Fig. 1A), and they served as isogenic controls for the measurements of PM2-specific effects in the electrochemical measurements. The nonspecific binding of the virus to resistant E. coli (Fig. 1A) did not deviate from virus
decay during the assay, which was demonstrated with virus material in the absence of cells.

In one-step growth experiments (MOI, 10) the PM2-infected bacterial culture lyses effectively (53). Ten minutes postinfection (p.i.) ~95% of cells were infected (MOI, 10). With an increase in the MOI, the fraction of infected cells increased (99% at an MOI of 140). To determine how many virus particles bind per CFU, receptor saturation experiments were carried out. The maximum number of bound particles was ~180 per 1 ER72M2 CFU, achieved at an MOI of ~300 (Fig. 1B). BAL-31 bound over 400 particles per CFU at the highest MOI used (850) with no signs of saturation. Thin-section electron microscopy of BAL-31 and ER72M2 cells during the first 10 min of infection (Fig. 1C to E) showed only a few DNA-containing virus particles associated with the cell surface when an MOI of 160 was used. Counts of visible virus particles on the cell surface revealed approximately 20 DNA-containing particles per 100 cell sections. The cell-associated particles had a mean diameter of ~40 nm, while PM2 has a ~51-nm diameter in thin-section electron microscopy (Fig. 1C to E) (54). In addition, no empty virus capsids were detected in association with the cells.

Binding inhibition assays were used to search for the PM2 receptor. Isolated LPSs from sensitive cells had no effect on PM2 adsorption (data not shown). Neither did the addition of the soluble or insoluble fractions from disrupted sensitive cells. In addition, extensive protease treatment of the host cells to digest the surface proteins did not alter the capacity of the cells to adsorb phages (data not shown).

Release of virion components follows phage adsorption. Incorporation of L-[35S]methionine to capsid and LC-associated proteins was studied after separation of the labeled virion proteins in a polyacrylamide gel. About 60% of the radioactivity associated with the capsid (proteins P1 and P2), and the rest (40%) associated with the LC proteins. Extraction of the nucleic acid and lipids from the 33P-labeled virions showed that ~87% of the radioactivity was associated with nucleic acid and the rest (13%) resided in the lipid fraction.

Association of the labeled viruses to ER72M2 cells was examined by rate zonal centrifugation (Fig. 2). This allowed the radioactivity associated with the nonadsorbed particles to be distinguished from that released or bound to host cells. The mixing of L-[35S]methionine-labeled PM2 (MOI, 10) with ER72M2 cells resulted in the formation of two separate fractions. Sixty percent of the radioactivity was found free at the top of the gradient, and 40% associated with the cells (Fig. 2A). In these conditions ~75% of PM2 particles were bound, as detected by the adsorption assay or reduction of the virus peak (Fig. 2A). An autoradiogram of the supernatant and cell-associated fractions of the adsorption mixture separated by tricine-SDS-PAGE revealed that the major coat protein P2 was found in the soluble fraction (inset in Fig. 2A). Due to the low specific activity obtained, we could not demonstrate the association of the LC proteins with the cells. Sedimentation analysis of the 33P-labeled PM2 mixed with ER72M2 cells (MOI, 10) showed that ~35% of the radioactivity associated with the cells while the majority (65%) was released (Fig. 2B). In this experiment ~80% of the particles associated with the cells (see above). An autoradiogram of the supernatant and

FIG. 1. Phage PM2 adsorption. (A) Time-dependent adsorption to BAL-31 (black circles), ER72M2 (black triangles), A28 (open circles), and E. coli HB101 (open triangles) cells. The amount of nonadsorbed phages was measured after different times p.i. Fifty PM2-resistant ER72M2 cell lines were tested using a 10-min adsorption time, and the results did not deviate from those obtained for E. coli, as indicated by a grey square. (B) The number of PM2 particles adsorbed to ER72M2 (open circles) and BAL-31 (black circles) cells at different MOIs (15-min adsorption time). (C to E) Thin sections of PM2-infected cells (MOI, 160). (C) Two DNA-containing virus particles in association with ER72M2 cells after 10 min of infection are denoted by arrowheads. Bar, 200 nm. (D) A higher magnification of a free virion and one adsorbed DNA-containing particle on an ER72M2 cell after 2 min of infection. (E) A cell-associated DNA-containing particle on a BAL-31 cell after 2 min of infection. Bar in panel E, 50 nm for panels D and E.
cell-associated fraction demonstrated that the viral DNA resided in both fractions (Fig. 2B).

Receptorless *Pseudoalteromonas* sp. strain A28 supports virus reproduction. *Pseudoalteromonas* sp. strain A28 harbors a plasmid, pAS28 (52), which shares significant sequence similarity with the early operon of PM2 (62, 63). The adsorption test revealed that PM2 could not bind to the A28 cell surface (Fig. 1A). Further, analysis of the effects of PM2 on A28 cells, as determined by measuring ion fluxes, revealed that the phage had no effect on envelope permeability of A28 cells (Fig. 3). Interestingly, transfection of purified PM2 genomes into A28 spheroplasts produced plaques if the transfected cells were plated on a lawn of PM2-sensitive cells. This shows that A28, a close relative to the two *Pseudoalteromonas* hosts, does not have the receptor for PM2, but after the genome has entered the cell, viruses are produced. No other tested pseudoalteromonads or *Alteromonas macleodii* (Table 1) was able to produce virus in this assay, except the host cells.

The outer membrane of marine pseudoalteromonads is highly permeable. Pseudoalteromonads are gram-negative cells. The OMs of the best-studied gram-negative bacteria, *E. coli* and *Salmonella enterica*, form a permeability barrier to lipophilic compounds such as TPP+ or ionophoric antibiotics (for a review, see reference 73). However, the OM permeability of pseudoalteromonads to TPP+ was rather high (Fig. 3A). Ac-
cumulation of TPP⁺ differed between the studied strains. BAL-31 and A28 accumulated considerably higher amounts of TPP⁺ compared to ER72M2, indicating a higher membrane voltage (ΔΨ). However, higher OM permeability to lipophilic compounds in the case of BAL-31 and A28 also is possible, as they accumulated a higher amount of PCB⁻ and were more sensitive to GD compared to the ER72M2 strain (Fig. 3B).

In spite of the high OM permeability of the studied three Pseudoalteromonas strains, the CM-depolarizing and intracellular K⁺-releasing effect of GD was low. In addition, the depolarizing effect of the polycationic antibiotic PMB was rather low on ER72M2, but somewhat higher in the case of BAL-31 (data not shown). However, the combination of GD and PMB induced very effective depolarization of the CM and an efficient release of intracellular K⁺ (Fig. 3 and 4). The presence of TPP⁺ or PCB⁻ ions had no effect on the viability of the cells or the amount of infective centers observed when BAL-31 and ER72M2 were infected with an MOI of 10.

EDTA is known to chelate divalent cations away from the LPS layer, increase the OM permeability to lipophilic compounds, and induce the additional accumulation of lipophilic cations in gram-negative cells (71). However, in the case of Pseudoalteromonas strains, the addition of up to 500 μM EDTA to the cell suspension (in 50 mM Tris-HCl [pH 8.0] containing 450 mM NaCl) caused only partial release of accumulated TPP⁺ and K⁺ (data not shown).

**PM2 induces depolarization of the CM.** Phage-induced effects on the CM can be detected by measuring the efflux of the membrane voltage indicator TPP⁺ ion from the infected cells. The release of TPP⁺ from infected BAL-31 and ER72M2 cells began 20 to 22 s after addition of phage particles and continued for approximately 2 min, indicating depolarization of the CM (Fig. 3A). The phage-induced leakage of accumulated TPP⁺ was MOI dose dependent (Fig. 4A). Approximately five-times-higher numbers of infectious particles induced effects with similar amplitudes on BAL-31 cells when compared to ER72M2 cells. Four-times-higher numbers of superinfecting phages, added 6 min after the initial infection, had no effect on the external concentrations of the three ions studied (Fig. 3, ER72M2 strain).

Since the combination of GD and PMB very effectively depolarized the CM, it was possible to calculate the membrane voltage. The ΔΨ of ER72M2 and BAL-31 cells before infection was ~165 and ~187 mV, respectively (calculated from Fig. 3A). After infection, before GD addition, ΔΨ was ~153 mV for ER72M2 and ~177 mV for BAL-31. At a high MOI (150), the CMs of ER72M2 cells were depolarized from ~167 to ~142 mV in 5 min (Fig. 4A). After infection, the membrane voltage stayed at the lower level and repolarization of the CM was not detected. The accuracy of the calculations depends on viable count measurements. Light microscopy of growing bacteria revealed that both strains did not separate the dividing cells, and the cells were seen predominantly as doublets. When this is taken into account, the effective ΔΨ is ~18 mV lower than the calculated one.

**PM2 induces strong binding of PCB⁻ to cell membranes and K⁺ leakage.** In addition to depolarization of the CM, PM2 induced high accumulation of PCB⁻ by infected BAL-31 and ER72M2 cells (Fig. 3B). The decrease of PCB⁻ concentration in the medium started 12 to 15 s after the addition of phages...
and continued for approximately 1.5 min. The amplitude of the phage-induced effect on the accumulation of PCB \(^{−}\) increased with the MOI (Fig. 4B). PM2 virus particles bound rather low amounts of PCB \(^{−}\). However, at high MOIs, the binding of PCB \(^{−}\) to the viral membranes can be observed immediately after the phage addition as a fast uptake of this anion.

Cells initially accumulated potassium ions, but after approximately 1 min of incubation, a slow spontaneous leakage of intracellular K\(^{+}\) started (Fig. 3C). GD and PMB caused elimination of the K\(^{+}\) gradient on the CM, allowing us to calculate the intracellular K\(^{+}\) concentration during infection (calculated from Fig. 3C). The maximal calculated intracellular K\(^{+}\) concentration (just before the spontaneous leakage starts) was \(~2.0\) and \(~2.3\) M for ER72M2 and BAL-31, respectively. However, when the inaccuracy in viable count measurements (see \(\Delta\psi\) calculations) is taken into account, the effective intracellular K\(^{+}\) concentration is about half of the calculated one. The addition of infectious PM2 particles stimulated the efflux of K\(^{+}\) ions starting \(~15\) s after infection and lasting for \(~2\) min (Fig. 3C). The amplitude of this efflux was proportional to the MOI used (Fig. 4C). After this K\(^{+}\) efflux stimulation, the rate of K\(^{+}\) leakage from infected ER72M2 and BAL-31 cells was close to the rate of spontaneous leakage, which was also observed for A28 cells (Fig. 3C). The calculated intracellular potassium ion concentrations of ER72M2 and BAL-31 during the first 2 min of infection decreased from \(~1.7\) to \(~1.5\) M and from \(~2.0\) to \(~1.75\) M, respectively (see also above).

**Additional energy considerations.** In order to estimate the role of cell energetics in infection, the effects of energy transformation-modifying compounds were studied. The addition of glucose did not induce additional accumulation of either TPP \(^{−}\) or K\(^{+}\) ions, but some release of the accumulated TPP \(^{−}\) was registered. The addition of glycerol, lactose, or sucrose to the incubation medium had no effect on the accumulation of TPP \(^{−}\) or K\(^{+}\). NaN\(_3\) eliminates bacterial growth by blocking respiration and uncoupling oxidative phosphorylation due to inhibition of cytochrome oxidase and membrane H\(^{+}\)-ATPase (42). In the presence of 20 mM NaN\(_3\), the amount of accumulated TPP \(^{−}\) was somewhat lower but the cells were able to maintain membrane voltage and keep the K\(^{+}\) gradient. In these conditions, the amplitude of PM2-induced TPP \(^{−}\) and especially PCB \(^{−}\) and K\(^{+}\) fluxes was even higher than without NaN\(_3\) (data not shown).

The measurements of ATP-dependent light emission by the luciferin-luciferase system showed that the calculated intracellular concentration of ATP was rather high, \(~8\) mM. However, the effective concentration was \(~4\) mM (see \(\Delta\psi\) calculations). The concentration of intracellular ATP in ER72M2 cells did not decrease during infection (calculated concentration 8.3 mM at 3.5 min p.i. and 9.3 mM at 10 min p.i.).

**Purified PM2 protein P1 inhibits phage adsorption.** The outer layer of the PM2 virion consists of two major structural proteins, P1 and P2. To evaluate whether these capsid proteins could interfere with the interaction between the phages and the cells, the proteins were isolated from viral particles (see reference 54) and added to the adsorption mixture. Clear adsorption inhibition was observed (data not shown). To test which of the proteins was responsible for the effect, they were separately expressed from plasmids and purified (Fig. 5A for P1; for P2, data not shown). The recovery of purified proteins P1 and P2 was \(~1\) and \(~4\) mg/liter of culture, respectively. N-terminal amino acid sequence analysis of the purified proteins gave sequences MVKKKLA and MRSLNLNSI, equivalent to the P1 and P2 sequences, respectively, as deduced from the virus genome. In addition to the information obtained during purification by size exclusion chromatography (P1, \(~40\) kDa, and P2, \(~65\) kDa), the multimericity of isolated proteins was determined by rate zonal centrifugation (P1, \(~55\) kDa, and P2, \(~85\) kDa). Recombinant P1 and P2 behaved as the corresponding proteins from the virus particle (54), indicating that the recombinant P1 is a monomer and P2 is a trimer in solution.

The ability of these proteins to interfere with PM2 adsorption was analyzed by two different methods. In the adsorption test, the presence of increasing amounts of P1 blocked adsorption (Fig. 5B). In the presence of P1, phage-induced effects on the envelope permeability of ER72M2 cells with respect to...
PM2 has a temporal effect on OM permeability. The addition of GD at different time points before and after the addition of PM2 particles in the presence of calcium (Fig. 6) revealed that cells were sensitive to GD effects (PCB- binding and K+ leakage) only during the first 2 min of infection, but not after that time or before infection. This indicates the presence of a temporal virus-induced patch on the OM, which is permeable to lipophilic GD molecules.

In the absence of calcium, ER72M2 cells had considerably higher membrane voltage (~212 mV; see above). Under these conditions, PM2 alone did not cause depolarization of the CM (~209 mV), PCB- accumulation, or K+ leakage (Fig. 7A to C). However, GD added 3 min or later after addition of the phage particles induced strong depolarization of the CM, leakage of K+, and accumulation of PCB-, indicating that the transient nature of OM permeability was abolished (Fig. 7A to C). The amplitude of phage PM2-induced binding of PCB- depended on the concentration of CaCl₂ at the beginning of infection (Fig. 7D). The maximal effect was achieved with 10 to 15 mM CaCl₂. Higher concentrations of Ca²⁺ had deleterious effects on infection.

Small structural PM2 protein P7 has muralytic activity. The lytic activities associated with bacteriophage PM2 particles were examined by analyzing highly purified virions by zymogram analysis. After separation of viral proteins by SDS-PAGE (16% polyacrylamide–SDS gel containing peptidoglycan), proteins were allowed to renature at different temperatures (4, 15, 28, and 37°C). After renaturation at 4 or 15°C, one clear PM2-derived zone was detected (Fig. 8). The molecular mass of the PM2 structural protein with lytic activity was below 17 kDa, which is the mass of the PRD1 lytic enzyme P15 (Fig. 8).

The internal LC particle contains proteins P3 to P10 (48, 53). Isolated LC particles were analyzed in zymograms, and the protein with muralytic activity was present in this preparation (data not shown). In order to determine which protein was responsible for this activity, the structural proteins P3, P4, P5, P6, P7, and P8 were expressed in E. coli (see Table 2) at two different temperatures, 18 and 37°C. After expression, the cells were disrupted and fractionated into soluble and insoluble fractions. Expression of proteins P3, P5, P6, and P7 could be detected in cell extracts with Coomassie blue staining. P5 was detected in the soluble fraction (18°C). P3 (37°C), P6 (18°C and 37°C), and P7 (18°C and 37°C) were found in the insoluble fraction. Expression of proteins P4 (4.4 kDa) and P8 (7.3 kDa) could not be detected by this method. The produced proteins were identified by N-terminal amino acid sequence analysis. The obtained sequences, MNTSVPTS, MKKAHMF, ANFLT KNF, and MINKTTIK, were identical to the sequences deduced from the genes for PM2 proteins P3, P5, P6, and P7, respectively. The first methionine of recombinant protein P6 is removed after translation, which is also the case with the virus-derived protein.

To analyze whether the cell extracts containing recombinant PM2 proteins could produce peptidoglycan-hydrolyzing activity similar to that observed in viral specimens, zymogram analysis was performed. Cell extracts of the bacteria carrying the cloning vectors without inserts were used as negative controls to reveal the cell-derived lytic activities. Screening of the cell extracts in zymograms analyses using different renaturation temperatures showed that protein P7 (expressed at 37°C) had similar enzymatic activity as that observed in PM2 virions and in the LC particles (Fig. 8). According to these data, we conclude that protein P7 is a lytic enzyme. It is the smallest characterized structural protein (3.7 kDa) in the PM2 virion, containing one putative membrane-spanning helix (53, 62). However, a data bank search of the PM2 proteins did not reveal any matches to known sequences for lytic activities (PROSITE) (34).

DISCUSSION

Only two Pseudoalteromonas isolates carrying a high number of receptors on the cell surface have been shown to act as hosts for PM2 (Fig. 1) (53). Pseudoalteromonas sp. strain A28 harboring a plasmid (pAS28) (52), but no receptor, was shown to support PM2 growth after purified phage genomes were transfected into the cell. This finding is consistent with the observation that the replication and regulatory regions of the PM2 genome and plasmid pAS28 share significant similarity (63). In this study, we showed that an abundant receptor molecule is nonproteinaceous or is a protein that is very resistant to proteolysis. Extracts of sensitive cells did not interfere with the virus, indicating that the receptor or receptor complex is functional only on the cell surface, not being extractable, as is that of phage PRD1 (28).

In the native virion, P1 monomers are grouped as pentamers at the fivefold vertices by the N-terminal domain, which con-
nects the complex to the capsid formed from P2 trimers (48). Previously, it has been demonstrated that the lack of the C-terminal portion of P1 results in an inability of the virus to infect the host (54). We showed that the monomeric protein P1 readily associated with the cellular receptor, interfering with virus adsorption (Fig. 5). This demonstrates that the initial interaction between the host and PM2 is mediated by P1, which is known to build up the spike structure of the virion. The trimeric viral coat protein P2 did not interfere with binding of PM2 to its primary receptor.

The adsorption experiments (Fig. 1A and B) and thin-section electron microscopy (Fig. 1C to E) of infected cells gave contradicting results, as only very few DNA-containing particles were seen to be in contact with cells in spite of the high adsorption rate and high MOI used. Interestingly, the diameter of the cell-associated particles corresponds to the LC, not to the virion (Fig. 1D). In addition, no empty particles were detected in association with the cells, even 10 min p.i., when all the DNA entry-derived electrochemical events were over (Fig. 3). Moreover, the mixing of PM2 with its host cell leads to the dissociation of the particle (Fig. 2). Our previous results have demonstrated that the capsid is a metastable structure easily dissociated by calcium depletion, freezing and thawing of the particles, or by storage (53, 54). In contrast to these results, the empty viral capsid stays associated with the cell in the case of numerous dsDNA phages, including internal membrane-containing bacteriophages PRD1 and Bam35 (61, 78). The gross morphology of the PM2 virion resembles that of PRD1 (6, 39), which uses the internal membrane vesicle for DNA delivery by transforming it into a tube-like structure (38). No tubular membrane structures have been detected for PM2. Accordingly PM2 entry diverges considerably from those viruses.

The envelope permeability of _Pseudoalteromonas_ cells deviates from that determined for the other well-studied enteric bacteria _E. coli_ and _S. enterica_ (73). In contrast to enterobacteria, the OM of _pseudoalteromonads_ was permeable to TPP\(^{-}\) (Fig. 3). The intracellular concentration of solutes in bacteria has been shown to increase in conjunction with increasing external salt concentration (46, 49). High intracellular K\(^{+}\) concentrations observed in _pseudoalteromonads_ presumably are a consequence of the extracellular osmolarity in its natural environment, seawater (and in the growth medium).

The dependence of the depolarizing effect of external GD on PM2 addition and its temporal nature (Fig. 6) indicate virus-induced changes in the OM. We envision that the LC interacts with the OM, creating a window for GD entry into the periplasm, where it causes depolarization in the CM. Presumably, due to the diffusion of lipids and movement of LPS

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**FIG. 7.** Phage-induced effects on envelope permeability of ER72M2 cells with respect to TPP\(^{-}\) (A), PCB\(^{-}\) (B), and K\(^{+}\) (C) ions in the absence of CaCl\(_2\). The measurements were carried out as described for Fig. 2 except that the incubation medium contained 50 mM Tris-HCl (pH 8.0) and 450 mM NaCl. Curve 1 shows noninfected cells, and curve 2 shows cells infected with an MOI of 50. GD and PMB were added at final concentrations of 5 and 100 \(\mu\)g/ml, respectively. (D) Phage PM2-induced accumulation of PCB\(^{-}\) by ER72M2 cells in the presence of different CaCl\(_2\) concentrations as indicated. The measurements were carried out as described for Fig. 1. The cells were infected with an MOI of 50.
molecules on the plane of the membrane, this permeability window closes within a few minutes. If there is a fusion between the virus membrane and the host OM, then the supercoiled DNA and its associated proteins, as well as the internal surface of the LC, will face the peptidoglycan. The interaction between protein P4 and the PM2 DNA has been proposed, and it seems likely that P4 is associated with the viral membrane inside the vesicle (54, 64, 83, 84). However, after entering the cytoplasm, no genome-associated proteins are needed for proper initiation of virus replication (62, 91).

Many bacteriophages have proteins with lytic activity (55, 58). Furthermore, PM2-derived endolysin activity has been characterized (90). In addition to playing a role in host cell lysis, lytic enzymes are often structural virion components. These proteins locally digest the murein sacculus during penetration into the cell. Examples of such proteins are the two murolytic enzymes, P7 and P15 of PRD1 (21, 80, 81), gp16 of bacteriophage T7 (65), base-plate protein gp5 of T4 (51, 69), P5 of phage φ6 (20), and the recently described terminal protein gp3 of bacteriophage φ29 (66). Here, we demonstrated that a small LC-associated integral membrane protein, P7, harbors lytic activity (Fig. 8). It is conceivable that P7 locally opens the murein layer, allowing the entering DNA to reach the host CM.

PM2-induced effects on the CM (TPP+ and K+ leakage and PCB− binding) started some 15 to 20 s after phage addition and were MOI dependent (Fig. 3 and 4). These effects lasted approximately 2 min. There was only a moderate decrease in the membrane voltage during DNA penetration and no decrease in the cellular ATP concentration. DNA penetration through the CM obviously causes only a temporal destabilization in the membrane with no severe adverse effects on the cellular energy balance. Similar to the OM, there is a well-controlled pore in the CM for the penetration of DNA comprised of virus or host proteins or both. The PM2-induced relatively strong binding of PCB− to cellular membranes is an indication of pore formation in the bacterial envelope, as has been shown for bacteriophages T4 and lambda (14, 29, 41). This PM2-induced pore is permeable to potassium ions but not to ATP. After DNA passage, the pore is sealed, indicating a gating phenomenon.

Calcium controls the infection process and stability of PM2. The LC-associated proteins P6 and P10 and the gene product of open reading frame h contain a putative calcium-binding site (48, 54, 62). In the absence of calcium, the infection process seems to stop at the CM stage, since the OM was permeabilized but depolarization of the CM did not take place. Bacteriophage T5 DNA penetration through the CM also is controlled by calcium ions (13), pointing to a similar mechanism.

Combining these and previous results allowed us to delineate an entry pathway for PM2. It deviates considerably from previously described phage DNA delivery mechanisms. The binding of the virion to the cell surface receptor triggers a cascade of events where the viral coat is dissociated and the LC most probably fuses with the bacterial OM. This event is rapid (~2 min) but also occasionally leads to failure, as a fraction of the viral DNA is released. The murein-hydrolyzing enzyme in the phage membrane aids in peptidoglycan penetration. The penetration of the CM is a controlled gated event where a transient lowering of the membrane voltage is detected. Obviously, unique delivery and packaging mechanisms for the highly supercoiled circular PM2 genome have evolved.

ACKNOWLEDGMENTS

The Helsinki Graduate School in Biotechnology and the Molecular Biology and Viikki Graduate School in Biosciences are acknowledged for fellowships to H.M.K. and R.H.H., respectively. This work was supported by research grants 1202108 and 1202855 from the Academy of Finland (to D.H.B. and J.K.H.B.). R.D. is a Lithuanian State Fellowship holder. Pia Rydman is thanked for the guidance with the zymogram analysis.

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