Isolation and Characterization of a Generalized Transducing Phage for *Pseudomonas aeruginosa* Strains PAO1 and PA14

Jonathan M. Budzik,1 William A. Rosche,2 Arne Rietsch,3 and George A. O'Toole4*

Dartmouth College1 and Department of Microbiology and Immunology, Dartmouth Medical School,4 Hanover, New Hampshire 03755; Faculty of Biological Science, The University of Tulsa, Tulsa, Oklahoma 74104;2 and Department of Microbiology and Molecular Genetics, Harvard Medical School, Boston, Massachusetts 021153

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A temperate, type IV pilus-dependent, double-stranded DNA bacteriophage named DMS3 was isolated from a clinical strain of *Pseudomonas aeruginosa*. A clear-plaque variant of this bacteriophage was isolated. DMS3 is capable of mediating generalized transduction within and between *P. aeruginosa* strains PA14 and PAO1, thus providing a useful tool for the genetic analysis of *P. aeruginosa*.

*Pseudomonas aeruginosa* is an opportunistic human pathogen that infects individuals with weakened immune systems, such as hospitalized patients and those suffering from severe burns or other traumatic skin damage or from cystic fibrosis. *P. aeruginosa* is also a model organism for the study of bacterial biofilms (3). Two of the best-characterized strains of *P. aeruginosa* are PAO1 (7) and PA14 (11).

While bacteriophages are a useful tool for the genetic analysis of bacteria, we were not aware of any bacteriophage capable of mediating generalized transduction for strain PA14. Furthermore, bacteriophages with broader host ranges might also be capable of transductional gene exchange between strains (1, 8). We report the identification and characterization of a bacteriophage of *P. aeruginosa* that is capable of generalized transduction within and between *Pseudomonas* strains PAO1 and PA14.

**Isolation and initial characterization of phage DMS3.** Based on work with *Myxococcus* spp., we rationalized that we could trigger the lytic phase of a temperate bacteriophage present in *Pseudomonas* isolates by prolonged starvation (C. Manoil, personal communication). To test this hypothesis, ~70 clinical isolates identified as *P. aeruginosa* but otherwise uncharacterized were obtained from the Dartmouth Hitchcock Medical Center. The clinical isolates were grown in Luria-Bertani (LB) medium for 48 h at 37°C, the cells were subsequently pelleted by centrifugation at 17,000 × g for 5 min, and the supernatant was transferred to a clean tube. To kill the remaining bacteria, several drops of chloroform were added to the supernatant and the mixture was vortexed for 30 s. The resulting potential phage lysates were stored at 4°C.

A phage lysate derived from one of the *P. aeruginosa* clinical isolates was found to be capable of infecting *P. aeruginosa* strains PAO1 and PA14 and four *P. aeruginosa* clinical isolates, as demonstrated by the zones of lysis resulting from phages spotted onto bacterial lawns. Bacterial lawns were produced by adding ~5 × 10^9 CFU of *P. aeruginosa* from an overnight LB broth culture to 3 ml of LB top agar (0.8%) (12). The inoculated top agar was overlaid on an LB agar plate and allowed to solidify. Figure 1A shows the zones of lysis resulting from DMS3 spotted onto a lawn of PA14. Zones of lysis were also observed for DMS3 spotted onto lawns of PAO1 (Fig. 1D) and four clinical isolates (data not shown). The bacteriophage capable of inflicting these bacterial strains was plaque purified and designated DMS3. DMS3 has a stable titer (as measured by PFU) over a period of several years when stored at 4°C in the LB medium used to prepare the lysate.

Transmission electron microscopy (TEM) was performed to determine phage morphology. TEM grids (copper; size, 300 mesh) were coated with 0.4% Formvar and then floated on 10-μl spots of phage lysates followed by negative staining for 4 min using 20 μl of 2% phosphotungstate (pH 4.0). After being stained, the grids were rinsed with water and air dried. The grids were viewed using a transmission electron microscope (JEM 100CX EM; JEOL Ltd.) at ×500,000 magnification. TEM showed that DMS3 has a tail approximately 110 nm in length and an icosahedral head approximately 50 nm in diameter (Fig. 2).

The phage chromosome was isolated as reported previously (12) to determine the size and nature of the nucleic acid. After the experiment was begun with 0.5 ml of a phage lysate (~10^10 PFU/ml), sodium dodecyl sulfate was added to achieve a concentration of 0.5% and proteinase K was added to achieve a concentration of 0.05 mg/ml. The solution was heated for 30 min at 65°C followed by four successive extractions with a 1:1 phenol-chloroform solution. The nucleic acid was then ethanol precipitated. Phage nucleic acid was identified as DNA on the basis of its sensitivity to digestion by restriction endonucleases. The nucleic acid isolated from the phage could be cleaved by restriction enzymes DpnII and AluI (New England Biolabs, Beverly, Mass.), which exclusively digest double-stranded DNA (dsDNA), suggesting that the nucleic acid of DMS3 is dsDNA. To determine the size of the phage genome, pulsed-field gel electrophoresis (PFGE) was performed as reported previously (4). Isolated phage DNA was treated with S1 nuclelease (0 to 10 U) for 45 min at 37°C to linearize any circular DNA. Reactions were stopped with addition of cold EDTA (50 mM). Samples were electrophoresed in 1% PGFE-certified agarose (Bio-Rad) in 0.5× Tris-borate-EDTA. Electrophoresis was performed in a CHEF-DR II apparatus (Bio-Rad) at...
14°C with a linearly ramped switching time of 1 to 12 s. A constant voltage of 5 V/cm (150 V) was used for 20 h. After electrophoresis, the gel was stained with SYBR Gold (Molecular Probes) and visualized on a STORM 860 apparatus (Amersham). The genome size of the phage is estimated on the basis of PFGE results to be between 50 and 65 kb (Fig. 3).

As previously reported (9), DMS3 is susceptible to host-controlled modification, as evidenced by efficiency of plating. As shown in Table 1, the titer of a DMS3 lysate was lower on a strain other than the one on which it was originally grown, suggesting that the phage DNA might be modified to escape restriction. Host-controlled modification of the phage was observed when it was grown on \emph{P. aeruginosa} strain PAO1 or PA14. Although the basis of this host-controlled modification is unclear, for practical purposes these results suggest that phage grown on PA14 should be used to prepare lysates on strain PA14 and that phage grown on PA01 should be used to prepare lysates on strain PA01.

Type IV pili are the receptors for phage DMS3. A number of surface factors have been implicated as bacteriophage receptors. The ability of DMS3 to form plaques on a strain carrying a mutation in \emph{algC} (Fig. 1E), which is required for the synthesis of the lipopolysaccharide O-antigen and alginate, indicates that neither of these molecules serves as a receptor for this bacteriophage (2, 5). While DMS3 could form a zone of lysis on wild-type \emph{P. aeruginosa} PA14 (Fig. 1A), no such plaques formed on a \emph{pilA} or \emph{pilD} mutant (Fig. 1B and C). Both the PilA and PilD proteins are required for the synthesis of type IV pili. Therefore, DMS3 utilizes the type IV pilus as its receptor.

**FIG. 1.** Plaque formation by DMS3. Five microliters of a lysate of phage DMS3 was spotted in duplicate onto lawns of \emph{P. aeruginosa} wild-type and mutant strains. The zone of lysis (or clearing) on the plate indicates that DMS3 was able to infect the wild-type \emph{P. aeruginosa} PA14 strain (A). In contrast, the phage was incapable of producing a zone of lysis on \emph{P. aeruginosa} PA14 mutants defective in type IV pilus biogenesis (B and C). Both \emph{P. aeruginosa} PAO1 (D) and the \emph{P. aeruginosa} PAO1 \emph{algC} mutant (E) supported formation of zones of lysis.

**FIG. 2.** Electron micrograph of phage DMS3. A transmission electron micrograph of DMS3 negatively stained with phosphotungstate is shown.

**FIG. 3.** PFGE of phage DMS3 DNA treated with S1 nuclease. Isolated phage DNA was treated with S1 nuclease (lane 1, 0 U; lane 2, 0.5 U; lane 3, 1 U; lane 4, 10 U), which linearizes circular DNA. Lanes labeled M1 and M2 show size markers.
TABLE 1. DMS3 is susceptible to host-controlled modification

<table>
<thead>
<tr>
<th>Strain on which the phage lysate was prepared</th>
<th>Lawn on which phage titration was performed</th>
<th>Titer (PFU/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PA14</td>
<td>PA14</td>
<td>$5 \times 10^{11}$</td>
</tr>
<tr>
<td>PA14</td>
<td>PAO1</td>
<td>$3 \times 10^{6}$</td>
</tr>
<tr>
<td>PAO1</td>
<td>PAO1</td>
<td>$1.1 \times 10^{8}$</td>
</tr>
<tr>
<td>PAO1</td>
<td>PA14</td>
<td>$1.6 \times 10^{7}$</td>
</tr>
</tbody>
</table>

DMS3 mediates generalized transduction within and between *P. aeruginosa* strains. Phage lysates were prepared, as reported previously (8), by combining 100 µl of phage (~$1 \times 10^9$ PFU) with 100 µl of the donor bacterial strain (~$5 \times 10^8$ CFU from an overnight, LB-grown culture) and calcium chloride (10 mM). After a 15-min incubation at room temperature, the mixtures were combined with 3 ml of LB top agar (0.8%) and overlaid onto fresh LB plates. After ~14 to 16 h of incubation at 37°C, 4 ml of LB liquid medium was added to the LB top agar containing cells and phage and the top agar was scraped from the surface of the agar plate and transferred (along with the added LB liquid medium) to a 50-ml conical tube. An additional 3 ml of LB liquid medium was added to this mixture, which was then shaken slowly at room temperature for 4 h to allow release of the phage from the top agar. The lysates were centrifuged for 5 min at 17,000 × g to remove bacteria and agar, the supernatants were transferred to a new tube, and the remaining bacterial cells were lysed by the addition of several drops of chloroform. The chloroform-containing lysates were vortexed for 30 s and stored at 4°C. The resulting phage titers were typically ~$10^{10}$ to $10^{11}$ PFU/ml.

Transductions were performed as reported previously (8), with tetracycline or gentamicin resistance genes as the selectable markers carried by donor strains. Typically, 100 µl of an overnight LB-grown culture (~$5 \times 10^8$ CFU) was mixed with 100 µl of a phage lysate with a titer of $10^{10}$ to $10^{11}$ PFU/ml and calcium chloride was added to achieve a 10 mM concentration. After incubation for 15 min at 37°C, transduction mixtures were plated on LB agar medium supplemented with tetracycline (150 µg/ml) or gentamicin (100 µg/ml) as appropriate; transductants typically appeared after 48 h of incubation at 37°C. DMS3 mediated transduction within *P. aeruginosa* strains PAO1 and PA14 at an efficiency of ~$5 \times 10^{-9}$ transductants/PFU. Representative transduction frequencies for a variety of markers are shown in Table 2. The addition of EGTA (10 mM) or citrate (10 mM) or altering the multiplicity of infection did not improve transduction frequencies. We were unable to demonstrate that phage DMS3 was capable of mediating generalized transduction of markers with strain PAK. At least 15 different genetic markers from strains PA14 or PAO1 mapping across the chromosome, including alleles mapping to the *flgK*, *pilU* (a hyper-piliated strain), *sadB* (a biofilm mutant), *det-3* (a biofilm mutant), *lasR*, and *lasI* genes, were transduced using phage DMS3. Several of these transduction events were subsequently confirmed by PCR mapping and sequencing (10) or by phenotypic testing (e.g., flagellum-mediated swimming motility or biofilm development). These data demonstrate that DMS3 is capable of generalized transduction.

We also observed cross-strain transductions between strains PAO1 and PA14. The frequency of cross-strain transductions between strains PAO1 and PA14 was ~$2 \times 10^{-9}$ transductants/PFU. These frequencies were determined using three markers: *lasI*, *flgK*, and *sad-6* (a mutation in an uncharacterized gene required for motility; see Table 2). Cross-strain transductions were performed using fresh lysates prepared from single-plaque plaques propagated on the desired donor strain. Fresh lysates (with titers of ~$10^{11}$ PFU/ml) used the day they are prepared are essential for successful cross-strain transductions. The frequency of transduction events mediated by DMS3 appears to be at the borderline at which transductions can be detected; thus, it is helpful to perform multiple, replicate transductions.

Phage immunity tests of lysogens were performed as reported previously (9); the results of these tests suggested that DMS3 is a temperate (class 1) bacteriophage capable of lysogeny (6). This presents a problem in that transductants may be lysogens, which would prevent their use in subsequent transductions. Therefore, all transductants were assessed for phage-mediated immunity by the cross-streaking assay described previously (12), and only those transductants susceptible to phage infection were used for subsequent studies.

### Isolation of a clear-plaque variant of DMS3

A clear-plaque variant of DMS3 is useful in that it would form a more readily discernible plaque and allow for more-robust visualization of phage sensitivity in cross-streak assays. Phage DMS3 was mutagenized with UV (~2.5 J/m²), resulting in a ~100-fold reduction of PFU. The mutagenized phage was plated on *P. aeruginosa* PAO1, and a single clear-plaque isolate was identified among ~6,500 plaques screened with a dissecting microscope. This clear-plaque isolate is still capable of mediating generalized transduction.

### Summary

We have identified and characterized a generalized transducing phage that is capable of mediating transduction within and between strains of *P. aeruginosa*. Our data indicate that this temperate phage contains a dsDNA chromosome of ~50 to 65 kb, infects via type IV pili, and has an icosahedral phage-head morphology. A clear-plaque variant of this bacteriophage was isolated. DMS3 is a useful genetic tool for genetic studies of *P. aeruginosa* strains PAO1 and PA14.
We are grateful to J. Schwartzman for providing us with the *P. aeruginosa* clinical isolates and to C. Manoil for suggesting the approach of obtaining phage from bacterial isolates. We also thank Chuck Daghlian and Louisa Howard at the Rippell Electron Microscopy Facility at Dartmouth College for their assistance with electron microscopy.

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