Genomic and Transcriptional Mapping of PaMx41, Archetype of a New Lineage of Bacteriophages Infecting Pseudomonas aeruginosa

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ABSTRACT
Previously, a collection of virulent phages infecting Pseudomonas aeruginosa was isolated from open water reservoirs and residual waters. Here, we described the comparative genomics of a set of five related phages from the collection, the physical structure of the genome, the structural proteomics of the virion, and the transcriptional program of archetypal phage PaMx41. The phage genomes were closely associated with each other and with those of two other P. aeruginosa phages, 119X and PaP2, which were previously filed in the databases. Overall, the genomes were approximately 43 kb, harboring 53 conserved open reading frames (ORFs) and three short ORFs in indel regions and containing 45% GC content. The genome of PaMx41 was further characterized as a linear, terminally redundant DNA molecule. A total of 16 ORFs were associated with putative functions, including nucleic acid metabolism, morphogenesis, and lysis, and eight virion proteins were identified through mass spectrometry. However, the coding sequences without assigned functions represent 70% of the ORFs. The PaMx41 transcription program was organized in early, middle, and late expressed genomic modules, which correlated with regions containing functionally related genes. The high genomic conservation among these distantly isolated phages suggests that these viruses undergo selective pressure to remain unchanged. The 119X lineage represents a unique set of phages that corresponds to a novel phage group. The features recognized in the genomes and the broad host range of clinical strains suggest that these phages are candidates for therapy applications.

IMPORTANCE
Pseudomonas aeruginosa is an opportunistic pathogen that causes stubborn nosocomial infections that are frequently resistant to multiple antibiotics. Bacterial viruses (bacteriophages or phages) represent a natural mechanism for pathogenic bacterial control. Here, a group of virulent phages, previously shown to infect a broad range of clinical P. aeruginosa strains, was characterized at the genomic and molecular levels. These phages belong to a unique and tightly related group. In addition, we conducted a transcriptional study of an archetypal phage of this group to characterize the role of many unknown coding sequences based on expression temporarilites. These results contribute to our knowledge of 119X-like phages and, in general, provide information concerning P. aeruginosa podophage diversity and lytic cycles.

The renewed interest in the study of genetic diversity in phage populations has been prompted by the accessibility of massive DNA sequencing platforms, which has radically increased the number of phage genome sequences reported in the data banks (1). Presently, genome sequencing is an invaluable tool in phage characterization, which is fundamental at the first stages of the functional annotation of new coding sequences (2). Homologous sequences or structures in data banks provide accurate clues concerning the potential function of coding sequences. In addition, the genome sequence is the most important criterion used to understand relationships between phage groups (3). Pseudomonas aeruginosa is a Gram-negative bacterium that is globally distributed in a wide range of environments and is one of the most common opportunistic human pathogens (4, 5). Phages that infect this bacterium are frequently isolated and characterized because of their potential application as therapeutic tools (6). Pseudomonas-tailed phages are the most abundant group deposited in databases. Among these organisms, 38% are podoviruses, which show icosahedral heads and short tails (7). The International Committee on Taxonomy of Viruses classified virulent podophages that infect P. aeruginosa into three genera and the phage 119X as an unassigned species.

Several studies have focused on a more comprehensive characterization at the genomic, transcriptional, and structural proteomics levels of the three podophage genera, Phikmvvirus (8, 9), Luz24virus (10, 11), and Lit1virus (12, 13). At the genomic level, the sizes ranged from 41.6 to 74.9 kb, and the GC content was typically lower (52% to 62%) than the average of many genomes of P. aeruginosa strains (66%). Direct terminal repeats are common in the three groups, ranging from 184 bp in Luz24 (10) to 665 bp in Lit1 phages (12). Genomes are organized into coding blocks harboring functionally related genes that typically correlate with the temporal nature of genome transcription observed during the lytic cycle (9, 11, 13).

There is little available information concerning 119X and its diversity and structural features. The present study provided an essential phylogenetic perspective that can lead to the recognition of a novel lineage of Pseudomonas phages.

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similarly identified were glaringly, this group of phages was clustered in a single group, and a partial sequence of one of these phages (PaMx32) suggested that these organisms are similar to 119X (16). Interestingly, these podophages showed the broadest host range among Mexican strains, productively infecting 30% to 60% of the more than 100 clinical isolates tested (16); however, these isolates were unable to infect the type strains PAO1 and PA14. To obtain knowledge about this group of phages, the genomes of five independent isolates were sequenced. This work describes and compares these five isolates, revealing a closely related group that includes 119X and PaP2, which are different from phages recognized previously. Because of their general similarity, we used phage PaMx41 as a model to describe the virion genome topology, protein composition, lytic cycle, and transcriptional timing of the functionally different gene clusters of the group. Differences at the genomic and transcriptional levels between P. aeruginosa podophages are discussed.

MATERIALS AND METHODS

Bacterial strains and phage propagation. Bacteriophages PaMx33, PaMx35, PaMx41, PaMx43, and PaMx46 were isolated from open water reservoirs and residual waters (16). A clinical strain, Pseudomonas aeruginosa Ps26, was used as the host bacterium because these phages were unable to adequately infect the type strains PAO1 or PA14 and the sequenced strains LESB58 and 39016. Host bacteria were grown overnight with shaking at 37°C in lysogeny broth (LB) medium. The phages were propagated using a standard agar overlay method, eluted from the agar, precipitated in 1.4 M NaCl and 16% (wt/vol) polyethylene glycol 8000 (PEG 8000) at 4°C, and subsequently purified through CsCl gradient centrifugation as previously described (17). Virion particles were stable for over 1 year of storage at 4°C.

Genome sequencing and assembly. Bacteriophage genomes were determined using high-throughput DNA sequencing using SOLID technology (Life Technologies, Grand Island, NY, USA). The reads were preprocessed using the Applied Biosystems de novo assembly accessories. Phage genomes were de novo assembled from 50-bp sequencing reads using Velvet v1.1 (18).

Genome composition and functional annotation. The genes were predicted using GenMark v1.1 (19), and ribosome binding sites were identified using rbs_finder.pl (20). Putative functions were assigned to predicted open reading frames (ORFs) based on the results of (i) blastp searches against GenBank nonredundant protein databases, (ii) structural similarities using HHpred (21) and E-TASSER (4), and (iii) a search of the protein conserved domains using InterPro (22). Putative P. aeruginosa promoter sequences (TTGACG-N15,18-TATAA [nucleotides in uppercase are present in more than 40% of the sequences; nucleotides in lowercase are present in more than 30% but less than 40% of the sequences]) (23) were predicted using Pattern Locator (24), PromoterHunter (25), and Neural Network Promoter Prediction (26) tools. Only those sequences detected using three tools were accepted. The E-CAI server was used to calculate the global genomic GC content and to determine the average percentage of codons ending in A+T+G.

Phage genome comparative analysis. The homology among the five newly reported phages and with 119X (GenBank accession number NC_007807.1) and PaP2 (GenBank accession number NC_005884.1) was observed via blastn against the nucleotide collection of NCBI. Pairwise genome comparisons at the nucleotide level were performed using MUMmer v3.0 (28) with a break length of 120 nucleotides. Single nucleotide polymorphisms (SNPs) were identified based on the global comparison of the seven phages using the progressiveMauve algorithm (29). The protein sequences of predicted ORFs were compared with all of the bacteriophage protein sequences deposited in the NCBI nonredundant protein database using blastp v2.2.30. An ORF was classified as a homolog between genomes when its blastp E value was lower than 1e−3 and the length of the shortest ORF among homologs was at least 35% of the longest sequence.

Analysis of virion structural proteins. PaMx41 structural proteins from CsCl-purified virions were separated using Tris-glycine sodium dodecyl sulfate-12% polyacrylamide gel electrophoresis (SDS-PAGE) using Precision Plus Protein Dual Xtra prestained protein standards (Bio-Rad, Hercules, CA) as molecular weight markers. The gel bands were excised and destained, and an in-gel protein digestion was performed using trypsin. The peptides were extracted and analyzed using a nanoscale liquid chromatography-electrospray ionization-tandem mass spectrometry (NanoLC-ESI-MS/MS) system. Protein identification against the virus nonredundant NCBI protein database (v1.69p; Matrix Science, London, United Kingdom) was conducted using Mascot search algorithms. The search parameters were set to monoisotopic mass values, allowing one missed cleavage, with a peptide mass tolerance of 1.2 Da and a fragment mass tolerance of 0.6 Da. The modifications were deamidation (NQ), oxidation (M), and propionamide (C).

Analysis of genome ends. Sequence coverage was assessed after mapping the sequencing reads on the PaMx41 genome using the SHRIMP package (30) and visualization using Tablet software (31). The in silico restriction pattern of PaMx41 was predicted using NEBcutter (32). Virion DNA of phage PaMx41 was digested using the Bal 31 exonuclease. The samples were removed at 0, 15, 30, and 45 min after the addition of the enzyme. The reaction was stopped with the addition of 50 mM EGTA and incubation at 70°C for 10 min. The samples were purified through phenol-chloroform extraction and resuspended in 10 μL of water. The DNA samples were digested with BglI endonuclease at 37°C for 2 h. The restriction pattern was visualized using 1% agarose gel electrophoresis.

One-step growth curve. To determine the phage growth characteristics, three independent single-step growth experiments were performed. Briefly, the host strain, Pseudomonas aeruginosa Ps26, was grown in liquid LB medium at 37°C until an optical density at 600 nm (OD600) of 0.4 (approximately 1 × 10^7 cells) was reached. One milliliter of culture was concentrated and suspended in 0.1 ml of LB that was supplemented with 10 mM MgCl_2, and an aliquot containing 1 × 10^8 phages was added to the culture, corresponding to a multiplicity of infection (MOI) of 0.1. Subsequently, the phages were adsorbed for 10 min at room temperature. The sample was diluted 1:1,000 in LB and incubated at 37°C for 120 min. The samples were collected at different times after infection for phage quantification. The samples were split in two, and one fraction was immediately plated on lawns of host bacteria, while the second fraction was treated with chloroform to kill the bacterial cells, infected or not, and the phage was quantified on lawns of host bacteria. The burst size was determined as the phage titer at 85 min (titer of directly plated samples) divided by the titer of infected cells at 20 min of infection (titer of directly plated samples minus titer of chloroform-treated samples at 20 min).

RNA and Northern blot methods. RNA was extracted from Ps26 cells that were infected with phage PaMx41 at an MOI of 10. Briefly, 50 μl of host culture at an OD_600 of 0.4 was harvested and resuspended in 1 ml of LB with 10 mM MgCl_2. An aliquot containing 5 × 10^6 phages was added to the cell suspension, and the phages were adsorbed for 10 min. The cells were washed with fresh LB, resuspended in 100 ml of LB, and incubated at 37°C. The samples were collected at 0, 15, 30, 45, 60, and 75 min after infection, supplemented with 0.3 mg/ml of rifampin and 0.5 mg/ml of chloramphenicol, and harvested on ice. RNA was obtained from the bacterial pellets through phenol-chloroform extraction and treated with RNase-free DNase I according to the manufacturer’s recommendations (Thermo Fisher Scientific, Waltham, MA). Total RNA was resuspended in...
TABLE 1 Genome sizes and nucleotide compositions of the five sequenced phages and two homologs in the GenBank database

<table>
<thead>
<tr>
<th>Phage</th>
<th>GenBank accession no.</th>
<th>Length (bp)</th>
<th>GC content (%)</th>
<th>C3+G3 (%)&lt;sup&gt;a&lt;/sup&gt;</th>
<th>No. of ORFs</th>
</tr>
</thead>
<tbody>
<tr>
<td>PaMx33</td>
<td>KU884561</td>
<td>43,265</td>
<td>45.25</td>
<td>41.37</td>
<td>54</td>
</tr>
<tr>
<td>PaMx35</td>
<td>KU884562</td>
<td>43,733</td>
<td>45.04</td>
<td>40.87</td>
<td>56</td>
</tr>
<tr>
<td>PaMx41</td>
<td>KU884563</td>
<td>43,490</td>
<td>45.21</td>
<td>41.52</td>
<td>55</td>
</tr>
<tr>
<td>PaMx43</td>
<td>KU884564</td>
<td>43,223</td>
<td>45.33</td>
<td>41.96</td>
<td>54</td>
</tr>
<tr>
<td>PaMx46</td>
<td>KU884565</td>
<td>43,266</td>
<td>45.24</td>
<td>41.41</td>
<td>55</td>
</tr>
<tr>
<td>PaP2</td>
<td>NC_005884.1</td>
<td>43,783</td>
<td>45.36</td>
<td>42.78</td>
<td>55</td>
</tr>
<tr>
<td>119X</td>
<td>NC_007807.1</td>
<td>43,365</td>
<td>44.89</td>
<td>40.67</td>
<td>53</td>
</tr>
</tbody>
</table>

<sup>a</sup>C3 + G3 refers to codons ending in G + C.

diethyl pyrocarbonate (DEPC)-treated water and stored at −80°C until further use. A total of 15 μg of each RNA sample was separated on a 1.2% agarose electrophoresis gel supplemented with 20 mM guanidine thiocyanate using Tris-borate-EDTA (TBE) as a running buffer. Formamide was used as a loading buffer. Subsequently, the samples were blotted onto Hybond–N+ membranes (GE Healthcare Biosciences, Little Chalfont, Buckinghamshire, United Kingdom) as previously described (33). RNA was fixed to the membrane through UV cross-linking for 5 min. The DNA fragments that were obtained after digesting PaMx41 genomic DNA were used as probes. Probes A and B were obtained using BglII; probes C, F, and G were obtained using BamHI; probes D and J were obtained with NcoI; probe E was obtained with BclI; probe H was obtained with AgeI; and probe K was obtained with SpeI. The restriction fragments were purified using QIAquick gel extraction kit (Qiagen, Germany). The DNA fragments were used as the templates for radiolabeling reactions with [α-<sup>32</sup>P]dATP (PerkinElmer, Boston, MA) using the random primer method (Random primer labeling kit; Amersham, Little Chalfont, Buckinghamshire, United Kingdom). The radiolabeled probes were purified using Centri-Sep spin columns (Life Technologies, Carlsbad, CA). Prehybridizing and hybridizing procedures were conducted in 7% SDS–0.5 M NaH2PO4 (pH 7.2)–2% dextran sulfate (34). The membranes were washed twice at 60°C with 1× SSC (1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate)/0.1% SDS, dried, and exposed. The film was developed after overnight exposure with an intensifying screen at −80°C.

Accession number(s). The complete genome sequences of the five phages were deposited in GenBank under accession numbers KU884561 (PaMx33), KU884562 (PaMx35), KU884563 (PaMx41), KU884564 (PaMx43), and KU884565 (PaMx46).

RESULTS AND DISCUSSION

Genomic characterization. Phages PaMx33, PaMx35, PaMx41, PaMx43, and PaMx46 are members of a podovirus group whose genomes displayed similar restriction and cross-hybridization patterns (16). Each phage lysed 30 to 44 strains of a collection of approximately 100 clinical <i>P. aeruginosa</i> isolates (16). To compare their genomic features, the complete sequences of the five genomes were assembled and annotated (see Materials and Methods). Three homologous phage genomes were identified, 119X, PaP2 (14), and PaMx32 (16). The genomic sequence of 119X was deposited in GenBank in 2006 (accession number NC_007807.1), but the phage was isolated in Australia approximately 50 years ago (7). PaP2 was isolated in China and its genomic sequence was deposited in GenBank in 2004 (accession number NC_005884.1), and PaMx32 is a member of the present group whose genome was partially sequenced (GenBank accession number JQ067091.1). The seven completely sequenced phage genomes were compared and analyzed. The genomes were approximately 43 kb long (Table 1) and contained 56 different ORFs (Table 1). Fifty-three ORFs were common to all phages, with five in insertions in various genomes (Fig. 1B). These sequences contained 45% GC content (Table 1 and Fig. 1A), which was in contrast with the approximately 66% GC content of <i>P. aeruginosa</i> strains (35). These differences impact the bias of codon usage in 119X-like phages, which contain 59% A3 + T3-ending codons versus 23% A3 + T3-ending codons for various other <i>P. aeruginosa</i> strains (see Materials and Methods). The consequences of this difference need further investigation.

Comparative genomics. The phage genome sequences were compared to evaluate their level of similarity. The results showed a high degree of conservation, even for phages isolated from distant global locations and time periods (from 99.95% to 89.5% identity). High levels of conservation have been reported in genomes of <i>Pseudomonas</i> (36) and <i>Brucella</i> (37) virulent phages isolated from distant places. Using progressiveMauve (29) and the genome of PaMx43 as a reference, we observed five large indels: three insertions (including one duplication) and one deletion (Fig. 1B; see also Table S2 in the supplemental material).

Additionally, 35 indels of 1 to up to 23 nucleotides were identified (see Table S1 in the supplemental material). Most of the 1-bp insertions and deletions were located in noncoding regions, and those located within coding regions generated translational frameshifting and were positioned close to the 5′ and 3′ ends. The 5′ indels shifted the beginning of the protein to the next start codon, whereas the 3′ indels produced a premature stop codon, and both cases resulted in shorter proteins but might conserve function because the core sequence remained unchanged. In addition, short indels that were longer than 1 bp within ORFs were in-frame variants that predictably exerted little effect on the putative proteins. Additionally, a total of 3,956 single nucleotide polymorphisms (SNPs) were observed in the global genome comparison; however, the number of SNPs in pairwise alignments varied from 25 to 2,270 (see Table S2 in the supplemental material). Notably, the PaMx43 genome had the lowest number of SNPs (610) relative to that of PaP2 from China (see Table S2). Thus, except for the ORF31 duplication in PaP2, this phage was more related to PaMx43 than to any other Mexican phage described here. The observed variations suggest a strong preference of the genomes to conserve functions despite their geographical or temporal isolation.

Although at the nucleotide level no other sequence showed similarity to the genomes of 119X-like phages, distant relationships were observed at the protein level (38). By comparing the amino acid sequences of the putative proteins encoded in the analyzed genomes, it was found that only seven sequences were shared between different phages (see Materials and Methods): large terminase, endolysin, MarzG-like protein, DNA polymerase, helicase, ORF24, and ORF35. No more than two homologous proteins were shared with the same phage. However, several proteins...
probes, and the mRNAs shown below overlapped a region of probes ORFs, phogenesis, and lysis modules (Fig. 1B shows the putative assigned long to nucleotide, DNA, and RNA metabolism, packaging, morphogenesis, and gray, unknown function. The identified functions be-
tween the upstream ORFs (ORF14 and ORF37, respectively). The assign-
codons of ORF15 and ORF36 overlapped with the stop codons of
that were encoded in 119X-like phages matched the hypothetical proteins observed in large contigs from metagenome data (see Table S3 in the supplemental material) that were obtained from sludge and groundwater (39, 40), suggesting that related phages might be observed in a wide range of water reservoirs. The unique nature of most of the coding sequences from 119X-like phages facilitated the discrimination of this group.

Genome functional annotation. We observed 53 ORFs con-
served in each of the seven genomes analyzed (Table 1) and only three accessory ORFs, namely, a, b and c, in various genomes (Fig. 1B). Ribosome binding sites were observed for all of the predicted ORFs, except for ORF15, ORF36, and ORF47. Indeed, the start codons of ORF15 and ORF36 overlapped with the stop codons of the upstream ORFs (ORF14 and ORF37, respectively). The assignment of functions to the recognized ORFs was based on the pre-
dicted amino acid sequences and structural similarities that were shared with characterized proteins (see Materials and Methods). A total of 27 ORFs were not homologous to any other sequence, and 13 ORFs resembled proteins without assigned functions from metagenomics data and other phage and bacterial origins. Only 16 ORFs possessed functionally conserved domains that were similar to proteins with a known function. The identified functions be-
tween the nucleotide, DNA, and RNA metabolism, packaging, morphogenesis, and lysis modules (Fig. 1B shows the putative assigned functions, and Table S3 in the supplemental material summarizes the criteria that were used for function assigment).

Transcription program and regulatory sequences of the PaMx41 genome. The lytic cycle of viruses is an orderly gene expression program that is coordinated with the availability of specific proteins needed at different times of infection (41). The di-
versity of strategies used by different groups of phages to achieve successful propagation makes it hard to predict the transcriptional timing of a specific phage (42). To obtain knowledge concerning the timing of the lytic cycle and improve the annotation of 119X-
like phages, we evaluated the temporal pattern of gene expression for PaMx41. One-step growth curve assays of phage PaMx41 (see Fig. S1 in the supplemental material) showed that the eclipse pe-
ned after infection through Northern blot analyses using 11 restriction
were identified using both probes, and the mRNAs shown below overlapped a region of probes I, J, and K were identified using the three probes.

![Consensus genome map with functional annotation of 119x-like phages and transcription program of PAMx41.](aem.asm.org/.../b22017544.jpg)
were revealed with probe A (Fig. 1C; see also Fig. S2 in the supplemental material), which corresponded to the gene cluster next to the left terminus (ORF1, ORFa, ORF2, and ORF3). The products of the four early ORFs lack similarity with other known proteins but may be related to host gene shutoff transcriptional regulation or antitermination (43, 44). Interestingly, E1 but not E2 was a transient transcript observed only at 15 min of infection (see Fig. S2). Phage early mRNA shutdown has been reported in T7 (45) and in the phage PhiKZ; early mRNA shutdown is observed after the early stage of infection (46). Probes G, H, I, J, and K detected 11 middle mRNAs covering the leftward-oriented cluster in the right arm of the genome (Fig. 1B and C), expressing functions associated with DNA replication and nucleic acid metabolism. As the 2-kb early region is duplicated at both ends of the genome, it is likely that middle transcription may be physically linked to early transcription from right-end duplication.

The late transcripts included seven mRNAs that were revealed using probes B to F (Fig. 1C) and comprise morphogenesis, virion structural, and lysis ORFs (Fig. 1; see also Fig. S2 in the supplemental material). A short signal was observed at a position corresponding to approximately 0.9 to 0.7 kb in almost all of the autoradiograms, irrespective of the probe used, and was considered to be nonspecific, so it was not taken into account. Most of the phage mRNAs remained detectable until the end of the infection, but during lysis at 75 min, most of the signals disappeared (see Fig. S2), which likely reflects mRNA degradation at the start of cell lysis.

The three temporal classes of mRNAs that are observed above suggest a classical phage transcriptional program. However, because the mRNA ends were not identified, it was difficult to infer the location of the promoters originating the transcripts. Therefore, we conducted a search of P. aeruginosa σ70 promoter consensus sequences. Only one putative promoter that was recognized by the host RNA polymerase and was identified in the negative strand of the PaMx41 genome upstream of ORF3 (at 1,915 to 1,950 bp; AGGCCTGACTTCTAAGGCAAGACATTAGCACCC; −35 and −10 are shown in italics). Although, the transcription results represent a specific case of PaMx41, the putative promoter was conserved in the seven compared phages. Thus, 119x-like phages may depend on the host RNA polymerase to initiate the transcription of early genes. σ70 promoter sequences were not identified in the rest of the genome, and an RNA polymerase homolog was not observed among the predicted ORFs. We hypothesized that the specificity of the host RNA polymerase is changed by an early stage of infection (46).

Identification of the virion genome termini. The sequencing reads of the five sequenced genomes may be assembled circularly. However, in tailed phages, this topology may also result from direct terminal redundancy of a linear genomic DNA segment, where the sequencing coverage of the redundant segment is twice that of the nonredundant portion (48). Indeed, we detected a 2-kb segment showing approximately twice the sequencing coverage as that of the rest of the genome (Fig. 2A). The coverage values suggested that the duplication of approximately 2 kb might represent the chromosome ends of PaMx41, and the hypothesized model is depicted in Fig. 2B.

To examine the suggested genome topology, we conducted restriction assays of the genomic PaMx41 DNA using the endonuclease BglI. The model shown in Fig. 2B indicates that 11 bands are expected from a linear topology. The restriction fragment corresponding to the right end of the terminally redundant genome should be approximately 2.7 kb instead of the 580 bp expected in the linear nonredundant genome (Fig. 2B). The restriction assay (Fig. 2C) showed a total of 11 bands, including the left-end and right-end fragments (5 and X in Fig. 2C, lane 2), which was consistent with the direct terminal redundancy topology. Moreover, a time course treatment of the PaMx41 DNA with exonuclease Bal 31, followed by BglI restriction, resulted in the rapid loss of the end fragments and sequential reduction of the internal fragments (Fig. 2C, lanes 3 to 5). The outcome of such treatment was consistent with the linear topology for the virion genome. Taken together,
PaMx41 (Fig. 1). Based on structural similarities, we identified the clusters between these phages. Thus, gp 10 variations may be responsible for the host range differences. To our knowledge, portal proteins or recombination proteins have never been associated with receptor recognition; this information is still rare in the databases. Most of the putative proteins that are associated with a specific function, or even the experimentally identified virion proteins, remain without an assigned function. The transcriptional study of this group revealed a classical gene expression program organized into early, middle, and late genomic modules. The uniqueness of the early genes and lack of recognizable middle and late promoter sequences in this group of phages suggested that these sequences possess particular elements of the phage-bacteria interaction. The results facilitated the integration of 119X-like phages as a unique lineage within the Podoviridae family.

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**REFERENCES**


**FIG 3** PaMx41 virion structural proteins. The protein components of PaMx41 purified virions were resolved using 12% SDS-PAGE and stained with Coomassie brilliant blue. Observed bands were excised from the gel and MS-MS treated, and the data were analyzed and correlated with the predicted amino acid composition encoded by phage genes. Sequence coverage is the percentage of the protein amino acid sequence covered by the matching peptides. Molecular mass (kDa) was calculated from the gene sequence. The Mascot score is ~10log10 the probability of matching the mass value calculated for the corresponding gene.
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