Genome sequence and characterization of *Streptomyces* phage Pablito, representing a new species within the genus *Janusvirus*

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Streptomycetes are ubiquitous soil bacteria. Here we report the complete and annotated genome sequence and characterization of *Streptomyces* phage Pablito, isolated from a soil sample in Haarlem, the Netherlands using *Streptomyces lividans* as host. This phage was able to infect a diverse range of *Streptomyces* strains, but none belonging to the sister genus *Kitasatospora*. Phage Pablito has double-stranded DNA with a genome length of 49,581 base pairs encoding 76 putative proteins, of which 26 could be predicted. The presence of a serine integrase protein indicated the lysogenic nature of phage Pablito. The phage remained stable over a wide range of temperatures (25–45 °C) and at pH ≥ 7.0, but lost infectivity at temperatures above 55 °C or when the pH dropped below 6.0. This newly isolated phage is closely related to *Streptomyces* phage Janus and Hank144 and considered a new species classified in the genus *Janusvirus*, within the subfamily *Arquattrovirinae*.

*Streptomyces* spp. are Gram-positive, spore-forming bacteria that thrive in almost all soil environments. They are well-known for their ability to produce a wide array of clinically-relevant antibiotics, such as streptomycin, neomycin, chloramphenicol and many more¹–³. Most research has been focused on the industrial use of these bacteria and their secretion of secondary metabolites⁴,⁵. However, it remains largely unknown how bacteriophages can recognize, attach and infect these multicellular bacteria. *Streptomyces* have an unusual lifecycle, which starts when spores encounter a favorable environment. This triggers germination and outgrowth into a so-called vegetative mycelium. After a period of vegetative growth, aerial hyphae are formed that can develop into chains of spores. These spores can be dispersed to more favorable environments to establish new colonies. Previous research has shown that *Streptomyces* phages can only infect young vegetative mycelia, but are unable to infect spores⁶,⁷. However, most bacteriophage research has been focused on unicellular bacteria, while the interactions and underlying mechanisms of bacteriophages infecting multicellular bacteria, like *Streptomyces*, remains largely unexplored. Besides the phage growth limitation system⁸,⁹, chemical defense¹⁰ and recently discovered cell wall-deficiency¹¹, many more defense mechanisms against phage attack might be present in *Streptomyces*. In addition, relatively few *Streptomyces* phages have been properly sequenced and characterized in comparison to other genera. Therefore, it is important to isolate new phages infecting multicellular bacterial species, allowing us to expand our knowledge about the immense viral diversity.

In this paper, we describe a newly discovered Actinobacteriophage, called *Streptomyces* phage Pablito, which was isolated from a soil sample in the Netherlands. This phage belongs to the genus *Janusvirus*, within the subfamily *Arquattrovirinae*. Phage Pablito is a temperate phage with an icosahedral capsid head of 56.1 ± 1.3 nm and a long non-contractile tail of 163.5 ± 0.4 nm. Finally, we provide a whole genome analysis of this newly identified phage. Together, these results shed light on phage taxonomy and can expand the knowledge on phage-*Streptomyces* interactions in the future.

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Materials and methods

Bacteriophage isolation. Phage Pablito was isolated from a soil sample obtained from the National Park Zuid-Kennemerland in the Netherlands (N52°23′31″, E4°34′49″). To this end, 0.5 g soil was mixed with spores from the host S. lividans strain 1326 in 10 ml Difco Nutrient Broth (DNB) (BD Biosciences) supplemented with 0.5% glucose and 4 mM Ca(NO₃)₂ and incubated overnight at 30 °C, while shaking at 130 RPM. After 24 h, the culture was centrifuged for 10 min at 4000 g and the supernatant containing phages was filtered through a 0.22 μm filter (Millipore). The supernatant was subsequently spread onto a DNB agar plate, followed by applying an overlay (in DNB soft agar) containing 10³ spores ml⁻¹ from S. lividans. The plate was incubated at 30 °C for 24–48 h before plaque formation was assessed. Single plaques were purified by streaking them twice to a fresh double agar overlay plate containing the bacterial host. High titre single viral stocks were obtained by picking purified plaques and incubating them with S. lividans in DNB medium for 10 min before plating on double-agar overlay DNB plates. Viral lysates were collected by flooding the plate with 24 ml DNB and gently rocking the plate for 2 h. The supernatant was filtered through a 0.22 μm filter. Collected phages were stored at 4 °C.

Host range analysis. The host range of phage Pablito was determined using serial dilutions on double agar overlay plates. In short, 3 µl of the phage dilution was spotted in duplicate on a bacterial lawn of different Streptomyces and Kitasatospora strains from the Microbial biotechnology (MBT) collection at Leiden University, which represents an in-house collection of isolated actinomycetes. The plates were incubated at 30 °C for at least 24 h before lysis was observed. The presence of a lysis zone was scored as a positive result. The efficiency of plating (EOP) for a given strain was calculated relative to the host strain S. lividans when plaques were observed.

Stability of phage Pablito. A one step-growth curve analysis was performed in triplicate to calculate the latent period and phage burst size. In short, 10⁵ spores ml⁻¹ of S. lividans were allowed to germinate in 10 ml DNB medium at 30 °C for 5 h. At T = 0, the culture was infected with phage Pablito at Multiplicity of Infection (MOI) = 0.01 and 100 µl was sampled at 10 min intervals up to 180 min. The samples were filtered, diluted and immediately plated on double agar plates. Burst size was calculated using the following formula:

\[
\text{Burst size} = \frac{\text{average of free phages after burst (T = 90 to T = 180)} - \text{average of free phages before burst (T = 10 to T = 80)}}{\text{number of phages at T = 0}}
\]

The thermal stability of phage Pablito was determined by incubating 1 ml of lysate (10⁶ PFU ml⁻¹) at 25, 30, 37, 45, 55 and 65 °C. The samples were filtered after 1 h of incubation and immediately plated on double agar plate to determine the phage titre. For pH stability, 10 µl of a phage Pablito suspension (10⁶ phages ml⁻¹) was added to 990 µl DNB medium adjusted to different pH values (2–9) using 6 M HCl or 1 M NaOH. The tubes were then incubated at 30 °C for 1 h before the samples were filtered through a 0.22 μm filter and plated using the double agar overlay method. All measurements were performed in triplicate.

DNA extraction and phylogenetic analysis. DNA of phage Pablito was isolated using the Phage DNA Isolation Kit from Norgen (Thorold, Canada) according to the manufacturers protocol. After isolation, multiple samples were pooled and DNA was concentrated using the Concentrator Plus (Eppendorf) for 20 min. Whole-genome sequencing and de novo assembly of phage Pablito was performed by BaseClear using Illumina NovaSeq PE150 sequencing with a mean coverage of 118.21 (Leiden, the Netherlands). The complete genome sequence of Streptomyces phage Pablito was deposited at GenBank under accession number OK412919. A linear map of the genome was constructed by Geneious Prime 2020.1.2 and PhageTerm was used to determine genome ends.

Phylogenetic analysis was performed using the putative protein sequence of the major capsid head protein. The protein sequences of Streptomyces phage Janus, Hank144, Pablito and Joe were acquired from NCBI and aligned using MUSCLE. A Maximum Likelihood phylogenetic tree was constructed through Molecular Evolutionary Genetics Analysis Version X (MEGAX) using 100 bootstrap replicates to estimate the genus of Phage Pablito.

Electron microscopy. Two liquid cultures of S. lividans infected with phage Pablito were filtered, combined and precipitated overnight with 5×PEG solution (20% PEG8000 in 2.5 M NaCl) at 4 °C and centrifuged for 70 min at 3500 g after which the supernatant was discarded. Of the remaining pellet, 3 µl was placed on a glow discharged 200 mesh carbon coated copper grid (EMS) and allowed to set for 30 s before excess sample solution was removed by filter paper. The phages were stained with 2% uranyl acetate for 45 s and air-dried for an additional 30 min. The grids were observed using a single tilt specimen holder inside a 120 kV Talos L120C TEM with a Lab6 electron source and Ceta detector at the Netherlands Center for Nanoscopy (NeCEN, Leiden).

Results and discussion

Morphology and host range determination of Pablito. Bacteriophage Pablito produced clear small plaques on the S. lividans host strain using the double agar overlay method (Fig. 1a). Transmission electron microscopy (TEM) was used to reveal the morphology of phage Pablito. This demonstrated that phage Pablito has an icosahedral capsid head of 56.1 ± 1.3 nm (n = 3) and a long non-contractile tail of 163.5 ± 0.4 nm (n = 3) (Fig. 1b). The long, flexible tail together with the icosahedral capsid head were the first indication that this phage is a siphovirus belonging to the class Caudoviricetes, which was further confirmed by whole-genome sequencing (see below).

A host range assay on 10 actinomycetes revealed that all eight Streptomyces strains tested could be infected by phage Pablito (Table 1). Depending on the host strain, some lysis zones showed a turbid morphology, which
is common for several Streptomyces phages\(^{16}\), and was the first indication that phage Pablito might be a temperate phage. The EOP for a strain relative to the host strain *S. lividans* from which the phage was isolated, is given when plaques were countable. On the contrary, the two species that remained uninfected (i.e. MBT66 and *K. viridifaciens*) belong to the sister genus *Kitasatospora*, members of actinomycetes which are morphologically similar to *Streptomyces* but generally classified by their resistance against *Streptomyces* phages\(^{17,18}\).

**Genomic analysis.** Whole genome sequencing and de novo assembly of phage Pablito revealed a double-stranded DNA genome of 49,581 base pairs. While its host *S. lividans* has relatively high G + C content of 72%, *Streptomyces* phage Pablito has a G + C content of 66.4% and a 3′-cohesive termini of CGC CGT GTCTT (Fig. 2). The genome contains 76 potential coding sequences (CDSs), of which 50 encode for hypothetical proteins, while the function of 26 CDSs could be predicted. The latter were categorized as morphogenesis proteins, nucleotide metabolism and DNA replication proteins, lysogenic module and a bacterial lysis protein as indicated in Fig. 2.

Phylogenetic analysis (Fig. 3a) based on the putative major capsid head protein revealed that *Streptomyces* phage Pablito is similar to *Streptomyces* phage Janus and *Streptomyces* phage Hank144, while *Streptomyces* phage Joe is more distantly related. Based on the criteria from the Bacterial Viruses Subcommittee, 70% nucleotide identity is established as the cut-off for genera, while 95% identity assigns phages to the same species\(^{19}\). As calculated by the intergenomic distance calculator VIRIDIC\(^{20}\), *Streptomyces* phage Pablito shows 70.0 and 72.5% relatedness to *Streptomyces* phages Janus and Hank144, respectively (Fig. 3b). Since the relatedness of *Streptomyces* phage Pablito is lower than 95%, this phage is considered a new species according to the International Committee on Taxonomy of Viruses (ICTV)\(^{20}\). Because *Streptomyces* phage Janus and *Streptomyces* phage Hank144 are both temperate phages and whole genome sequencing revealed the presence of a serine integrase protein in *Streptomyces* phage Pablito, we conclude that phage Pablito is a temperate phage as well\(^{21}\). Together, these data indicate that *Streptomyces* phage Pablito is a newly discovered species that can be included in the genus

| Bacterial strain          | Origin       | Infectivity | Lysis zone morphology | EOP  |
|---------------------------|--------------|-------------|-----------------------|------|
| *Streptomyces lividans*   | MBT collection | +           | Clear                 | 1    |
| *Streptomyces coelicolor* | MBT collection | +           | Turbid                |      |
| *Streptomyces griseus*    | MBT collection | +           | Turbid                |      |
| *Streptomyces venezuelae* | MBT collection | +           | Turbid                |      |
| *Streptomyces albus*      | MBT collection | +           | Clear                 | 0.3  |
| *Kitasatospora viridifaciens* | DSM40239 | −           | None                  |      |
| MBT13                     | MBT collection | +           | Turbid                |      |
| MBT61                     | MBT collection | +           | Turbid                |      |
| MBT66                     | MBT collection | −           | None                  |      |
| MBT86                     | MBT collection | +           | Clear                 | 1.5  |

**Table 1.** Susceptibility of actinomycete strains to infection by phage Pablito as determined by plaque assays.

**Figure 1.** Morphology of phage Pablito. (a) Plaque morphology of phage Pablito after 24 h of infection in *S. lividans* using the DNB double agar overlay plate. (b) Representative TEM images of phage Pablito. Phage particles were stained with 2% uranyl acetate.
Janusvirus. A taxonomic proposal has been submitted to ICTV to officially recognize phage Pablito as part of the genus Janusvirus as a new species called "Janusvirus Pablito". The Actinobacteriophage Database (https://phagesdb.org/) places phage Pablito in Cluster BD, Subcluster BD2. In addition, phage Pablito is peripherally related to two prophages found in Streptomyces finlayi strain NBSH44 and Streptomyces venezuelae strain ATCC 14583.

One-step growth curve. Streptomyces phage adsorption was found to be maximal for germlings. We therefore incubated S. lividans spores first for 5 h in DNB medium before infecting the culture with bacteriophages. The one-step growth curve of phage Pablito showed a relatively long latent period of 80 min, with a burst size of 22 virions per cell (Fig. 4). However, another rise of viral titer was observed around 150 min, which can happen when multiple phages bind to one germinated spore, as a sign of delayed adsorption or as an indication of a lysogenic infection cycle. The highest titre obtained was around $10^6$ PFU ml$^{-1}$ after 24 h. A different range of MOI, longer incubation periods and even flooding of a confluent plaque plate did not result in an increase of the total number of phages ml$^{-1}$.

Phage stability. We also assessed the effect of temperature and pH on phage viability. Phage Pablito was stable up to 45 °C without a reduction in viability (Fig. 5a). However, the infectivity significantly decreased when the phage was incubated for an hour at 55 °C or higher. Phage Pablito was relatively stable at pH 7.0–9.0, while infectivity steeply declined when the pH was lower than 6.0 (Fig. 5b). No plaques were found when the pH was below 4.0, while only a small amount of activity was retained at pH 5.0 and pH 6.0.

Conclusion
This study presents Streptomyces phage Pablito as a temperate siphophage being able to infect diverse Streptomyces strains. Genomic analysis and TEM images showed that this phage could be classified as a new species in the genus Janusvirus within the subfamily Arquattrovirinae. Members of this subfamily/genus have long non-contractile tails and icosahedral capsid heads. Phage Pablito showed a burst size of 22 virions per cell and was stable at temperatures ranging from 25 to 45 °C and a pH between 7.0 and 9.0. This phage can be used in future studies to shed new light on bacteriophage-Streptomyces interactions and disclose how this phage affects soil community.
Data availability
The genome of *Streptomyces* phage Pablito is available on NCBI (GenBank Accession No. OK412919).

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**Author contributions**

The experimental plan was designed by V.O., A.B. and D.C.. The phage genetics were analyzed by V.O. and A.M.K.. The paper was written by V.O. and the final manuscript was shaped, edited and reviewed by all authors.

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**Competing interests**

The authors declare no competing interests.

**Additional information**

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