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### **Title**

Comparative genomics of Cluster O mycobacteriophages.

### **Permalink**

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# **Journal**

PloS one, 10(3)

### **ISSN**

1932-6203

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# **Publication Date**

2015

### DOI

10.1371/journal.pone.0118725

Peer reviewed







Citation: Cresawn SG, Pope WH, Jacobs-Sera D, Bowman CA, Russell DA, Dedrick RM, et al. (2015) Comparative Genomics of Cluster O Mycobacteriophages. PLoS ONE 10(3): e0118725. doi:10.1371/journal.pone.0118725

Academic Editor: Mark J van Raaij, Centro Nacional de Biotecnologia - CSIC, SPAIN

**Received:** December 11, 2014 **Accepted:** January 13, 2015

Published: March 5, 2015

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**Data Availability Statement:** Phage genome sequences are available in GenBank. The accession numbers for these sequences are provided in Table 1.

**Funding:** This work was supported by grants from the National Institutes of Health (GM51975) and from the Howard Hughes Medical Institute (54308198). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Competing Interests:** The authors have declared that no competing interests exist.

RESEARCH ARTICLE

# Comparative Genomics of Cluster O Mycobacteriophages

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### **Abstract**

Mycobacteriophages – viruses of mycobacterial hosts – are genetically diverse but morphologically are all classified in the Caudovirales with double-stranded DNA and tails. We describe here a group of five closely related mycobacteriophages – Corndog, Catdawg, Dylan, Firecracker, and YungJamal – designated as Cluster O with long flexible tails but with unusual prolate capsids. Proteomic analysis of phage Corndog particles, Catdawg particles, and Corndog-infected cells confirms expression of half of the predicted gene products and indicates a non-canonical mechanism for translation of the Corndog tape measure protein. Bioinformatic analysis identifies 8–9 strongly predicted SigA promoters and all five Cluster O genomes contain more than 30 copies of a 17 bp repeat sequence with dyad symmetry located throughout the genomes. Comparison of the Cluster O phages provides insights into phage genome evolution including the processes of gene flux by horizontal genetic exchange.

### Introduction

The bacteriophage population is vast, dynamic, and old, spanning considerable genetic diversity [1–3]. Phages of phylogenetically distant hosts typically share little nucleotide sequence similarity and few genes encoding proteins with amino acid sequence similarity [4]. Phages also typically encode a high proportion of genes with no sequence similarity to proteins outside of the phages of that particular host, and the global phage population likely harbors the largest reservoir of unexplored sequence information [5]. Phages of a single common host may also show substantial nucleotide sequence variation, although the diversity is expected to be dependent on the diversity of the bacterial population within the environment from which those phages are isolated [6].

Mycobacteriophages—viruses of mycobacterial hosts—display considerable genetic diversity and GC% content [7, 8]. Comparative genomics of over 290 fully sequenced mycobacteriophage genomes shows that they can be divided into groups of closely-related genomes referred to as clusters, several of which can be further divided into subclusters. [7]. There are currently 20 clusters (A-T) and nine singleton phages (those without any close relatives), and ten of the clusters are subdivided into subclusters (phagesdb.org). The diversity of these phages varies among these various groups, with some containing closely related genomes sharing >90% of their genes, whereas others are highly diverse. The genomes are typically mosaic in their architectures, with individual genes or groups of genes present in a multitude of different genomic contexts [9].

Mycobacteriophage Corndog was isolated using *M. smegmatis* mc<sup>2</sup>155 as a host and was previously described as a singleton phage with an unusual prolate head [9]. The vast majority of mycobacteriophages have siphoviral morphologies, most of them with isometric heads. The exceptions are Corndog and the phages in Cluster I, although their dimensions differ; the length:width ratio of the capsids is 2.5:1 and 4:1 for Cluster I phages and Corndog respectively [8]. Corndog is also unusual in that the viral genome contains an atypically short (4-base) 3' single strand extension, and appears to use non-homologous end joining to recircularize the genome upon infection, a process likely facilitated by a phage-encoded Ku protein [10]. Corndog does not infect *M. tuberculosis* or *M. smegmatis* Jucho, and plates at a greatly reduced efficiency on *M. smegmatis* MKD8 relative to *M. smegmatis* mc<sup>2</sup>155 [6]. The genome was noted to



contain several unusual features including genes coding for methylases and glycosylases within the structural genes, a DNA Polymerase Beta clamp, and an AAA ATPase [9]. Corndog does not encode an integrase and stable lysogens have not been reported [8].

Here we describe four mycobacteriophages—Catdawg, Dylan, Firecracker, and YungJamal—with strong nucleotide sequence similarity to phage Corndog such that all five genomes constitute Cluster O. These genomes are sufficiently similar that dividing the cluster into subclusters is not warranted, and all five exhibit the prolate capsid morphology described for Corndog [9]. Genome comparisons reveal several notable features including putative transcriptional promoters and an unusual 17 bp repeated motif present more than 30 times in each genome. Proteomic analysis of purified Corndog virions and Corndog infected cells identifies about half of the predicted gene products including many small non-structural proteins of unknown function and one previously unannotated gene. Additional proteomic analysis of an unpurified lysate of Catdawg virions identifies a similar proportion of the predicted gene products.

### **Results**

# Five mycobacteriophages constitute Cluster O

Mycobacteriophage Corndog was isolated in 2001 [9] and until 2012 was designated as a singleton phage without any close relatives [11]. Since 2012, four phages—Catdawg, Dylan, Firecracker, and YungJamal—have been found that are related to Corndog and constitute Cluster O (Table 1, Fig. 1). They were isolated in the Science Education Alliance Phage Hunters Advancing Genomics and Evolutionary Science (SEA-PHAGES) program [12], the Mycobacterial Genetics Course held at the University of KwaZulu Natal (UKZN MGC) and the Phage Hunters Integrating Research & Education (PHIRE) Program at the University of Pittsburgh. The five Cluster O phages have similar genome lengths (69.8-72.1 kbp) and all contain unusually short (4-nucleotide) 3' single-stranded terminal extensions (Table 1). They have 122–128 predicted protein-coding genes and do not contain tRNA or tmRNA genes (Table 1). The five genomes are closely related at the nucleotide level (Fig. 1) and share high levels of average nucleotide identity (Table 2) that do not warrant division into subclusters. The Cluster O phages are not closely related to other mycobacteriophages although there is nucleotide sequence similarity to Subcluster I1 phages such as Brujita and to a lesser extent subcluster F1 phages such as GUmbie (Fig. 1). The GC% contents are similar to M. smegmatis (which is 67.4% GC; Table 1) as are the codon usage profiles (data not shown).

All five Cluster O phages have similar virion morphologies and are members of the Siphoviridae containing long, flexible non-contractile tails approximately  $248\pm8$  nm in length. However, they have unusual prolate heads with a length of  $165\pm2$  nm and width of  $38\pm1$  nm (length:width ratio of 4:1; Fig. 2).

Table 1. Cluster O Mycobacteriophages.

| Phage Name  | Accession # | Genome Length (bp) | GC%  | Overhang Sequence | # ORFs | Location             |
|-------------|-------------|--------------------|------|-------------------|--------|----------------------|
| Catdawg     | KF017002    | 72108              | 65.4 | GTGT              | 128    | Radnor, PA USA       |
| Corndog     | AY129335    | 69777              | 65.4 | GTCT              | 124    | Pittsburgh, PA USA   |
| Dylan       | KF024730    | 69815              | 65.4 | GTGT              | 122    | Durban, South Africa |
| Firecracker | JN698993    | 71341              | 65.5 | GTGT              | 127    | Santa Cruz, CA USA   |
| YungJamal   | KJ829260    | 70214              | 65.3 | GTCT              | 124    | Pittsburgh, PA USA   |

doi:10.1371/journal.pone.0118725.t001



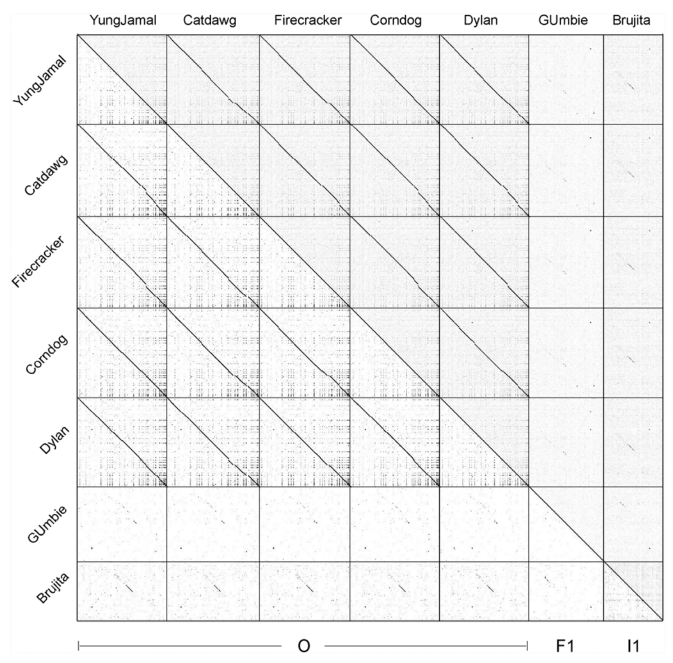


Fig 1. Dotplot comparison of Cluster O mycobacteriophages. The five Cluster O phages along with GUmbie (Subcluster F1) and Brujita (Subcluster I1) were compared using Gepard [13] and the dotplots displayed at two different levels of sensitivity and contrast in the upper right and lower left triangles.

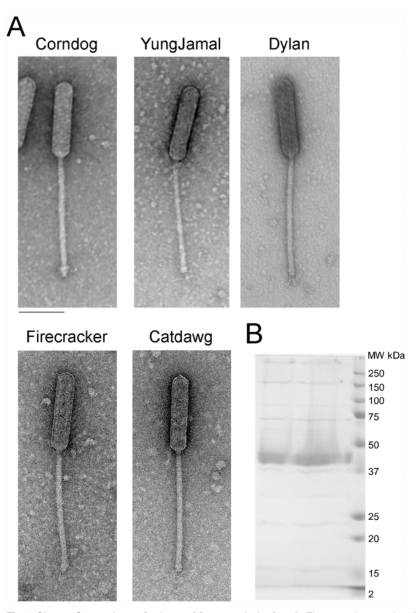
# Cluster O Genome Organizations

The five Cluster O genomes share similar organizations but differ with a variety of small insertions and deletions corresponding to one or a small number of genes (S1 Fig.; Figs. 3–7). The genomes contain three blocks of genes that likely correspond to transcriptional units. The first is a group of 10–12 leftwards-transcribed genes of mostly unknown functions at the left end of the genomes. The second is a large group of rightwards-transcribed genes (e.g. Corndog 11–72)



Table 2. ANI values for cluster O phages.

|             | Catdawg | Corndog | Dylan | Firecracker | YungJamal |
|-------------|---------|---------|-------|-------------|-----------|
| Catdawg     | 1       | 0.977   | 0.978 | 0.973       | 0.977     |
| Corndog     | 0.977   | 1       | 0.987 | 0.987       | 0.991     |
| Dylan       | 0.978   | 0.987   | 1     | 0.987       | 0.982     |
| Firecracker | 0.973   | 0.987   | 0.987 | 1           | 0.985     |
| YungJamal   | 0.977   | 0.991   | 0.982 | 0.985       | 1         |



**Fig 2. Cluster O mycobacteriophage virion morphologies. A.** Electron micrographs of Cluster O phages. Scale bar corresponds to 100 nm. **B.** SDS-PAGE analysis of Corndog virions.

doi:10.1371/journal.pone.0118725.g002



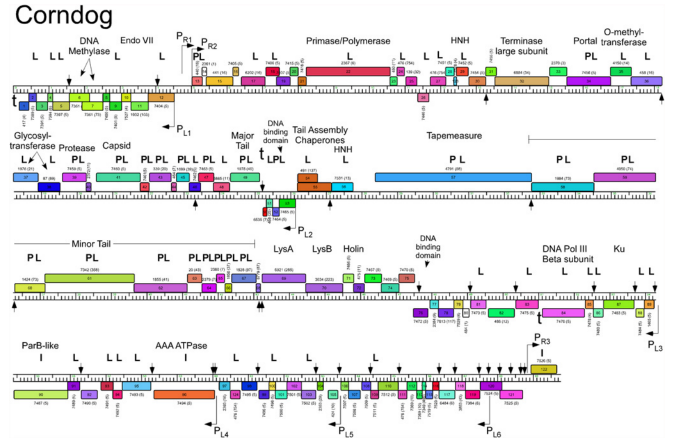


Fig 3. Genome map of Mycobacteriophage Corndog. The genome of phage Corndog is represented as a scale bar (major intervals: 1 kbp) with predicted genes shown as boxes either above (rightwards transcribed) or below (leftwards transcribed). Gene number is shown within each box and the phamily designation is shown either above or below with the number of phamily members shown in parentheses. Putative gene functions are indicated. The positions of putative SigA-like promoters ( $P_{L1}$ — $P_{L6}$  and  $P_{R1}$ — $P_{R3}$ ) are shown as large arrows and terminators (t) are indicated. Small vertical arrows show the locations of the palindromic repeat 5'-TGTTCGGNNNCCGAACA. Gene products identified by mass spectrometry (with at least two high confidence peptides per product) in twice CsCl banded particles (P) or from a once-banded lysate (P) are indicated, as well as three additional proteins identified in infected cells (P) not identified in the other samples. Proteins gp11, gp33, gp77, and gp102 had multiple high quality spectra (P2, P3, and P4 respectively) of a single peptide each.

containing the virion structure and assembly genes as well as the lysis cassette, although this is interrupted by up to four instances of a small number of small leftwards-transcribed genes. A third set of  $\sim$ 50 genes (e.g. Corndog 75–124) is transcribed leftwards, and a single gene at the extreme right end of the genomes is transcribed rightwards (Figs. 3–7).

Database comparison and HHPred searches reveal putative functions for fewer than 20% of the genes, although additional virion structure and assembly proteins are predicted based on synteny (Figs. 3–7). Unusually, the large terminase subunit gene is displaced ~14 kbp from the left cohesive end and an O-methyltransferase gene, two glycosyltransferase genes and a putative N-acetylglucosaminyltransferase gene are located between the portal and the capsid maturation protease genes. Of the small leftwards-transcribed genes within the virion structural operon, only one—a putative DNA binding protein (e.g. Corndog 53)—has a predicted function. Five genes within the long leftwards-transcribed region encode proteins with predicted functions including a DNA binding protein, a beta clamp subunit of DNA Polymerase III, a Ku-like protein, an AAA ATPase, and a ParB-like domain protein.



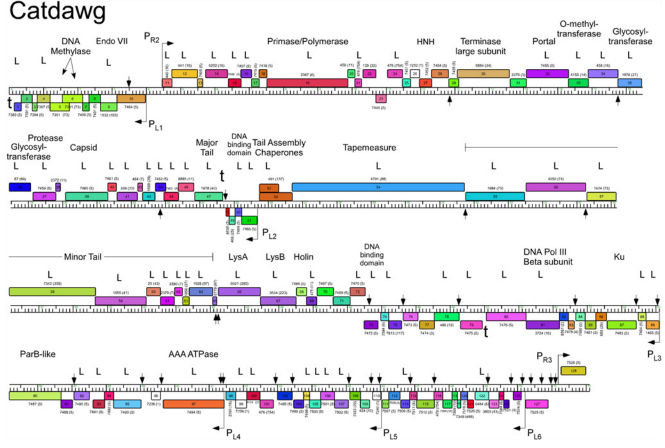
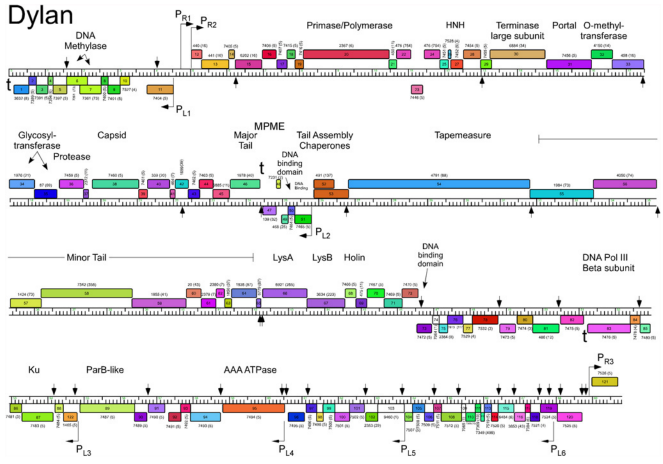


Fig 4. Genome map of Mycobacteriophage Catdawg. The genome of phage Catdawg is represented as a scale bar (major intervals: 1 kbp) with predicted genes shown as boxes either above (rightwards transcribed) or below (leftwards transcribed). Gene number is shown within each box and the phamily designation is shown either above or below with the number of phamily members shown in parentheses. Putative gene functions are indicated. The positions of putative SigA-like promoters ( $P_{L1}$ — $P_{L6}$  and  $P_{R1}$ — $P_{R3}$ ) are shown as large arrows. Small vertical arrows show the locations of the palindromic repeat 5′-TGTTCGGNNNCCGAACA. Catdawg proteins identified in a phage lysate using LC-MS/MS with at least two high confidence peptides per product are indicated (L).

### Predicted gene expression elements

The prediction of mycobacteriophage promoter locations is complicated because while some are related to mycobacterial SigA promoters [14–16], others appear not to be [17]. However, all five Cluster O phages contain at least eight strongly predicted SigA-like promoters, two rightwards facing ( $P_{R2}$ — $P_{R3}$ ) and six facing leftwards ( $P_{L1}$ — $P_{L6}$ ); Corndog, Dylan, and Yung-Jamal have an additional rightwards-facing promoter ( $P_{R1}$ ) upstream of  $P_{R2}$ .  $P_{L1}$  and  $P_{R2}$  transcribe divergently from the intergenic region located ~5 kbp from the left end and both are predicted to express leaderless mRNAs with the transcription +1 site coinciding with the first base of the first codon of the downstream gene. These intergenic regions are generally much more AT-rich than the rest of the genomes. Promoter  $P_{L2}$  that transcribes the leftward facing gene in the structural operon is similarly organized with respect to the start codon of the downstream gene (e.g. Corndog 53). Four leftwards promoters are situated within the long span of leftwards transcribed genes at the right side of the genomes, suggesting that these constitute at least four separate operons;  $P_{L6}$  is within coding regions (e.g. Corndog 120) but is strongly predicted (5′-TGTCAA—17 bp—TAGAAT).





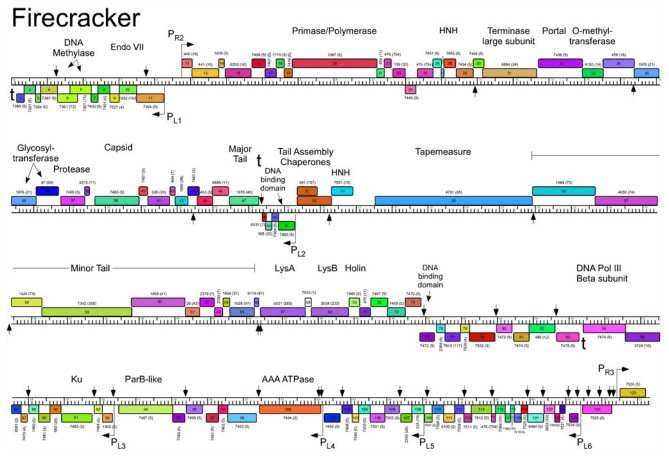
**Fig 5. Genome map of Mycobacteriophage Dylan.** The genome of phage Dylan is represented as a scale bar (major intervals: 1 kbp) with predicted genes shown as boxes either above (rightwards transcribed) or below (leftwards transcribed). Gene number is shown within each box and the phamily designation is shown either above or below with the number of phamily members shown in parentheses. Putative gene functions are indicated. The positions of putative SigA-like promoters (P<sub>L1</sub>—P<sub>L6</sub> and P<sub>R1</sub>—P<sub>R3</sub>) are shown as large arrows. Small vertical arrows show the locations of the palindromic repeat 5′-TGTTCGGNNNCCGAACA.

The Cluster O genomes have three motifs with the potential to form stem-loop RNA structures that play roles in modulating transcription [18]. The first is located at the extreme left ends of the genomes (Corndog coordinates 62–101) such as to terminate leftwards transcription. It contains a 13 bp stem-loop (with a 1 bp bulge) followed by 5'-TTTGT. The second is to the right of the major tail subunit gene (e.g. Corndog 49; coordinates 25166–25195) and has a 12 bp stem (with a 1 bp bulge), is followed by 5'-TTTCT and likely acts as terminator of rightwards transcription. The third is located between Corndog genes 83 and 84 (Corndog coordinates 51076–51107) and forms a predicted RNA structure with an 18 bp stem and an associated T-rich region that could act as a terminator of leftwards transcription.

### A conserved repeated sequence in Cluster O mycobacteriophages

The dot plot genome comparison (Fig. 1) suggests the presence of a small repeated sequence present many times in each of the Cluster O genomes. The conserved 17 bp sequence contains a 7bp inverted repeat separated by 3 bp (5'-TGTTCGGNNNCCGAACA) and is present 34 times in Corndog (Fig. 8) and similarly in the other Cluster O phages. The inverted repeat sequences are invariant among the 34 Corndog sites (there are three additional sites varying at





**Fig 6. Genome map of Mycobacteriophage Firecracker.** The genome of phage Firecracker is represented as a scale bar (major intervals: 1 kbp) with predicted genes shown as boxes either above (rightwards transcribed) or below (leftwards transcribed). Gene number is shown within each box and the phamily designation is shown either above or below with the number of phamily members shown in parentheses. Putative gene functions are indicated. The positions of putative SigA-like promoters ( $P_{L1}$ — $P_{L6}$  and  $P_{R1}$ — $P_{R3}$ ) are shown as large arrows. Small vertical arrows show the locations of the palindromic repeat 5′-TGTTCGGNNNCCGAACA.

one position), and although there is variation in the central three nucleotides, 5'-TTT (or 5'-AAA) is the most common, present in 29 of the 34 sites (Fig. 8). However, there is little evidence to support meaningful site orientation based on the central trinucleotide asymmetry, at least with regards to the direction of transcription; for example, of the 23 sites within the leftwards operon at the genome right end—Corndog genes 76–121–14 have 5'-TTT and 6 have 5'-AAA on the top strand (Fig. 8).

Most of the sites are in similar positions in all five genomes, although there are informative departures of two types. First, there are several instances where there is apparent loss of a site because of a single base change in one of the repeats. One example is a site in Corndog, Dylan, Firecracker and YungJamal immediately to the left of the methylase genes (e.g. Corndog 6; Fig. 3), which in Catdawg, has a single base change in the lefthand 7 bp segment. The change is non synonymous for the downstream gene (e.g. Corndog 5), and the sequence diverges downstream of it. A second example is the loss of a site in Catdawg in the 3' end of the larger tail chaperone gene (e.g. Catdawg 53, Fig. 4) because of a change at one position that is synonymous for the reading frame. A second type of departure is where recombination between sites appears to have contributed to insertions or deletions. One example is the presence of a ~550 bp segment between Catdawg genes 95 and 97 that is flanked by two of the repeats. In the other



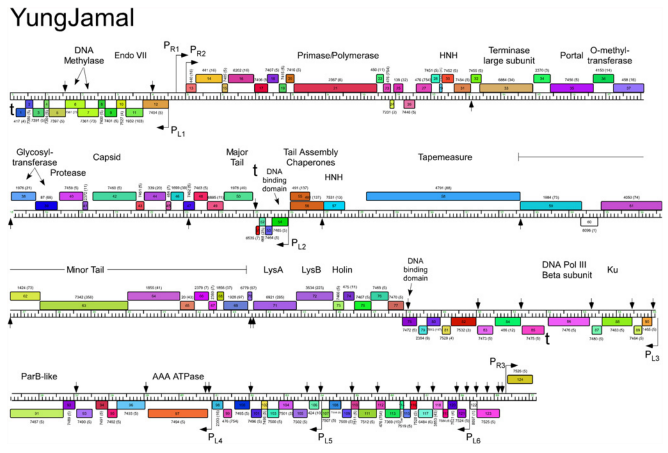


Fig 7. Genome map of Mycobacteriophage YungJamal. The genome of phage YungJamal is represented as a scale bar (major intervals: 1 kbp) with predicted genes shown as boxes either above (rightwards transcribed) or below (leftwards transcribed). Gene number is shown within each box and the phamily designation is shown either above or below with the number of phamily members shown in parentheses. Putative gene functions are indicated. The positions of putative SigA-like promoters ( $P_{L1}$ — $P_{L6}$  and  $P_{R1}$ — $P_{R3}$ ) are shown as large arrows. Small vertical arrows show the locations of the palindromic repeat 5'-TGTTCGGNNNCCGAACA.

four genomes there is only a single copy of the repeat, and a simple explanation is that Catdawg represents the ancestral state with the other genomes having a deletion resulting from recombination between the two repeats. In a second example, the region immediately downstream of the  $P_{L6}$  promoter in Corndog appears to represent the ancestral state with all other genomes having a deletion created by recombination between the two Corndog repeats immediately downstream of  $P_{L6}$ .

Fourteen of the Corndog repeats are within short intergenic regions and several others are close to the 5' end of the coding region and the annotated start site choice has yet to be confirmed (see below; Fig. 8). Eleven of the sites are clearly within coding regions (in Corndog genes 12, 36, 46, 55, 68, 76, 108, 111, 117, 120, and 121). However, the intergenic sites are not randomly distributed across the genome, and they are predominantly (11 of 14 in Corndog) in the leftwards-transcribed region of Corndog genes 76–121 (Fig. 3). The site symmetry suggests these represent binding sites for dimeric regulatory proteins, and we note there are three predicted DNA binding proteins encoded in each of the genomes (e.g. Corndog gp53, gp76, and gp90). However, the possible regulatory consequences are not clear. Although four of the sites are near predicted promoters, most are not, and a transcriptional regulatory function for these repeats seems unlikely. The site is not present in *M. smegmatis* mc²155 or *M. tuberculosis* 



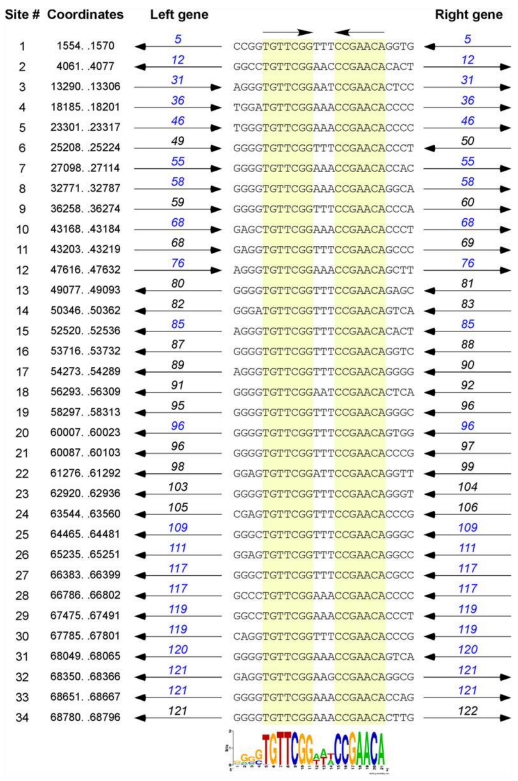


Fig 8. Conserved repeats sequences in the Corndog genome. The Corndog genome contains multiple repeats of a 17 bp sequence composed of two 7 bp inverted motifs separated by three base pairs. The 34 sites are aligned, showing the top strand (and flanking 4 bp) with the 7 bp motifs are highlighted in yellow; the coordinates shown correspond to the 17 bp sequence. The genes flanking the repeat (black) or the genes containing the repeat (blue) and their directions of transcription are shown. Fourteen of the 34 sites (# 6, 9, 11, 13, 14, 16, 17, 18, 19, 21, 22, 23, 24, and 34) are located between open reading frames, ten (#1, 3, 7, 8, 15, 20, 28, 29, 31, and 33) are within open reading frames but close to the 5' end of the gene (and could be intergenic if the start site is



not correctly identified), and ten (#2, 4, 5, 10, 12, 25, 26, 27, 30, and 32) are in the middle or towards the 3' ends of genes (and the gene is not shown). An additional three sites containing a single base change are not shown. The weblogo at the bottom shows alignment of all 34 sites and related sites identified by MEME [19]; both orientations are compiled due to the inverted repeat such that the flanking 4 bp is shown only on the left. Note that the central three nucleotide spacer is A/T rich, with the most common sequence being AAA or TTT (29 of the 34 sites). There is a slight preference for the orientation of the site to be such that the AAA is on the top strand when the site is transcribed in the rightwards direction. The flanking four nucleotides are G/C rich.

doi:10.1371/journal.pone.0118725.g008

genomes, or the genomes of other mycobacteriophages; there are two copies in *Mycobacterium*  $sp\ 05'1390\ [\underline{20}]$ .

# Identification of Cluster O phage proteins by SDS-PAGE and mass spectrometry

SDS-PAGE analysis of Corndog virion proteins shows a prominent band of 40 kDa and at least six minor proteins (Fig. 2B). Further analysis of CsCl-purified (twice banded) Corndog virions by LC-MS/MS identified twenty-one proteins with high confidence (≥2 peptides/protein Fig. 3, Table 3). All of these are encoded by genes in the interval 34–67 with the exception of gp13 (Fig. 3) and include the capsid (gp41) and major tail subunits (gp49), portal (gp34), protease (gp39), putative tail capping and head-tail connector proteins (gp42, gp43, gp45, gp47), tapemeasure protein (gp57) and minor tail proteins (gp58—gp67), as well as gp52 which is of unknown function and transcribed opposite to the other virion genes (Fig. 3). We note that other proteins encoded within this region including the O-methyltransferase (gp35), the glycosyltransferases (gp36, gp37) and the N-acetylglucosaminyltransferase (gp38) were not identified in the virions. LC-MS/MS of Corndog particles purified through a single round of CsCl banding identified all of the same proteins and another 36 Corndog-encoded proteins that are presumably contaminants from lysed cells (Table 3). For an additional four proteins (gp11, gp3, gp77, and gp102) we identified multiple spectra (2, 2, 2, and 4 respectively) but only from a single unique peptide each. We also analyzed extracts of Corndog-infected cells by LC-MS/ MS and identified an additional three gene products (gp90, gp96, and gp122) not found in the other samples (Fig. 3, Table 3). The proportion of predicted products identified by LC-MS/MS (48%) is somewhat lower than for similar experiments with mycobacteriophage Patience (75%) [21]. We also analyzed an unpurified lysate of Catdawg by LC-MS/MS using both chymotrypsin and trypsin cleavage (Table 4). A total of 63 proteins were identified (49% of total predicted), with a profile that is similar but not identical to the Corndog proteins.

The LC-MS/MS analysis unfortunately provides few clues as to the basis of the prolate capsids of the Cluster O phages. The capsid subunits (Corndog gp41) are predicted to be structurally similar to the isometric HK97 capsid subunit by HHPred [22] analysis, including the Nterminal 102-residue delta domain that is cleaved and lost during capsid maturation [23, 24]. The LC-MS/MS analysis reveals very few Corndog capsid subunit peptides from either purified particles or late-infected cells, perhaps reflecting poor trypsin digestion of the high molecular weight covalently crosslinked protein seen by SDS-PAGE (Fig. 2B), as seen in HK97 [25]. However, two of the six Corndog virion capsid peptide spectra identified correspond to the delta domain suggesting that it may remain during capsid maturation. Poor recovery of capsid peptides could also result from modifications whose masses are not readily predictable—such as complex sugar additions—and escape LC-MS/MS deconvolution. Major capsid subunit peptides were well-represented in the Catdawg sample, but many of these could have come from unassembled procapsids. We note that six Corndog proteins (gp5, gp17, gp52, gp59, gp61) and five Catdawg proteins (gp14, gp33, gp46, gp56 and gp58) have N-terminally acetylated peptides all at a threonine encoded by the second codon. The functional consequences of this—if any are not known.



Table 3. Corndog peptides identified by mass-spectrometry.

| Coordinates | Product/ Function             | Corndog Particles <sup>1</sup> |         | Infected cells | Total Peptides <sup>2</sup> | Start site Confirmed <sup>3</sup> |
|-------------|-------------------------------|--------------------------------|---------|----------------|-----------------------------|-----------------------------------|
|             |                               | 1x CsCl                        | 2x CsCl |                |                             |                                   |
| 28380–32765 | gp 57 tapemeasure             | 1266                           | 93      | 100            | 1459                        | See text                          |
| 36294–37142 | gp 60 minor tail protein      | 506                            | 38      | 32             | 576                         | Confirmed                         |
| 22037–22642 | gp 43                         | 444                            | 63      | 60             | 567                         | Confirmed                         |
| 32803–34521 | gp 58 minor tail protein      | 525                            | 26      | 13             | 564                         | Reassigned                        |
| 37139–39649 | gp 61 minor tail protein      | 328                            | 61      | 59             | 448                         | Confirmed, acetyl                 |
| 15549–16778 | gp 34 portal                  | 358                            | 35      | 49             | 442                         | Insufficient data                 |
| 39642–41132 | gp 62 minor tail protein      | 280                            | 35      | 28             | 343                         | Confirmed                         |
| 25493–25684 | gp 52                         | 175                            | 35      | 49             | 259                         | Confirmed, acetyl                 |
| 19596–20261 | gp 39 capsid mat. protease    | 209                            | 22      | 23             | 254                         | Consistent                        |
| 24316–25131 | gp 49 major tail              | 174                            | 43      | 19             | 236                         | Confirmed                         |
| 34518–36254 | gp 59 minor tail protein      | 187                            | 23      | 5              | 215                         | Confirmed, acetyl                 |
| 41144–41545 | gp 63 minor tail protein      | 144                            | 17      | 12             | 173                         | Confirmed                         |
| 41960–42184 | gp 65                         | 104                            | 3       | 0              | 107                         | Confirmed                         |
| 41549–41950 | gp 64 minor tail protein      | 89                             | 9       | 8              | 106                         | Confirmed                         |
| 23417–23842 | gp 47                         | 54                             | 5       | 10             | 69                          | Insufficient data                 |
| 56866–57207 | gp 93                         | 52                             | 0       | 15             | 67                          | Confirmed                         |
| 20547–21779 | gp 41 major capsid            | 54                             | 6       | 5              | 65                          | Confirmed                         |
| 21779–22027 | gp 42                         | 56                             | 5       | 0              | 61                          | Confirmed                         |
| 42385–43071 | gp 67                         | 34                             | 5       | 1              | 40                          | Confirmed                         |
| 5024–5311   | gp 13                         | 31                             | 2       | 0              | 33                          | Confirmed                         |
| 42197–42385 | gp 66                         | 25                             | 3       | 0              | 28                          | Processed?                        |
| 22796–23155 | gp 45                         | 15                             | 5       | 6              | 26                          | Insufficient data                 |
| 17352–18221 | gp 36                         | 22                             | 0       | 0              | 22                          | Confirmed                         |
| 1111–1581   | gp 5                          | 19                             | 0       | 0              | 19                          | Confirmed, acetyl                 |
| 60504–60770 | gp 124                        | 18                             | 0       | 0              | 18                          | Confirmed                         |
| 26207–26761 | gp 54 tail assembly chaperone | 18                             | 0       | 0              | 18                          | Insufficient data                 |
| 11715–12149 | gp 27                         | 16                             | 0       | 0              | 16                          | Consistent                        |
| 12449–12778 | gp 29                         | 16                             | 0       | 0              | 16                          | Consistent                        |
| 6409–7074   | gp 17                         | 15                             | 0       | 0              | 15                          | Confirmed, acetyl                 |
| 23835–24287 | gp 48                         | 10                             | 1       | 4              | 15                          | Insufficient data                 |
| 53972–54247 | gp 89                         | 13                             | 0       | 0              | 13                          | Confirmed                         |
| 57197–57469 | gp 94                         | 12                             | 1       | 0              | 13                          | Reassigned                        |
|             | gp 46                         | 12                             | 0       | 0              | 12                          | Insufficient data                 |
| 23152–23472 |                               | 11                             | 0       |                |                             | Insufficient data                 |
| 18907–19527 | gp 38 glycosyltransferase     |                                | 0       | 0<br>8         | 11<br>11                    | Insufficient data                 |
| 51112–52290 | gp 84 DNA pol Beta subunit    | 3                              |         |                |                             |                                   |
| 18218–18910 | gp 37 glycosyltransferase     | 10                             | 0       | 0              | 10                          | Insufficient data                 |
| 25320–25493 | gp 51                         | 10                             | 0       | 0              | 10                          | Insufficient data                 |
| 3790–4524   | gp 12                         | 9                              | 0       | 0              | 9                           | Consistent                        |
| 7098–7472   | gp 18                         | 9                              | 0       | 0              | 9                           | Insufficient data                 |
| 971–1111    | gp 4                          | 8                              | 0       | 1              | 9                           | Confirmed                         |
| 53737–53964 | gp 88                         | 8                              | 1       | 0              | 9                           | Insufficient data                 |
| 11362–11653 | gp 26                         | 7                              | 0       | 0              | 7                           | Reassigned                        |
| 62493–62897 | gp 103                        | 6                              | 0       | 0              | 6                           | Insufficient data                 |
| 64188–64442 | gp 109                        | 6                              | 0       | 0              | 6                           | Confirmed                         |
| 52540-52812 | gp 86                         | 5                              | 1       | 0              | 6                           | Confirmed                         |

(Continued)



Table 3. (Continued)

| Coordinates | Product/ Function         | Corndog Particles <sup>1</sup> |         | Infected cells | Total Peptides <sup>2</sup> | Start site Confirmed <sup>3</sup> |
|-------------|---------------------------|--------------------------------|---------|----------------|-----------------------------|-----------------------------------|
|             |                           | 1x CsCl                        | 2x CsCl |                |                             |                                   |
| 57466–58266 | gp 95                     | 5                              | 0       | 1              | 6                           | Insufficient data                 |
| 61585–61767 | gp 100                    | 5                              | 0       | 0              | 5                           | Insufficient data                 |
| 49097–49528 | gp 81                     | 5                              | 0       | 0              | 5                           | Insufficient data                 |
| 50364-50996 | gp 83                     | 5                              | 0       | 0              | 5                           | Insufficient data                 |
| 16775–17359 | gp 35 O-methyltransferase | 4                              | 0       | 0              | 4                           | Confirmed                         |
| 52318-52539 | gp 85                     | 4                              | 0       | 0              | 4                           | Insufficient data                 |
| 8227-10587  | gp 22                     | 3                              | 0       | 0              | 3                           | Insufficient data                 |
| 10777-10983 | gp 24                     | 3                              | 0       | 0              | 3                           | Insufficient data                 |
| 54424-55932 | gp 90 ParB-like           | 0                              | 0       | 3              | 3                           | Insufficient data                 |
| 55929–56267 | gp 91                     | 3                              | 0       | 0              | 3                           | Insufficient data                 |
| 58355-60055 | gp 96 AAA-ATpase          | 0                              | 0       | 3              | 3                           | Insufficient data                 |
| 68942-69664 | gp 122                    | 0                              | 0       | 2              | 2                           | Insufficient data                 |
| 439–651     | gp 2                      | 2                              | 0       | 0              | 2                           | Insufficient data                 |
| 27158–27757 | gp 56 HNH endonuclease    | 2                              | 0       | 0              | 2                           | Confirmed                         |
| 2707–3042   | gp 9                      | 2                              | 0       | 0              | 2                           | Confirmed                         |

<sup>&</sup>lt;sup>1</sup>Corndog virion particles were purified through one (1x) or two (2x) CsCl equilibrium density gradients.

In general, the LC-MS/MS analysis provides information about the translational start sites, and for 26 Corndog genes the annotated start site is confirmed (Table 3), and in 4 others the data is consistent with the predicted start but does not discern between the predicted start site and other possible start sites. For three genes (Corndog 26, 58, and 94) the LC-MS/MS data support re-annotation of the start sites (to positions 11,653, 32,803, and 57,469 respectively; Table 3). For one protein, Corndog gp66, 28 peptide spectra were obtained, but all correspond to the C-terminal 34 residues of the predicted 62-residue product suggesting that it may be post-translationally processed (Table 3). For its Catdawg homologue (gp63), 58 spectra were recovered all of which—with one exception that could be derived from an uncleaved precursor—are in the same C-terminal moiety. We also identified peptides for a previously unannotated Corndog gene (124) encoded between genes Corndog 97 and 98 (Table 3).

LC-MS/MS data confirms annotated start sites for 26 Catdawg genes and in nine others the data is consistent with the predicted start does but does not discern between the predicted start site and other possible start sites (<u>Table 4</u>). For one gene (Catdawg *122*) the LC-MS/MS data support re-annotation of the start site to position 69163 (<u>Table 4</u>).

Alignment of the Cluster O genome maps (S1 Fig., Figs. 3–7) shows an evident disparity in the annotation of the tape measure protein (*tmp*) genes. In Catdawg and Dylan the predicted translational start site overlaps the termination codon of the upstream tail assembly chaperone gene, and the LC-MS/MS data are consistent with the annotated Catdawg tmp start site (Table 4). However, in Corndog, Firecracker, and YungJamal, an HNH gene is inserted between the tail assembly chaperone and *tmp*, resulting in *tmp* being annotated to begin at the first available start codon ~ 600 bp downstream, leaving a non-coding gap (Fig. 9A). However, LC-MS/MS of Corndog proteins identified many peptide spectra corresponding to the upstream region of the *tmp* ORF indicating that translation begins upstream. The most N-

<sup>&</sup>lt;sup>2</sup>Table is sorted by total number of peptides assigned by stringent criteria. See text for details and thresholds.

<sup>&</sup>lt;sup>3</sup>Translation start sites are indicated as confirmed, consistent with the annotation, warranted reassignment of the start site (shown in coordinates), or insufficient data to confirm; acetyl, if more than 50% N-terminal peptides acetylated.



Table 4. Identification of Catdawg proteins by mass spectrometry.

| Coordinates    | Product/Function                  | Chymotrypsin | Trypsin | Total Peptides <sup>1</sup> | Start site confirmed <sup>2</sup> |
|----------------|-----------------------------------|--------------|---------|-----------------------------|-----------------------------------|
| 23858 to 24673 | gp47 major tail                   | 2516         | 3056    | 5572                        | Confirmed                         |
| 26700 to 31724 | gp54 tape measure                 | 1798         | 1662    | 3460                        | Consistent                        |
| 20089 to 21321 | gp39 major capsid                 | 1634         | 510     | 2144                        | Confirmed                         |
| 31762 to 33480 | gp55 minor tail protein           | 935          | 963     | 1898                        | Confirmed                         |
| 15091 to 16320 | gp32 portal                       | 1039         | 1134    | 2173                        | Consistent                        |
| 21594 to 22184 | gp41                              | 641          | 1145    | 1786                        | Confirmed                         |
| 33524 to 35260 | gp56 minor tail protein           | 454          | 763     | 1217                        | Confirmed, acetyl                 |
| 35300 to 36148 | gp57 minor tail protein           | 586          | 628     | 1214                        | Confirmed                         |
| 36145 to 38655 | gp58 minor tail protein           | 637          | 399     | 1036                        | Confirmed, acetyl                 |
| 38648 to 40138 | gp59 D-ala-D-ala-carboxypeptidase | 363          | 424     | 787                         | Confirmed                         |
| 41391 to 42077 | gp64                              | 359          | 218     | 577                         | Confirmed                         |
| 22338 to 22697 | gp43                              | 170          | 124     | 294                         | Confirmed                         |
| 40150 to 40551 | gp60                              | 164          | 170     | 334                         | Confirmed                         |
| 19138 to 19803 | gp37 capsid maturation protease   | 102          | 266     | 368                         | Consistent                        |
| 23377 to 23829 | gp46                              | 78           | 139     | 217                         | Confirmed, acetyl                 |
| 40555 to 40956 | gp61                              | 124          | 180     | 304                         | Confirmed                         |
| 22959 to 23384 | gp45                              | 113          | 149     | 262                         | Insufficient data                 |
| 42246 to 43466 | gp66 lysA                         | 44           | 198     | 242                         | Confirmed                         |
| 41203 to 41391 | gp63                              | 20           | 38      | 58                          | Insufficient data                 |
| 40966 to 41190 | gp62                              | 3            | 66      | 69                          | Confirmed                         |
| 6395 to 6769   | gp15                              | 3            | 48      | 51                          | Insufficient data                 |
| 4747 to 5496   | gp12                              |              | 42      | 42                          | Insufficient data                 |
| 25226 to 25035 | gp50                              | 8            | 39      | 47                          | Insufficient data                 |
| 5733 to 6371   | gp14                              | 6            | 28      | 34                          | Confirmed, acetyl                 |
| 16894 to 17763 | gp34 glycosyltransferase          | -            | 34      | 34                          | Insufficient data                 |
| 4467 to 4754   | gp11                              | 3            | 18      | 21                          | Confirmed                         |
| 52190 to 51225 | gp81                              | 2            | 29      | 31                          | Consistent                        |
| 358 to 152     | gp1                               | 9            | 8       | 17                          | Confirmed                         |
| 1198 to 821    | gp4                               | 2            | 28      | 30                          | Insufficient data                 |
| 7521 to 9884   | gp19 DNA primase/polymerase       |              | 26      | 26                          | Insufficient data                 |
| 46085 to 46525 | gp72                              |              | 18      | 18                          | Insufficient data                 |
| 43468 to 44508 | gp67 LysB                         |              | 19      | 19                          | Consistent                        |
| 46901 to 46494 | gp73 HTH DNA binding protein      |              | 18      | 18                          | Confirmed                         |
| 25678 to 25223 | gp51                              |              | 21      | 21                          | Insufficient data                 |
| 11045 to 11479 | gp24                              |              | 23      | 23                          | Consistent                        |
| 64667 to 64260 | gp106                             | 2            | 8       | 10                          | Consistent                        |
| 65071 to 64667 | gp107                             | _            | 10      | 10                          | Confirmed                         |
| 66856 to 66599 | gp113                             |              | 15      | 15                          | Confirmed                         |
| 61447 to 59669 | gp97 AAA ATPase                   |              | 3       | 3                           | Insufficient data                 |
| 54718 to 54491 | gp88                              |              | 8       | 8                           | Confirmed                         |
| 22709 to 23014 | gp44                              |              | 10      | 10                          | Insufficient data                 |
| 16317 to 16901 | gp33 O-methyl transferase         | 5            | 6       | 11                          | Confirmed, acetyl                 |
| 59020 to 58220 | gp95                              | 3            | 9       | 9                           | Insufficient data                 |
| 18449 to 19069 | <del></del>                       | 3            | 3       | 5                           | Insufficient data                 |
| 55100 to 54726 | gp36<br>gp89                      | 3            | 5       | 8                           | Insufficient data                 |
|                |                                   |              |         |                             |                                   |
| 49941 to 49309 | gp79                              |              | 8       | 12                          | Confirmed                         |

(Continued)



Table 4. (Continued)

| Coordinates    | Product/Function              | Chymotrypsin | Trypsin | Total Peptides <sup>1</sup> | Start site confirmed <sup>2</sup> |
|----------------|-------------------------------|--------------|---------|-----------------------------|-----------------------------------|
| 44814 to 45113 | gp69                          |              | 12      | 6                           | Insufficient data                 |
| 69163 to 68768 | gp122                         |              | 6       | 2                           | Reassigned                        |
| 57514 to 57065 | gp92                          | 2            |         | 5                           | Insufficient data                 |
| 62494 to 62105 | gp100                         |              | 5       | 4                           | Insufficient data                 |
| 17760 to 18452 | gp35 glycosyltransferase      |              | 4       | 7                           | Insufficient data                 |
| 47213 to 46980 | gp74                          | 7            | 4       | Confirmed                   |                                   |
| 57961 to 57620 | gp93                          | 2            | 2       | 4                           | Insufficient data                 |
| 62098 to 61778 | gp99                          |              | 4       | 6                           | Insufficient data                 |
| 64263 to 63934 | gp105                         |              | 6       | 5                           | Consistent                        |
| 25749 to 26303 | gp52 tail assembly chaperone  |              | 5       | 6                           | Insufficient data                 |
| 21321 to 21569 | gp40                          |              | 6       | 3                           | Insufficient data                 |
| 3150 to 2665   | gp9 Endo VII protein          |              | 3       | 2                           | Insufficient data                 |
| 51220 to 50057 | gp80 DNA pol III beta subunit |              | 2       | 3                           | Insufficient data                 |
| 10086 to 10280 | gp21                          |              | 3       | 2                           | Insufficient data                 |
| 6762 to 7073   | gp16                          |              | 2       | 4                           | Confirmed                         |
| 49266 to 48541 | gp78                          |              | 4       | 2                           | Insufficient data                 |
| 63941 to 63762 | gp104                         |              | 2       | 2                           | Consistent                        |

<sup>&</sup>lt;sup>1</sup>Table is sorted by total number of peptides assigned by stringent criteria. See text for details and thresholds.

terminal peptides have the sequence N-AIHIDIYAHLQK and are not generated by tryptic digestion. There are no canonical translation start sites between these peptides and the most upstream termination codon (Fig. 9A), and the threonine codon immediately upstream of this peptide (5'-ACG) is in the corresponding position to the tmp 5'-ATG start codon in Dylan and Catdawg (Fig. 9A). We have been unable to identify any RNA-level splicing event that would suggest that the HNH gene is part of an intron (Fig. 9B) and the most likely possibilities are that either the 5'-ACG codon is used for translation initiation or that translation begins upstream and tmp translation involves a ribosome bypassing event [26]. We are not aware of any other mycobacterial genes initiating translation with ACG and attempts to sequence the tmp N-terminus by Edman degradation have failed, presumably due to modification; the five N-terminal residues from another protein (gp43) from the same gel were readily determined.

# Mobile Elements in Cluster O phages

We noted previously that Corndog contains a truncated version of a Mycobacteriophage Mobile Element (MPME) (encoding Corndog gp25) found in phage genomes within an assortment of clusters (27). MPMEs are small (~440 bp) and include a 123-residue ORF, and two types (MPME1 and MPME2) have been described [27]. Phage YungJamal shares the same sequence as Corndog, which includes the left inverted repeat (IR-L) and 363 bp of MPME1, whereas Catdawg and Firecracker contain a similar segment of the MPME element but have different flanking sequences reflecting deletions of the Corndog sequence. Dylan does not contain an MPME fragment at this site but also does not simply correspond to a pre-integration site either, as there is a 20 bp separation between the Corndog/Dylan homology and IR-L rather than the typical 6 bp [27].

<sup>&</sup>lt;sup>2</sup>Translation start sites are indicated as confirmed, consistent with the annotation, warranted reassignment of the start site (shown in coordinates), or insufficient data to confirm; acetyl, if more than 50% N-terminal peptides acetylated.



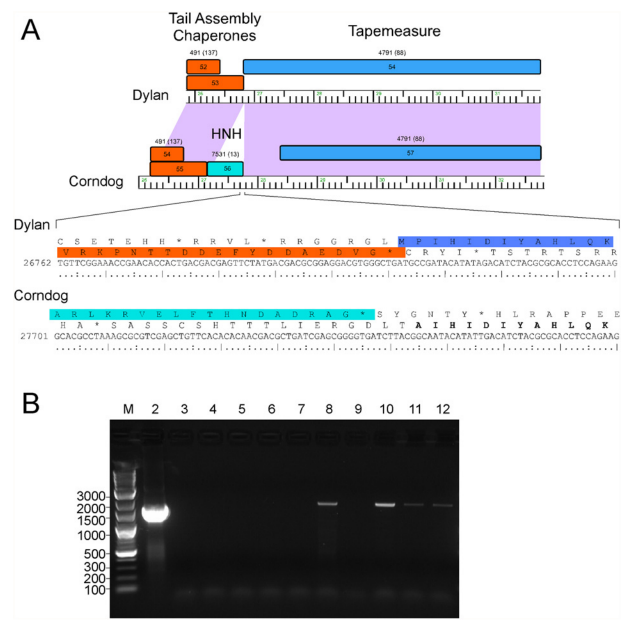


Fig 9. Unusual translation initiation of the Corndog tape measure protein gene. A. Two organizations of the tape measure genes are present in the Cluster O phages. In Dylan and Catdawg the *tmp* gene is predicted to start translation immediately downstream of the tail assembly chaperone genes that are translated via a programmed translational frameshift. In contrast, Corndog, Firecracker, and YungJamal have a non-coding gap prior to the *tmp* start site. However, LC-MS/MS identified Corndog peptides corresponding to this gap and the sequence of the most N-terminal peptides are shown in bold type. Translation presumably either initiates at the ACG threonine codon or starts further upstream and involves a ribosome bypass event. B. RT-PCR of Corndog transcripts. PCR products were generated using a Corndog lysate (lane 2) or RNA isolated from uninfected cells (lanes 3 and 4), or at different times after infected by Corndog: 30 min (lanes 5 and 6), 2.5 h (lanes 7 and 8), 3.5 h (lanes 9 and 10), and 4.5 h (lanes 11 and 12. Lanes 3, 5, 7, 9, and 11 are controls lacking reverse transcriptase. A DNA ladder marker (M) is shown with sizes in base pairs. Genomic DNA and unspliced RNAs generate an expected product of ~1.7 kbp. No smaller spliced products are observed.

Interestingly, Dylan contains a complete MPME element inserted to the right of the major tail subunit gene, and oriented in the opposite direction (i.e. IR-R proximal to the major tail subunit gene; Fig. 5). This MPME element is an apparent hybrid between MPME1 and MPME2 sequences presumably generated by recombination such that the 5' half corresponds



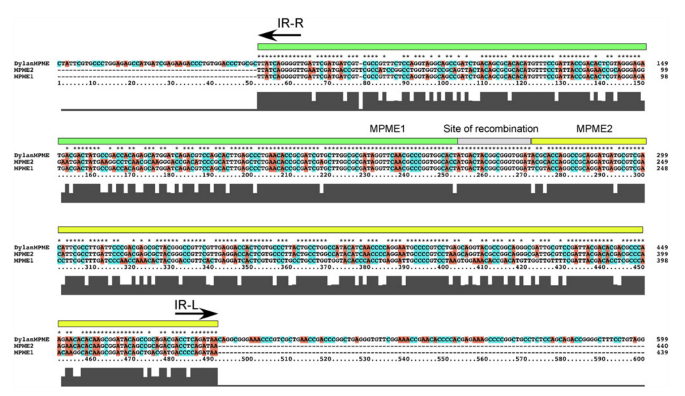


Fig 10. Dylan MPME element. Phage Dylan contains a Mycobacteriophage Mobile Element (MPME) inserted between genes 46 and 48. The Dylan MPME contains an open reading frame (47) that is transcribed leftwards, such that the MPME left inverted repeat (IR-R) is 48-proximal. Alignment of the Dylan MMPE sequence with MPME1 and MPMP2 [27] shows that one half (green box) is identical to MPME1 and the other half (yellow box) is identical to MPME2. The Dylan MPME is thus a hybrid of MPME1 and MPME2, presumably generated by homologous recombination with the intervening sequence (grey box).

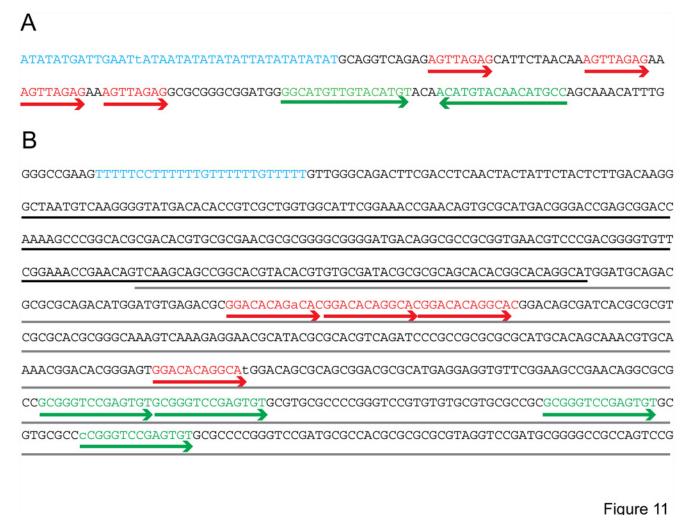
to MPME1 and the 3' half to MPME2 (Fig. 10). The IR-L of this MPME element (at coordinate 24957) is separated by 6 bp from sequence identity in Corndog (coordinate 25300) and the other phages, indicating this to be the site of the insertion. At the opposite end, there are 14 bp between IR-R and the shared sequences suggesting either differences in the pre-integration site or rearrangements associated with the insertion.

All five Cluster O genomes contain a homing endonuclease-like gene (HNH) gene upstream from the terminase (e.g. Corndog 29) implicated in DNA packaging [28], and two additional HNHs are present in subsets of the genomes. One of these corresponds to the insertion upstream of the tape measure protein gene as discussed above; the other is present in three of the genomes (Corndog, Catdawg, YungJamal) located downstream of the large terminase subunit gene (e.g. Corndog 33). Dylan and Firecracker lack this HNH gene and comparisons suggest a simple insertion 1–3 bp downstream of the terminase stop codon.

### Other features of Cluster O genomes

There are several other notable features of the Cluster O genomes. First, at the left ends of the genomes there are two adjacent leftwards-transcribed genes coding for domains of cytosine methyltransferases (Corndog genes 6 and 7 and their relatives). Corndog gp7 has a strong HHPred match to the N-terminal part of *Hae*III methylase as well as BLASTP matches to other methylases (including those not encoded by mycobacteriophages) extending across the entire protein span of gp7 (~195 residues) to within a few residues of the gp7 C-terminus. The 53 C-terminal residues of Corndog gp6 (and relatives) are predicted strongly by HHPred to





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**Fig 11. Sequence features of Cluster O genomes. A.** The AT rich element between Corndog genes *12* and *13* is highlighted in cyan, and two sets of flanking sequence repeats are shown in red and green. A similar arrangement of these sequences is observed in the other Cluster O phages. Residues in these sequence elements that differ across the phages (in the case of the AT rich element), or from the repeat consensus sequences are shown in lower case. **B.** A portion of Corndog genes *120* (underlined in black) and *121* (underlined in gray). The conserved T<sub>5</sub>CCT<sub>6</sub>GT<sub>6</sub>GT<sub>5</sub> sequence is shown in cyan and flanking sequence repeats are shown in green and red. Residues in these sequence elements that differ across the phages (in the case of the T rich element), or from the repeat consensus sequences are shown in lower case.

correspond to the three C-terminal alpha helices of *Hae*III methylase. However, the start site of gene 6 is ambiguous, and not only is it the strongest ribosome binding site associated with a start site located within the upstream (e.g. Corndog gene 7) open reading frame (Figs. 3–7), but there is also coding potential in the gene 6 frame in the overlap region, notwithstanding convincing conservation of the C-terminus of gp7 with numerous methylases. It is thus unclear whether two products are made that assemble to form a methylase active site—and if so where gp6 initiates from—or if a single product is expressed from a translational frameshift, a ribosome hop, or a spliced intron. However, RT-PCR analysis shows no evidence of splicing in this region (data not shown), and products of these genes were not identified by mass spectrometry. We note that similar arrangements of methylase gene segments are seen in other mycobacteriophages, and in phages of other hosts [29].



Secondly, the Cluster O phages encode several proteins with predicted transmembrane domains. Most contain only a single predicted membrane spanning domain and may not be membrane associated. However, downstream of the lysis cassette are two genes (e.g. Corndog 73 and 74) each encoding products with four predicted transmembrane domains that are strongly predicted to be membrane associated. Neither have relatives in other mycobacterio-phages, and their roles are unclear although they could also play a role in lysis.

The Cluster O genomes contain two AT-rich sequences, which are unusual among the GC rich mycobacteriophage genomes. The first, in the gap between the divergently transcribed operons on the lefthand side of the genome (i.e. Corndog genes 12 and 13) is a 39 nucleotide sequence consisting of 37 A or T residues that varies at only a single residue across the 5 Cluster O genomes. The second AT-rich sequence occurs at the far right hand side of the Cluster O genomes. In Corndog, this sequence lies within gene 120 whose central part is AT-rich and includes the sequence 5′-T<sub>5</sub>CCT<sub>6</sub>GT<sub>6</sub>GT<sub>5</sub>. Corndog 120 is poorly conserved among Cluster O genomes and we did not observe any peptides that could be encoded by this sequence in our MS data, raising the question of its assignment, but this AT-rich sequence is identical in all five phages. It is located 35 bp downstream of the putative P<sub>L6</sub> promoter and could play a regulatory role. Interestingly, a complex set of sequence repeats occurs to the right of each of these AT-rich elements (Fig. 11), and it is plausible that one or the other of these represents the phage origin of replication.

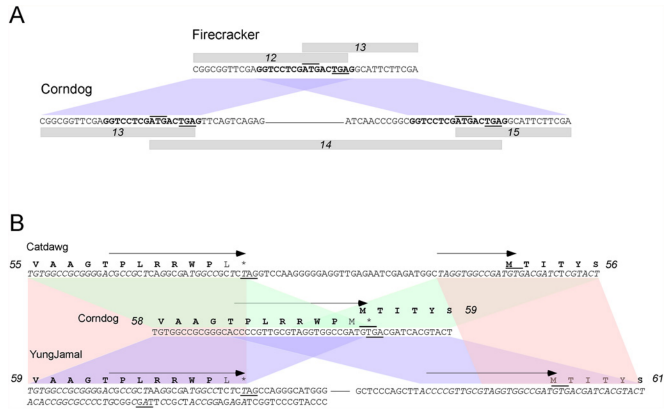
# Insights into phage genome evolution

Several regions of the Cluster O genomes differ in gene content as a consequence of deletions or insertions, typically by one or a small group of genes. These gene content differences occur in a variety of genomic contexts and apparently reflect relative recent horizontal exchange events rather than whole genome ancestries.

There are two examples of a gene present in one genome but absent from the other four genomes. Corndog gene 14 is small (126 bp) but HHPred analysis confidently predicts that gp14 folds similarly to the mycobacteriophage Pukovnik Xis protein [30] and is likely to be a DNA binding protein. Genome comparisons show that Corndog 14 is flanked by a 17 bp direct repeat present only once in the other genomes (Fig. 12A). Either Corndog represents the ancestral state from which gene 14 has been deleted by homologous recombination between the repeats, or Corndog has acquired 14 by recombination with a partner DNA carrying a sequence similar to the repeat.

A more complex relationship is seen with YungJamal gene 60, which is absent from the other four genomes. YungJamal 60 is transcribed in the leftwards direction, opposite to the tail genes that flank it and is of unknown function (Fig. 7). The gene is flanked by imperfect 24 bp direct repeats of which just 14 bp are conserved, and Corndog, Firecracker and Dylan each contain only a single copy of the repeat that is identical to the rightmost YungJamal copy (Fig. 12B). The base differences between the leftmost copy of the repeat in YungJamal and Corndog are such that the amino acid sequence of the products is maintained, with the exception of the C-terminal most residue (Fig. 12B). Catdawg differs from Corndog, Firecracker and Dylan in that it contains a small insertion including a partial second copy of the repeat. A plausible scenario is that Corndog, Firecracker and Dylan represent the ancestral state (and a canonical virion structural gene organization) into which YungJamal 60 was acquired by recombination, which subsequently underwent deletion to give the Catdawg structure. This then provides an evolutionary context for understanding the Catdawg genome that would not have been possible without the other Cluster O relatives.





**Fig 12. Insights into genome evolution. A.** Insertion of Corndog gene *14*. Cluster O genome comparisons show that Corndog gene *14* is missing from the other four related genomes. A 17 bp direct repeat (bold type) flanks Corndog gene *14* but is present only once at the junction of Firecracker gene *12* and *13* and their homologues in Dylan, Catdawg and YungJamal. Termination codons are underlined and translation start codons are overlined. Regions of nucleotide similarity are indicated by colored trapezoids. **B.** Insertion of gene *60* in YungJamal. YungJamal gene *60* encodes a protein of unknown function and is absent in all four other Cluster O genomes. YungJamal *60* is transcribed leftwards and is flanked by imperfectly conserved 24 bp inverted repeats (shown by arrows), but in which only 14 bp are conserved. However, Corndog (as well as Dylan and Firecracker) contains just a single copy of this repeat at the junction of genes Corndog *58* and *59*. Unusually the rightmost copy of the repeat in YungJamal (at the beginning of gene *61*) is identical to the Corndog sequence, whereas the leftmost repeat (at the end of gene *59*) is a degenerate copy in which most of the base changes are synonymous, except for the C-terminal residue. Termination codons are underlined and translation start codons are overlined; the sequences of both strands for the left component of YungJamal are indicated to show the termination codon (underlined) of YungJamal *60*. Catdawg lacks a homologue of YungJamal *60* but carries a small insertion relative to Corndog, Dylan, and Firecracker, and has part of the rightmost YungJamal repeat. Catdawg and YungJamal sequences shared with Corndog are shown in italic type. Sequences of nucleotide similarity are indicated by the colored trapezoids (Catdawg and Corndog, green; Corndog and YungJamal, purple; Catdawg and YungJamal, red).

Among the various other insertions and deletions, we note that Corndog 10 and its homologues in Firecracker and YungJamal are absent from Catdawg. The deletion in Catdawg reflects a loss of 281 bp relative to the other genomes, and an accompanying insertion of 15 bp of unknown origin. There are no obvious repeated sequences flanking the deletion and the mechanism involved is unclear.

### **Discussion**

The Cluster O mycobacteriophages are an interesting group of phages with several features not found in other phages of *M. smegmatis*. The most obvious of these is their prolate heads with a 4:1 length:width ratio. Prolate-headed phages within the *Caudovirales* are somewhat uncommon, with the best-studied being T4, although the length to width ratio of T4 is relatively small. However, phages with longer heads have been described for other hosts including phages of *Caulobacter* (length:width ratios of 3.5:1–4.5:1) [31] and *Lactobacillus* [32–34] and a model



has been described for the structural organizations of icosahedral prolate capsids [35]. It is notable that HHpred predicts a subunit fold that is very similar to that of HK97, which forms an isometric shell [36]. The genomic and proteomic analyses identified no unusual components of the particles, such as proteins that might specifically determine capsid length, as tape measure proteins do with tails. The prolate shape thus might be determined solely by the physical nature of the capsomers [35].

Mass spectrometry reveals an unexpected dearth of Corndog capsid peptides, as capsid monomers are expected to be the most abundant components of purified virions. Thirteen virion proteins had more peptides than the capsid subunit, including most of the minor tail proteins, the portal, and the proposed capsid protease. Although it is plausible that some peptides were not identified because of covalent crosslinking as in HK97, it is possible that the mature capsid subunits are modified such as to obscure the predicted peptide masses. Four genes between the portal and protease genes have plausible modification functions including an Omethyltransferase (Corndog gp35), glycosyltransferase proteins (gp36, gp37), and a putative N-acetylglucosaminyltransferase (gp38). All four were identified by LC/MS-MS in infected cells and could add complex methyl and glycan modifications to the capsid with unpredictable molecular masses.

The Cluster O phages carry an unusual array of 17 bp repeats of unknown function. They are located throughout the genomes but are more densely positioned towards the right genome ends. Many are intergenic, although about one-third of them are within coding regions. They differ from the Start Associated Sequences (SAS) repeats in the Cluster K phages [37] in not being closely linked to translational initiation sites, and are more similar in their distribution to the stoperator sites in the Cluster A phages [16, 38]. However the Cluster A stoperator sites are asymmetric and orientated with the direction of transcription, an important feature of their proposed function in termination of transcription and silencing [16]. Moreover, we have not been able to recover stable lysogens of Corndog or other Cluster O phages, and they do not encode an integrase or a parAB partitioning system (the parB-like domain proteins such as Corndog gp90 are unlikely to be involved with genome stability) and are not obviously temperate, at least in M. smegmatis mc<sup>2</sup>155. However, the sites clearly have dyad symmetry and are predicted to be bound by dimeric DNA binding proteins. Because the large majority of sites are not associated with predicted promoters, the DNA binding interaction must be involved in a process other than the regulation of transcription initiation. We also note that few, if any, of these short repeats are involved in any of the insertions, deletions or rearrangement observed between the five Cluster O genomes.

Finally, comparative genomics and LC-MS/MS resolve the oddity of an apparent extended non-coding gap in Corndog between the tapemeasure protein gene and the upstream gene, which was similarly predicted in the Firecracker and YungJamal genomes. All three also share the insertion of an HNH gene upstream of this apparent non-coding gap. LC-MS/MS analysis shows that translation does indeed begin upstream, although where translation initiates remains unclear, and we have been unable to determine the N-terminal sequence of the tape measure protein by Edman degradation (data not shown). Because there is no commonly used start codon (ATG, GTG, TTG) upstream of the most N-terminal peptides identified, *tmp* expression must use a non-canonical mechanism. Among the possibilities is the use of an unusual codon for translation initiation—perhaps the ACG codon immediately upstream of the N-terminal peptide—or by initiation of translation somewhere upstream coupled with a translational bypass event. Regardless of which non-canonical mechanism is used, there is no obvious reduction in the expression level of *tmp* in Corndog, and the three phages with this arrangement (Corndog, Firecracker, and YungJamal) grow similarly to Catdawg and Dylan that use an ATG start codon.



In summary, the Cluster O mycobacteriophages represent an interesting group of closely related phages with a variety of interesting genomic features. The identification of a variety of conserved features suggests novel and interesting regulatory features warranting experimental investigation.

### **Materials and Methods**

# **Electron Microscopy**

Cluster O phage samples were spotted on 400 mesh carbon coated copper grids, stained with 1% uranyl acetate, and imaged with a Morgagni TEM.

# Bioinformatic analyses

Bioinformatic analyses used DNA Master (<a href="http://cobamide2.bio.pitt.edu/">http://cobamide2.bio.pitt.edu/</a>), Aragorn [39], Gepard [13], HHpred [22], tRNAscan [40], and Phamerator [41]. The Phamerator database used for genomic comparisons was Mycobacteriophage\_292. Phams were built using BLASTP and/or ClustalW, with similarity cut-offs e-values of 10<sup>-50</sup> and 32.5% similarity or better as described elsewhere [41]. Transmembrane domains were identified using SOSUI [42], TopPred [43] and TMHMM [44]. Predicted SigA-like promoters were identified using promoter prediction in DNAMaster set to search for sigma-70 binding sites. The search parameters were as follows: site and merge methods set to geometric, -35 and-10 weights set to 1.0, and spacing weight set to 0.1. The top scoring promoters were evaluated for transcriptional direction of flanking genes and whether they were within or between predicted coding regions.

### SDS-PAGE

Corndog particles were concentrated and purified by CsCl density gradient ultracentrifugation. The visible phage band was dialyzed against two changes of phage buffer (10 mM Tris pH 7.5, 10 mM MgSO<sub>4</sub>, 20 mM NaCl, 1 mM CaCl<sub>2</sub>); 500  $\mu$ l of the dialyzed CsCl band was pelleted by centrifugation for 30min at 14000 rpm. The pellet was resuspended in 75  $\mu$ l of 20 mM DTT, then 2  $\mu$ l of 0.5 M EDTA and 1  $\mu$ l of 1 M MgSO<sub>4</sub> was added. The phage was disrupted by heating to 75°C for 2 mins, and then sonicated on ice six times for 30 seconds to disrupt the DNA. The sample was mixed with 25  $\mu$ l 4 x SDS sample buffer and heated in a boiling bath for 3 minutes at 95°C. The sample was electrophoresed through a 12% polyacrylamide gel containing SDS, and stained with Coomassie Brilliant Blue in methanol.

# Transcript analysis

A log phase *M. smegmatis* mc<sup>2</sup>155 culture was infected with Corndog particles at a multiplicity of infection (moi) of 3, and total RNA collected at various time points post-infection (30 min, 2.5 h, 3.5 h, and 4.5 h) using the Qiagen RNeasy Mini Kit (Qiagen). RNA was treated with DNase I (Invitrogen) and cDNA was generated using random hexamers and Maxima reverse transcriptase (Fermentas). PCR was used with the following primers to check the size of the cDNA product: (5' GAAGGTGCCTTCAAGACGGCCG 3') and (5' GCGACCACATCGCTGATGCTCTG 3'). A Corndog phage lysate was used as a positive control for PCR.

### Mass-spectrometry

LC-MS/MS analysis was performed on Corndog particles purified by either one or two rounds of banding by CsCl equilibrium density centrifugation. For LC-MS/MS analysis of infected cells, 5 mls of exponentially growing M. smegmatis  $mc^2155$  (OD<sub>600</sub> = 0.4) in 7H9 /ADC was concentrated to a 500  $\mu$ l volume via low-speed centrifugation, and infected with Corndog at a



multiplicity of infection (moi) of 100. Phage particles were allowed to adsorb for 15 minutes, then 4.5 mls of fresh 7H9 medium was added, and incubated further with shaking for three hours at 37°C; the OD<sub>600</sub> was monitored throughout to follow cell growth and lysis. At 165 minutes post-adsorption, a 1-millilter aliquot was removed from the culture, the cells were pelleted via centrifugation (1 min, 14K rpm in a microfuge), and the supernatant was removed. The cell pellet was frozen at—80°C, and then shipped overnight on wet-ice to the University of California, Davis Proteomics Core (UCDPC) http://proteomics.ucdavis.edu. There, the cells were lysed via a MagnaLyser, the insoluble fraction was removed, and the soluble proteins were precipitated, digested with Trypsin, and cleaned-up using a macro spin-column. The peptides were then separated using an Easy-LC II High-Pressure Liquid Chromatography HPLC system and loaded into a Q-exactive orbitrap mass spectrometer with a Proxeon nano-spray source (Thermo) for tandem ms analysis. Detected spectra and fragmentation profiles were matched against a database comprised of a six-frame translation of the Corndog genome, the annotated proteins of M. smegmatis mc<sup>2</sup>155, and UniProt using X!Tandem. Peptide matches were analyzed using Scaffold4. Settings used a peptide threshold of 95%, and protein FDR of 1%. For proteomic analysis of Catdawg, a 1 ml aliquot of a phage lysate was pelleted at 14K for 2 hours at 4°C, resuspended in 100 µl of 0.1 M phosphate buffer and shipped overnight on dry ice to MSBioworks (http://www.msbioworks.com/) for mass spectrometry analysis of phage proteins.

For N-terminal analysis of proteins, a Catdawg lysate were labeled with 200 mM TMPP in 20% acetonitrile [45]. Approximately 20  $\mu g$  of labeled proteins were resolved and separated on a 4–12% Bis Tris SDS-PAGE gel in MOPS buffer and the gel lanes excised into 20 equally sized segments. Gel segments were protease digested using either trypsin or chymotrypsin and analyzed by nano LC-MS/MS with a Waters NanoAcquity HPLC system interfaced to a Thermo-Fisher Orbitrap Velos Pro. Peptides were loaded on a trapping column and eluted over a 75  $\mu m$  analytical column at 350 nL/min. The mass spectrometer was operated in data-dependent mode, with MS performed in the Orbitrap at 60,000 FWHM resolution and MS/MS performed in the LTQ. The 15 most abundant ions were analyzed. Mascot DAT files were parsed into Scaffold for validation and filtered to create a non-redundant list. Filtering used a minimum protein value of 99% and peptide value of 50% (Prophet scores), and required at least two unique peptides per protein. Protease peptide data were merged for analysis. Peptide data from the two different proteases were merged using Scaffold4 for subsequent data analysis Settings used a peptide threshold of 95%, and protein FDR of 1%.

### Supporting Information

**S1 Fig. Comparison of Cluster O genome maps.** Genome maps of the five Cluster O phages, Corndog, Catdawg, Dylan, Firecracker and YungJamal were generated by Phamerator using the database mycobacteriophage\_292 (41). Genes are shown as boxes above (rightwards-transcribed) or below (leftwards-transcribed) the genome with gene names within the boxes. Phamily assignments for genes are shown above the boxes with the number of phamily members in parentheses. Shading between genomes shows pairwise nucleotide sequence similarity and spectrum colored with violet being the most similar, and red being the least similar but above the threshold BLASTN E value of 10<sup>-5</sup>. (PDF)



# **Acknowledgments**

We thank the Howard Hughes Medical Institute for administrative contributions to the Science Education Alliance (SEA) Phage Hunters Advancing Genomics and Evolutionary Science (SEA-PHAGES) and the Phage Hunters Integrating Research and education (PHIRE) programs.

### **Author Contributions**

Conceived and designed the experiments: SGC WHP GAH RAK GFH. Performed the experiments: DD MP JSL DAR RMD EP TC SC VM DJS. Analyzed the data: TA SHB SET DD AJL D. Bollivar RLD JRH GPK KRA MP JS D. Byrnes JFS VCW RP NR KAC UPG CKS KB D. Westholm SB CB GAH AMF CRG SDM LEH KK RAK CL CAB DJS WHP SGC RWH DAR RMD JSL EP DM D. Wodarski CDLS KE CSJ SLK SMM ALW TSZ MZ GFH. Contributed reagents/materials/analysis tools: SGC. Wrote the paper: SGC WHP DJS RWH GFH. Author contributions are listed in Table S1.

### References

- Hatfull GF, Hendrix RW. Bacteriophages and their Genomes. Current Opinions in Virology. 2011; 1, 298–303. doi: 10.1016/j.coviro.2011.06.009 PMID: 22034588
- Wommack KE, Colwell RR. Virioplankton: viruses in aquatic ecosystems. Microbiol Mol Biol Rev. 2000; 64(1):69–114. PMID: 10704475
- Abrescia NG, Bamford DH, Grimes JM, Stuart DI. Structure unifies the viral universe. Annu Rev Biochem. 2012; 81:795–822. Epub 2012/04/10. doi: <a href="https://doi.org/10.1146/annurev-biochem-060910-095130">10.1146/annurev-biochem-060910-095130</a> PMID: 22482909
- 4. Hendrix RW, Smith MC, Burns RN, Ford ME, Hatfull GF. Evolutionary relationships among diverse bacteriophages and prophages: all the world's a phage. Proc Natl Acad Sci U S A. 1999; 96(5):2192–7. Epub 1999/03/03. PMID: 10051617
- Hendrix RW. Bacteriophages. In: Knipe DM, Howley PM, editors. Fields Virology, Sixth Edition. Philadelphia: Lippincott Williams & Wilkins; 2013.
- Jacobs-Sera D, Marinelli LJ, Bowman C, Broussard GW, Guerrero Bustamante C, Boyle MM, et al. On the nature of mycobacteriophage diversity and host preference. Virology. 2012; 434(2):187–201. doi: 10.1016/j.virol.2012.09.026 PMID: 23084079
- Hatfull GF. Molecular Genetics of Mycobacteriophages. Microbiology Spectrum. 2014; 2(2):1–36. doi: 10.1128/microbiolspec.MGM2-0032-2013 PMID: 25328854
- Hatfull GF. The secret lives of mycobacteriophages. Adv Virus Res. 2012; 82:179–288. doi: 10.1016/ B978-0-12-394621-8.00015-7 PMID: 22420855
- Pedulla ML, Ford ME, Houtz JM, Karthikeyan T, Wadsworth C, Lewis JA, et al. Origins of highly mosaic mycobacteriophage genomes. Cell. 2003; 113(2):171–82. PMID: 12705866
- Pitcher RS, Tonkin LM, Daley JM, Palmbos PL, Green AJ, Velting TL, et al. Mycobacteriophage exploit NHEJ to facilitate genome circularization. Mol Cell. 2006; 23(5):743–8. PMID: 16949369
- Hatfull GF, Science Education Alliance Phage Hunters Advancing G, Evolutionary Science P, Kwa-Zulu-Natal Research Institute for T, Students HIVMGC, Phage Hunters Integrating R, et al. Complete genome sequences of 138 mycobacteriophages. J Virol. 2012; 86(4):2382–4. doi: 10.1128/JVI.06870-11 PMID: 22282335
- Jordan TC, Burnett SH, Carson S, Caruso SM, Clase K, DeJong RJ, et al. A broadly implementable research course in phage discovery and genomics for first-year undergraduate students. MBio. 2014; 5 (1):e01051–13. doi: 10.1128/mBio.01051-13 PMID: 24496795
- Krumsiek J, Arnold R, Rattei T. Gepard: a rapid and sensitive tool for creating dotplots on genome scale. Bioinformatics. 2007; 23(8):1026–8. PMID: 17309896
- Oldfield LM, Hatfull GF. Mutational Analysis of the Mycobacteriophage BPs Promoter PR Reveals Context-Dependent Sequences for Mycobacterial Gene Expression. J Bacteriol. 2014; 196(20):3589–97. doi: 10.1128/JB.01801-14 PMID: 25092027
- Nesbit CE, Levin ME, Donnelly-Wu MK, Hatfull GF. Transcriptional regulation of repressor synthesis in mycobacteriophage L5. Mol Microbiol. 1995; 17(6):1045–56. Epub 1995/09/01. PMID: 8594325



- Brown KL, Sarkis GJ, Wadsworth C, Hatfull GF. Transcriptional silencing by the mycobacteriophage L5 repressor. Embo J. 1997; 16(19):5914–21. Epub 1997/10/06. doi: 10.1093/emboj/16.19.5914
   PMID: 9312049
- Dedrick RM, Marinelli LJ, Newton GL, Pogliano K, Pogliano J, Hatfull GF. Functional requirements for bacteriophage growth: gene essentiality and expression in mycobacteriophage Giles. Mol Microbiol. 2013; 88(3):577–89. doi: 10.1111/mmi.12210 PMID: 23560716
- Czyz A, Mooney RA, Iaconi A, Landick R. Mycobacterial RNA polymerase requires a U-tract at intrinsic terminators and is aided by NusG at suboptimal terminators. MBio. 2014; 5(2):e00931. doi: 10.1128/ mBio.00931-14 PMID: 24713321
- 19. Bailey TL, Boden M, Buske FA, Frith M, Grant CE, Clementi L, et al. MEME SUITE: tools for motif discovery and searching. Nucleic Acids Res. 2009; 37(Web Server issue):W202–8. doi: 10.1093/nar/gkp335 PMID: 19458158
- 20. Kim BJ, Kim BR, Lee SY, Seok SH, Kook YH, Kim BJ. Whole-Genome Sequence of a Novel Species, Mycobacterium yongonense DSM 45126T. Genome announcements. 2013; 1(4). doi: 10.1128/genomeA.00604-13.
- Pope WH, Jacobs-Sera D, Russell DA, Rubin DH, Kajee A, Msibi ZN, et al. Genomics and proteomics of mycobacteriophage patience, an accidental tourist in the mycobacterium neighborhood. MBio. 2014;5(6). doi: 10.1128/mBio.02145-14.
- Soding J, Biegert A, Lupas AN. The HHpred interactive server for protein homology detection and structure prediction. Nucleic Acids Res. 2005; 33(Web Server issue):W244–8. Epub 2005/06/28. doi: 10.1093/nar/gki408 PMID: 15980461
- Oh B, Moyer CL, Hendrix RW, Duda RL. The delta domain of the HK97 major capsid protein is essential for assembly. Virology. 2014; 456–457:171–8. doi: 10.1016/j.virol.2014.03.022.
- Duda RL, Oh B, Hendrix RW. Functional domains of the HK97 capsid maturation protease and the mechanisms of protein encapsidation. J Mol Biol. 2013; 425(15):2765–81. doi: 10.1016/j.jmb.2013.05. 002 PMID: 23688818
- Popa MP, McKelvey TA, Hempel J, Hendrix RW. Bacteriophage HK97 structure: wholesale covalent cross-linking between the major head shell subunits. J Virol. 1991; 65(6):3227–37. PMID: 1709700
- Huang WM, Ao SZ, Casjens S, Orlandi R, Zeikus R, Weiss R, et al. A persistent untranslated sequence within bacteriophage T4 DNA topoisomerase gene 60. Science. 1988; 239(4843):1005–12. PMID: 2830666
- Sampson T, Broussard GW, Marinelli LJ, Jacobs-Sera D, Ray M, Ko CC, et al. Mycobacteriophages BPs, Angel and Halo: comparative genomics reveals a novel class of ultra-small mobile genetic elements. Microbiology. 2009; 155(Pt 9):2962–77.
- 28. Kala S, Cumby N, Sadowski PD, Hyder BZ, Kanelis V, Davidson AR, et al. HNH proteins are a wide-spread component of phage DNA packaging machines. Proc Natl Acad Sci U S A. 2014; 111 (16):6022–7. doi: 10.1073/pnas.1320952111 PMID: 24711378
- Stolt P, Grampp B, Zillig W. Genes for DNA cytosine methyltransferases and structural proteins, expressed during lytic growth by the phage phi H of the archaebacterium Halobacterium salinarium. Biological chemistry Hoppe-Seyler. 1994; 375(11):747–57. PMID: 7695837
- Singh S, Plaks JG, Homa NJ, Amrich CG, Heroux A, Hatfull GF, et al. The Structure of Xis Reveals the Basis for Filament Formation and Insight into DNA Bending within a Mycobacteriophage Intasome. J Mol Biol. 2014; 426(2):412–22. doi: 10.1016/j.jmb.2013.10.002 PMID: 24112940
- Gill JJ, Berry JD, Russell WK, Lessor L, Escobar-Garcia DA, Hernandez D, et al. The Caulobacter crescentus phage phiCbK: genomics of a canonical phage. BMC Genomics. 2012; 13:542. doi: 10.1186/1471-2164-13-542 PMID: 23050599
- Forsman P. Characterization of a prolate-headed bacteriophage of Lactobacillus delbrueckii subsp. lactis, and its DNA homology with isometric-headed phages. Arch Virol. 1993; 132(3-4):321-30. PMID: 8397503
- Riipinen KA, Forsman P, Alatossava T. The genomes and comparative genomics of Lactobacillus delbrueckii phages. Arch Virol. 2011; 156(7):1217–33. doi: 10.1007/s00705-011-0980-5 PMID: 21465086
- Ackermann HW. 5500 Phages examined in the electron microscope. Arch Virol. 2007; 152(2):227–43.
  PMID: 17051420
- Luque A, Reguera D. The structure of elongated viral capsids. Biophys J. 2010; 98(12):2993–3003. doi: 10.1016/j.bpj.2010.02.051 PMID: 20550912
- Hendrix RW, Johnson JE. Bacteriophage HK97 capsid assembly and maturation. Adv Exp Med Biol. 2012; 726:351–63. Epub 2012/02/03. doi: 10.1007/978-1-4614-0980-9\_15 PMID: 22297521



- Pope WH, Ferreira CM, Jacobs-Sera D, Benjamin RC, Davis AJ, DeJong RJ, et al. Cluster K Mycobacteriophages: Insights into the Evolutionary Origins of Mycobacteriophage TM4. PLoS ONE. 2011; 6(10):e26750. doi: 10.1371/journal.pone.0026750 PMID: 22053209
- Pope WH, Jacobs-Sera D, Russell DA, Peebles CL, Al-Atrache Z, Alcoser TA, et al. Expanding the Diversity of Mycobacteriophages: Insights into Genome Architecture and Evolution. PLoS ONE. 2011; 6(1):e16329. doi: 10.1371/journal.pone.0016329 PMID: 21298013
- Laslett D, Canback B. ARAGORN, a program to detect tRNA genes and tmRNA genes in nucleotide sequences. Nucleic Acids Res. 2004; 32(1):11–6. PMID: 14704338
- Lowe TM, Eddy SR. tRNAscan-SE: a program for improved detection of transfer RNA genes in genomic sequence. Nucleic Acids Res. 1997; 25(5):955–64. PMID: 9023104
- Cresawn SG, Bogel M, Day N, Jacobs-Sera D, Hendrix RW, Hatfull GF. Phamerator: a bioinformatic tool for comparative bacteriophage genomics. BMC Bioinformatics. 2011; 12:395. doi: 10.1186/1471-2105-12-395 PMID: 21991981
- Hirokawa T, Boon-Chieng S, Mitaku S. SOSUI: classification and secondary structure prediction system for membrane proteins. Bioinformatics. 1998; 14(4):378–9. PMID: 9632836
- **43.** Claros MG, von Heijne G. TopPred II: an improved software for membrane protein structure predictions. Computer applications in the biosciences: CABIOS. 1994; 10(6):685–6. PMID: 7704669
- Sonnhammer EL, von Heijne G, Krogh A. A hidden Markov model for predicting transmembrane helices in protein sequences. Proc Int Conf Intell Syst Mol Biol. 1998; 6:175–82. PMID: 9783223
- 45. Baudet M, Ortet P, Gaillard JC, Fernandez B, Guerin P, Enjalbal C, et al. Proteomics-based refinement of Deinococcus deserti genome annotation reveals an unwonted use of non-canonical translation initiation codons. Molecular & cellular proteomics: MCP. 2010; 9(2):415–26. doi: 10.1074/mcp.M900359-MCP200.