Prevalence of Bacillus anthracis-Like Organisms and Bacteriophages in the Intestinal Tract of the Earthworm Eisenia fetida

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Stable infection of Bacillus anthracis laboratory strains with environmental bacteriophages confers survival phenotypes in soil and earthworm intestinal niches (R. Schuch and V. A. Fischetti, PLoS One 4:e6352, 2009). Here, the natural occurrence of two such B. anthracis-infective bacteriophages, Wip1 and Wip4, was examined in the intestines of Eisenia fetida earthworms as part of a 6-year longitudinal study at a Pennsylvania forest site. The Wip1 tectivirus was initially dominant before being supplanted by the Wip4 siphovirus, which was then dominant for the next 3 years. In a host range analysis of a wide-ranging group of Bacillus species and related organisms, Wip1 and Wip4 were both infective only toward B. anthracis and certain B. cereus strains. The natural host of Wip4 remained constant for 3 years and was a B. cereus strain that expressed a B. anthracis-like surface polysaccharide at septal positions on the cell surface. Next, a novel metagenomic approach was used to determine the extent to which such B. cereus- and B. anthracis-like strains are found in worms from two geographical locations. Three different enrichment strategies were used for metagenomic DNA isolation, based either on the ability of B. cereus sensu lato to form heat-resistant spores, the sensitivity of B. anthracis to the PlyG lysis, or the selective amplification of environmental phages cocultured with B. anthracis. Findings from this work indicate that B. cereus sensu lato and its phages are common inhabitants of earthworm intestines.

The notorious animal pathogen Bacillus anthracis is considered to be a genetically monomorphic variant of the large, well-studied B. cereus sensu lato (s.l.) lineage of often ubiquitous soil organisms like B. cereus sensu stricto (referred to here as simply B. cereus), B. thuringiensis, B. pseudomycoides, B. weihenstephanensis, and B. mycoides (20, 21, 25, 38). Despite a high degree of chromosomal DNA sequence similarity and synteny within B. cereus s.l., it nonetheless has a highly pleomorphic group of saprophytes, intestinal symbionts, and both opportunistic and frank pathogens of many animals (19, 39). The niche-specific adaptive behaviors associated with B. cereus s.l. are often attributed to highly variable sets of potentially transmissible megaplasmids (20, 25). For example, the pXO1 and pXO2 plasmids of B. anthracis encode both the virulence genes directly responsible for anthrax and the transcriptional regulatory proteins involved in plasmid and chromosomal gene expression (23, 40). The presence of large plasmids alone, however, does not sufficiently explain the ability of B. cereus s.l. organisms to elaborate different behaviors. Rather, species-specific chromosomal loci, accounting for ca. 15% of the genes in each sequenced B. cereus s.l. genome (18), may also have significant phenotypic impacts. Included among the highly variable loci of B. cereus s.l. are a diverse range of mobile genetic elements, including small plasmids, insertion sequences, transposons, introns and, importantly, bacteriophages (21).

Bacteriophages have a very well-described impact on bacterial genetic diversity and behavior through the introduction of virus-encoded factors that alter bacterial phenotypes in a process called lysogenic conversion (7). Despite the presence of species-specific bacteriophages throughout the B. cereus s.l. lineage, either in chromosomes as prophages or as independently replicating episomes, the possibility of lysogenic conversion has, until recently (30), received little attention. Toward this end, we recently described a set of bacteriophages with the ability to enhance environmental survival phenotypes of B. cereus s.l. (30). Lysogeny with any of a series of soil-, fern rhizosphere-, and earthworm gut-derived phages was found to reduce or block sporulation, favor biofilm formation, and promote long-term survival in both soil microcosms and the intestinal tract of Eisenia fetida worms (30). This effect was observed with both laboratory B. anthracis strains and environmental B. cereus isolates. Furthermore, the earthworm colonization phenotype was driven solely by expression of phage-encoded bacterial sigma factors which can activate the expression of at least one (but likely more) B. anthracis chromosomal loci (30). Here, bacteriophages promote environmental functioning in a process by which phage-encoded transcription factors activate host-encoded phenotypes.

The initial goal of the present study was to describe two B. anthracis-infective phages originally found in the intestines of E. fetida worms from a forest leaf litter site in Pennsylvania. We followed the occurrence of these phages over 6 years and characterized one natural bacterial host. Based on the identification of B. cereus s.l. organisms and phages in the worm niche, metagenomic methods were developed and used to specifically examine the extent to which such bacteria and phages are found in worms at a separate location, in New York City.

MATERIALS AND METHODS

Bacterial strains and growth conditions. The majority of bacterial strains in the present study were previously described (29, 30, 32). Additional strains include Staphylococcus aureus Newman (3), B. cereus E33L (12), and B. thuringiensis Al Hakam (8). All organisms, except Escherichia coli, were grown on brain-
heart infusion (BHI; Difco) broth or agar plates at 30°C according to standard protocols. E. coli strain TOP10 (Invitrogen, Inc.) was grown on Luria-Bertani (LB; Sigma-Aldrich, Inc.) broth or agar plates at 37°C according to standard protocols.

**Virus manipulations.** Wy was obtained from the Felix d’Herelle Reference Center for Bacterial Viruses at Laval University in Quebec, Canada. Phage propagation, storage, and host range analyses were performed exactly as described previously (29). Phage plaque assays were always performed with a “no phage” control, to account for a potential background of induced prophage.

**Worm and soil collection and analysis.** E. fetida worms and surrounding soils were taken from two sites: a forest floor in Stroudsburg, Pennsylvania (PA), and an outdoor garden in New York City, New York (NYC). In PA, on the first Saturday of October over a 6-year period from 2003 until 2008, ~5 g of soil and two worms were removed from each of six positions at roughly three meter increments extending both north and south. In NYC, a total of 12 worms were collected once in June 2009. In all cases, worms and/or soil were taken from within ~12 cm of the surface. Soil from both the NYC garden and PA forest was always humus-like (i.e., dark brown, porous, and spongy), with a pH between 6 and 8. All samples were immediately placed in separate 50-ml Falcon tubes and transported to the laboratory for processing within 24 h. Recovery and identification of bacteriophages and bacterial organisms from worms and soil was performed as described previously (30) in the exact same manner every year. For PCR analyses of worm and soil samples, 25 µl of each environmental slurry was mixed with 25 µl of 0.5 M NaOH for 15 min at room temperature and diluted with 25 µl of 1 M Tris (pH 8.0) and 1.0 ml of distilled H2O to yield a PCR template. Then, 1.0 µl of each template was used in a 100-µl PCR containing the Taq PCR master mix (Qiagen, Inc.). Phase-specific DNA primers identical to those previously used (30). DNA samples were stored at ~20°C between analyses.

**Search for natural phage lysogens.** A bacterial culture-based method was used to search for the natural hosts of both Wip1 (from all worms sampled from 2003 to 2005) and Wip4 (from all worms sampled from 2006 to 2008). Extracted worm gut samples from each year were plated undiluted (and three 100-fold serial dilutions thereof) on 150-mm BHI agar plates. After overnight incubation, individual colonies exhibiting characteristic R. pseudocancillerus or B. cereus-like morphologies (37) were recovered, subcultured on BHI agar plates, and examined by PCR in the manner described with Wip1- and Wip4-specific PCR primers (lysogens are positive for these markers). PCR-positive organisms were subcultured twice and reexamined by PCR to confirm the presence of phage.

**Sequence analyses of RS1045.** The Wip4-positive strain RS1045 was used as DNA template source for 16S rRNA sequencing with primers 16S-up (5′-AGA GTTGTGATCCTGGCTCAG-3′) and 16S-down (5′-AGA AGAAAGGAGTGTGATCACTTC-3′) in a manner previously described (27). The plcR-papR locus of RS1045 was first amplified with the flanking primers PlcRup (5′-GCTCAATGATGTAGTGAATTCGGCTCGAG-3′) and PlcRdown (5′-GAAAGAGGTCGTTAGATGTCCT-3′), and the product was sequenced with PlcRup, PlcRdown, and the forward and reverse versions of PlcRmid1 (5′-GAGGGTGGCGGCTCTCCTAGGAGATTCTCG-3′) and PlcRmid2 (5′-GACGCTCTTTGACGGATCATG-3′). The resulting sequence was performed by GeneWiz, Inc. The resulting data were assembled by using SeqMan Pro (DNASTAR, Inc.) and analyzed by using BlastN.

**Genomic fingerprinting.** M13 fingerprint analysis was performed as described previously (2). Briefly, a M13 primer was used (5′-GAGGTGTTCCGCGCCTCT-3′) to generate size pattern variations that are diagnostic for bacterial species and phages. PCR products were denatured for 10 min at 95°C, followed by 40 cycles of 15 s at 96°C, 15 s at 36°C, and 2 min at 72°C. Resulting PCR products were separated on 1% agarose gels and visualized with ethidium bromide.

**Sequence alignments.** Multiple sequence alignments were obtained by using the CLUSTAL W server (http://www.ch.embnet.org/software/CLUSTALW.html) and visualized by using BOXSHADE 3.21 (http://www.ch.embnet.org/software/BOX_form.html).

**Phase-contrast and fluorescence microscopy.** The construction and purification of GFP-PlyG<sup>md</sup> was described previously (29, 30). For labeling, strains were transformed with a plasmid containing the GFP-PlyG<sup>md</sup> expression vector (using the pBAD TOPO TA expression kit of Invitrogen, Inc.) and transformed into Escherichia coli TOP10 cells. Colonies appearing on LB agar supplemented with 100 µg of ampicillin ml<sup>−1</sup> were randomly chosen for plasmid purification and DNA sequence analysis.

**Phylogenetic analysis.** For the production of a “spore-enhanced” metagenomic DNA library, a variation was introduced into the protocol prior to library construction. Isolation of worm gut material was first washed twice with PBS, resuspended in 0.5 ml of PBS, and heated at 95°C for 30 min to denature all nonsporozo forms. After heating, samples were placed on ice for 5 min, diluted into 2 ml of BHI and incubated with aeration at 37°C for 3 h. The 3-h incubation is sufficient to promote both spore germination and outgrowth (4), as well as the subsequent vegetative expansion of surviving forms (with clear samples becoming turbid). After this treatment, samples were washed twice with PBS (to remove remaining exogenous DNA) and resuspended in 0.5 ml of PBS prior to processing as described above for the analysis of metagenomic DNA.

For the “PhyG-released” metagenomic DNA library, worm gut material was first washed five times with PBS, resuspended in 0.5 ml of PBS, and mixed with 0.5 ml of a purified PhyG preparation (6.4-mg ml<sup>−1</sup> stock concentration) for 2 h at 37°C. Samples were then pelleted by centrifugation (4,000 rpm for 5 min in an Eppendorf Model 5415D Microcentrifuge), and the resulting supernatants were passed through a 0.22-µm-pore-size filter to remove contaminating material. The DNA-bearing lysate was extracted with phenol-chloroform and precipitated with ethanol prior to metagenomic library construction as described above.

For the production of *B. anthracis* phage-enhanced metagenomic DNA, another variation was used whereby phage-bearing supernatants of pellets introduced into the culture broth were incubated at 30°C with agitation. After an overnight incubation, potentially phage-bearing culture supernatants were collected, filtered to remove contaminating material, and used to infect another culture of *B. anthracis* DStere growing at 30°C with agitation. Duplicate sequences were recovered, filtered, extracted, with phenol-chloroform, and precipitated with ethanol prior to metagenomic library construction as described above.

Fifty clones were sequenced for each library. The number of resulting high-quality sequences varied with each library and included 47 for the total bacteria, 25 for the spore enhanced, 48 for the PhyG released, and 24 for the *B. anthracis* phage enhanced. Duplicate sequences were not observed. All missing sequences were clones with no library inserts.

**Nucleotide sequence accession numbers.** Sequences of both the entire plcR-papR locus and the partial 16S rRNA region of RS1045 were submitted to GenBank (under accession numbers GU142941 and GU142940, respectively). Sequences of the Wip1 and Wip4 lysins (GU142942 and GU142943, respectively), as well as those identified in the metagenomic analyses (GU142944, GU142945, GU142946, and GU142948), were submitted as well.

**RESULTS**

**Longitudinal survey: sample recovery.** Every first weekend of October over a 6-year period, two worms were recovered from each of six positions at 3-m increments extending both...
9 m north and east of a central, decaying tree stump. The 12 animals were returned to the laboratory, washed separately in distilled H₂O to remove adherent dirt particles, and processed for the recovery of gut contents. The contents were centrifuged, and the resulting supernatants were filtered to yield bacterium-free extracts containing worm-gut phages.

**Longitudinal survey: bacteriophage detection.** Bacteriophages in both worm and soil extracts were sought as PFU in two ways. First, extract aliquots were incubated for 16 h on BHI soft-agar overlays containing freshly plated lawns of organisms, including *B. anthracis, B. thuringiensis, B. cereus, Staphylococcus aureus, Streptococcus pyogenes,* and *Pseudomonas aeruginosa.* From each worm isolated throughout this experiment (except for in 2006 when no PFU were found), 1 × 10⁷ to 5 × 10⁸ PFU were recovered using the *B. anthracis* ΔSterne reporter. All other reporter organisms (including *B. thuringiensis* HD1, *B. cereus* ATCC 14579 and ATCC 10987, *S. aureus* Newman, and *P. aeruginosa* PAO1) yielded no PFU. In addition, all soil extracts were free of phages infecting the reporter organisms, including *B. anthracis* (see Table S1 in the supplemental material).

For the second method to detect PFU, each reporter organism was subjected to two 24-h infections with worm gut extracts. After this amplification step, infected culture supernatants were collected, filtered to remove bacteria, and titered on agar plates freshly overlaid with the corresponding reporter strain. This “phage-enrichment” method was previously used to detect otherwise undetectable phages in environmental samples (30). Here, 10⁶ to 10⁸ PFU per worm were found every year (including 2006) using the *B. anthracis* ΔSterne reporter. No plaques were observed on other reporter strains, including *B. cereus* and *B. thuringiensis.*

Surprisingly, no PFU were detected in the extracts of PA forest soil from which phage-bearing worms were isolated. Free intestinal phages should be released in worm excreta referred to as castings. Although released phages could have adsorbed to the humuslike soil or casting material, the preparation methods used here should have released some detectable phages into solution. Previously, we successfully isolated *B. anthracis*-infective phages (Slp1 and Frp1) from the soil using the same technique (30). To confirm here that the amplification step does indeed work, soil from three additional environments was examined (see Table S1 in the supplemental material). Although the primary extracts from these soils yielded no PFU on indicator strains (with two exceptions), the phage enrichment method did yield PFU for most *Bacillus* strains tested. Thus, while multiple soil samples do contain *Bacillus* phages, none are detectable in the soil containing PA worms in the present study.

**Longitudinal survey: bacteriophage identification.** To study *B. anthracis*-infective, worm intestinal phages, a single plaque was first obtained from the primary extract of a worm isolated in 2003 in PA. This phage, called Wip1 (for worm intestinal phage 1), was expanded and used for the identification of a unique phage DNA sequence based on a genetic screen for lysis activity (31). Thus, Wip1 genomic DNA was used to generate an inducible expression library for a lytic activity screen against *B. anthracis.* DNA sequences of 15 lytic clones revealed a common 657-bp open reading frame encoding a putative 218-residue protein similar to the glucosaminidases of *Tectiviridae* phages of Gram-positive bacteria (Fig. 1a). Two distinct Wip1 lysozyme gene-specific primer pairs were then used to screen both the primary gut extracts and the 125 resulting plaques derived from every worm between 2003 and 2008. Ultimately, Wip1-specific PCR products were observed with every gut extract and individual plaque isolated between 2003 and 2005. After 2005, Wip1 was never detected again. The genetic similarity of the Wip1 phages from 2003 to 2005 (25 phages per year) was confirmed by M13 genetic fingerprinting. DNA fingerprints of phage isolates from both 2004 and 2005 were identical to each other and to Wip1 from 2003 (Fig. 2a). Wip1 was, therefore, a common *B. anthracis* phage in the worm gut environment for 3 years (2003 to 2005).

The presence of Wip1-negative PFU after 2005 suggested the presence of a new dominant phage(s). To identify new phages, a high-titer lysate was first generated from a single plaque obtained from one worm in 2006. Genomic DNA was ultimately purified from the resulting phage (referred to as Wip4) and used to generate an expression library for a lysozyme activity screen as described above. Sequences of 10 lytic clones revealed a common 1,056-bp locus encoding a 351-residue protein most similar to N-acetylglucosaminidase in prophages of *B. thuringiensis,* *B. cereus,* *B. mycoides,* and *B. pseudomycoides* (Fig. 1b). Two distinct primer pairs specific for the Wip4 lysis gene were used in a PCR screen of all gut extract samples (from 2003 to 2008) and 150 plaques derived from each worm extract recovered after 2005. Although each pre-2006 sample was PCR negative for Wip4 (yet positive for Wip1), all of the 2006 to 2008 samples were Wip4-positive and Wip1-negative. M13 genetic fingerprints of the 2006 to 2008 phages were also identical (and distinct from the year 2003 to 2005 phages) (Fig. 2a), thus confirming a close genetic relationship. As a control, a PCR screen was also performed with primers specific to four previously described *B. anthracis* phages (Wb5, Frp2, Bcp1, and Wip2) known to be genetically distinct from Wip1 and Wip4 (30); these primers yielded no products using DNA templates obtained from worms between 2003 and 2008.

**Host range analysis.** Previously, phages corresponding to Wip1 and Wip4 were found to infect either of two lab strains of *B. anthracis,* but not the *B. cereus* strain ATCC 14579 (30). Since many *B. anthracis* phages (including Wγ and Wj8) can also infect certain *B. cereus* strains (29, 32), we decided to greatly expand our host range study to include more closely and distantly related organisms. For comparison, an analysis of the standard *B. anthracis* diagnostic phage used in clinical laboratories (1), Wγ, was included. High-titer phage preparations were generated on *B. anthracis* ΔSterne and tested for PFU formation on the strains in Table 1. Each phage was largely infective for only *B. anthracis* and the *B. cereus* strains ATCC 4342 and CDC52805. ATCC 4342 has previously been shown to support replication of phage Wγ (32). Wip1 was notably the most specific phage, infecting all three *B. anthracis* strains. Surprisingly, the Wγ diagnostic phage did not infect the environmental *B. anthracis* strain RS1615 and yielded plaques on ΔSterne only after 5 days. Thus, while Wip1 and Wip4 appear to be at least as selective as Wγ, Wip1 may be the superior diagnostic tool.

**Natural hosts of Wip1 and Wip4.** After the detection of Wip1 PFU and Wip1 loci by PCR in the worm gut, a natural
host of Wip1 was sought. Gut contents of multiple distinct worms (from 2003 to 2005) were first plated on BHI agar. Of the different \textit{B. cereus} s.l.-like colonies recovered, 100 were chosen each year and screened by PCR with Wip1-specific primers (data not shown). Despite the presence of “free” Wip1 in worm guts, no Wip1 lysogen was subsequently identified. A host may either have been obscured by background organisms, not expressed a \textit{B. cereus} s.l.-like colony morphology, or existed in a nonculturable state.

In contrast to Wip1, natural Wip4 lysogens were repeatedly identified in the guts of all worms from 2006 to 2008. Each of the worms from 2006 to 2008 was enriched for a \textit{B. cereus}-like strain that was PCR positive for all Wip4-specific primers and present at an estimated concentration of $\sim 6 \times 10^3$ CFU per worm. This is similar to previously reported values for steady-state colonization of the worm gut with vegetative forms of different \textit{B. cereus} s.l. strains (30). The M13 genetic fingerprints of Wip4 lysogens isolated from 2006 to 2008 were all identical (Fig. 2b), suggesting the same bacterial host was present for all 3 years.

In a previous study, a Wip4 lysogen was originally found to have some \textit{B. cereus}-like phenotypes, including colony morphology, beta-hemolysis, and resistance to the antibiotic fosfomycin (30). In the present study, DNA sequence analysis of the Wip4 lysogen strain (now referred to as RS1045) revealed the 5' half of the 16S rRNA locus to be 99% identical to sequences of only \textit{B. anthracis}, \textit{B. cereus} and \textit{B. thuringiensis}. The plcR-papR locus of RS1045, encoding a quorum-sensing system of \textit{B. cereus} s.l. (33), was also sequenced and found to be most similar (between 92 and 93% identical) to \textit{B. cereus} s.l. animal.

**FIG. 1.** (a) Sequence alignment of lysins encoded by the \textit{Tectiviridae} phages of \textit{B. cereus} s.l. The Wip1 lysin is compared to that of \textit{B. anthracis} phage AP50 (GenBank accession number EU408779) and \textit{B. thuringiensis} phages GIL16c (GenBank accession numberAY701338) and Bam35c (GenBank accession numberAY257527). (b) Sequence alignment of the Wip4 lysin with the lysins of prophages from \textit{B. thuringiensis} BGSC 4CC1 (Bt-pro), \textit{B. cereus} G9842 (Bc-pro), \textit{B. mycoides} Rock3-17 (Bmy-pro), and \textit{B. pseudomycoide} DSM 1244 (Bpm-pro) bearing the GenBank accession numbers ZP_04078846, YP_002446312, ZP_04160784, and ZP_04152425, respectively.
pathogenic strains, including \textit{B. anthracis}, \textit{B. thuringiensis} 97-27, \textit{B. cereus} 03BB102, and others (see Fig. S1 in the supplemental material). In addition, the protein sequences of PlcR and PapR were most similar (with 98 and 100\% identity, respectively) to the products of \textit{B. cereus} strain AH1273, a human blood infection isolate from Iceland (13). The last five residues of PapR encoded by RS1045 were VPFEY, indicating a phenotype present in all \textit{B. anthracis} strains (33).

RS1045 was observed to be a rod-shaped bacillus (Fig. 3a) that binds GFP-PlyG\textsuperscript{BD} at an exclusively sepal position (Fig. 3b). GFP-PlyG\textsuperscript{BD} consists of the C-terminal cell wall binding domain of the \gamma phage lysin, PlyG, fused to a fluorescent protein; the resulting molecule specifically directs fluorescence to the surface polysaccharides of \textit{B. anthracis} and some closely related \textit{B. cereus} strains such as \textit{ATCC} 4342 (29, 30). The sepal binding of GFP-PlyG\textsuperscript{BD} to RS1045, stands in distinct contrast to the whole-cell binding of \textit{B. anthracis} (Fig. 4d) and the polar-sepalt binding of \textit{ATCC} 4342 (Fig. 3e). Such binding patterns clearly reflect differences in the surface localization of \textit{B. anthracis}-like cell wall polysaccharides. Despite the unique binding of GFP-PlyG\textsuperscript{BD} to RS1045, the strain was nonetheless susceptible to the lytic action of PlyG in a liquid killing assay (Fig. 3e). Sensitivity to PlyG is a phenotypic specific to \textit{B. anthracis} and \textit{B. cereus} strains such as \textit{ATCC} 4342 (32) and, apparently, RS1045. Our findings here confirmed the presence of a \textit{B. anthracis}-like \textit{B. cereus} strain, RS1045, which also served as the source of one \textit{B. anthracis}-infective, worm gut bacteriophage (Wip4) identified in the present study.

**Bacterial flora of the worm gut environment.** Having identified \textit{B. anthracis} phages in PA earthworms, our next step was to assess the extent to which \textit{B. anthracis}-like organisms (like RS1045) were found in worms from another location—an outdoor garden bed in NYC. For this, we developed a novel three-tiered analytical method, distinct from the phage amplification- and bacterial culture-based methods used with PA worms. This method incorporated a metagenomic approach (28), and ultimately enabled specific access to \textit{B. cereus} s.l. organisms and phages in the worm gut microbiome.

Initially, the intestinal contents of 12 NYC worms were combined in a 3-ml PBS suspension and total DNA was extracted for construction of a plasmid vector-based library with insert sizes of 0.5 to 2 kb. Forty-seven DNA sequences were ultimately identified and assigned to a phylum, class, or genus (Fig. 4a) based on the most statistically significant similarity from a Blastn analysis. The total DNA pool from NYC worms was found to consist predominantly (62\%) of \textit{Gammaproteobacteria} sequences, including \textit{Shewanella}, \textit{Citrobacter}, \textit{Aeromonas}, \textit{Pseudomonas}, \textit{Salmonella}, \textit{Acinetobacter}, \textit{Thiomicrospira}, \textit{Klebsiella}, \textit{Enterobacter}, \textit{Marinomonas}, and others (Fig. 4b).
4a). *Aeromonas* spp. accounted for 20% of the sequences and was the most common genus identified (data not shown). While one sequence was most similar to the Gram-positive firmicute *Brevibacillus brevis*, no *B. cereus* s.l. homologs were observed. These findings differ slightly from a report for *E. fetida* from Long Island (28), in which 13% of the sequences were *Firmicutes* (although no *B. cereus* s.l. was reported).

*B. thuringiensis* and *B. mycoides* are the only previously described *B. cereus* s.l. organisms recovered from earthworms (in Denmark and Germany) (10, 14, 17).

Since the absence of *B. cereus* s.l. sequences from NYC worms could reflect a minority status in the worm gut, an initial enrichment for heat-resistant endospores was next performed prior to DNA isolation. As such, “spore-enriched” pools were generated by heating gut extracts at 95°C for 30 min to kill everything except spores; heated samples were then resuspended in BHI and grown at 37°C to promote the spore germination and outgrowth necessary for bacterial lysis, DNA recovery, and metagenomic library construction. Of 25 analyzed sequences, ~75% were now *B. cereus* s.l. group members, including *B. mycoides*, *B. pseudomycoides*, *B. weihenstephensis*, *B. cereus*, and *B. thuringiensis* (Fig. 4b). Surprisingly, 15% of the sequences were of non-spore-forming *Betaproteobacteria* that survived heat treatment.

The possibility of *B. anthracis*-like organisms being present as vegetative forms in the worm gut was addressed as part of the next enrichment strategy. Here, the sensitivity of vegetative *B. anthracis* and certain *B. cereus* strains to the lytic action of

FIG. 3. Susceptibility of RS1045 to the binding and lytic activities of the PlyG lysin. All images were obtained at 2,000× magnification. A phase-contrast micrograph of exponential-phase RS1045 is shown (a), with a corresponding fluorescence image (b), obtained with a 1-s exposure time, showing surface labeling with GFP-PlyG<sup>HD</sup>. Surface labeling of both *B. cereus* ATCC 4342 and *B. anthracis* ∆Sterne with GFP-PlyG<sup>HD</sup> is shown in fluorescence micrographs obtained with 1-s (c) and 0.1-s (d) exposures, respectively. (e) Sensitivity of RS1045 to the lytic activity of purified PlyG lysin. Aliquots of RS1045 were mixed with 100 μg of PlyG (■) or PBS alone as a control (▲), and the optical density was monitored for 15 min at room temperature in a SpectraMax Plus (Molecular Devices, Inc.) plate reader.
PlyG (32) was exploited prior to DNA isolation. PlyG treatment of gut extracts should specifically release only DNA from sensitive organisms into solution for subsequent metagenomic analysis. As expected, more than 90% of the inserts from the “PlyG-released” library were now from *B. cereus* s.l. organisms (Fig. 4c). Importantly, 19% of the inserts were 100% identical to a variety of *B. anthracis* loci. Thus, an organism similar to *B. anthracis* or *B. cereus* is a member of the NYC worm gut flora. These findings also show the specificity of phage lysins can be harnessed to focus metagenomic analyses on specific and/or rare organisms of interest.

**Bacteriophage flora of the worm gut environment.** Considering the presence of *B. anthracis*-like sequences, *B. anthracis*-infective phages should have been present in the NYC worms as well. To investigate this, worm intestinal extracts were first filtered to remove bacteria and retain free phages. When plated directly to a *B. anthracis* reporter, however, the filtrate yielded no PFU. Similarly, metagenomic DNA isolated from the filtrate also yielded no *B. anthracis* phage sequences (data not shown). While such findings could suggest the absence of *B. anthracis*-infective phages, they may simply be in an extreme minority or otherwise difficult to detect. A strategy to enrich for *B. anthracis*-infective phages was therefore used, whereby exponential phase /H9004 Sterne cultures were inoculated with worm gut extracts and incubated for 24 h. This infection cycle could amplify rare phages to detectable levels. Indeed, when NYC worm gut extracts were used to infect /H9004 Sterne overnight, the resulting supernatants became enriched for *B. anthracis*-infective phages (~10^6 PFU per ml). Sequence analysis of a metagenomic DNA library generated from phage-bearing supernatant revealed 39% of the inserts to be most similar to *B. anthracis* phages of /H9278 W siphovirus family, including Wβ and three distinct Wγ isolates (Fig. 4d) (11, 29). Interestingly, Wip4 (from PA) is genetically similar to the /H9278 W family as well. In all, these findings show phages infecting *B. anthracis* are indeed present in the NYC worms. Interestingly, the amplification step here also yielded sequences (51.9% of the total) similar to mobile genetic elements of Gram-negative organisms.

**DISCUSSION**

In the present study, the occurrence of *B. anthracis*-infective bacteriophages in earthworms recovered directly from the forest leaf litter in Pennsylvania was examined. Starting in 2003, phages infecting both *B. anthracis* and certain *B. cereus* strains were found in each of 12 different worms collected annually for 6 years over a 9-m radius around a central decaying tree stump. Interestingly, one phage species (Wip1) was replaced by another (Wip4) over time. RS1045 was found to be the host of Wip4 from 2006 to 2008 and could not be detected by PCR in the whole worm gut extracts recovered before 2006. The loss of spore-enriched (b), PlyG-released (c), and *B. anthracis* phage-enhanced (d) metagenomic DNA libraries are shown. The data are organized according to prokaryotic phyla, class, genus, species, or strain. Sequences that are similar to nothing in the database are classified as “unique.” Abbreviations: Bt, *B. thuringiensis*; weihensteph., *weihenstephanensis*; G, Gram-negative; Tn, transposon.
one phage and/or lysogen over another could reflect changing environmental conditions (i.e., soil humidity, temperature, and the nature of decaying organic matter in the soil) and an altered equilibrium in the worm gut microbiota. Nevertheless, a total of 72 E. fetida worms obtained over a 6-year period provided an environmental niche for at least two B. anthracis-infective phages.

The extreme host specificities of Wip1 and Wip4, suggested the presence of B. anthracis-like isolates (perhaps similar to B. cereus strains such as ATCC 4342 and CDC32805) in the intestines of worms from what is essentially an area where anthrax is not endemic. Indeed, RS1045, the Wip4 host, was found to be a B. cereus s.l. strain phylogenetically clustered with mammalian pathogens, including B. anthracis (based on its plcR-papR sequence). As with B. anthracis ΔSterne and B. cereus ATCC 4342, RS1045 expressed a B. anthracis-like surface polysaccharide structure and was sensitive to lysis after exposure to the PlyG lysin. RS1045 was unique, however, since the GFP-PlyGBD-binding location (and thus the location of a B. anthracis-like surface polysaccharide structure) was restricted to a septal position. Non-B. anthracis strains of B. cereus s.l. expressing a range of B. anthracis phenotypes and genotypes are present throughout the environment (12, 15, 21, 25, 36) and are likely the source of all B. anthracis-infective phages in areas where anthrax is not endemic, such as Pennsylvania and New York.

A second entirely distinct, metagenomic-based method was used in this work to supplement analysis of worm-associated B. cereus s.l. strains and phages. Here, the focus was on worms from a second nonendemic site: garden soil in NYC. Sequence analysis of 47 random clones from a metagenomic DNA library prepared from NYC worm intestines, revealed organisms of the Gammaproteobacteria as dominant intestinal forms. Similar results were found with worms from Long Island (28), Japan (34), and Germany (17). Although the percentage of Firmicutes in NYC worms was low compared to reports for Long Island worms, the organic matter on which worms feed can transiently influence the composition of intestinal flora (6). Initial decomposition processes favor Proteobacteria which can exploit soluble organic compounds (carbohydrates, amino acids, etc.), while a second successional phase favors Bacillus species which can degrade cellulose and hemicelluloses (6). The apparent ratio of Proteobacteria to Firmicutes can thus vary over time.

The initial absence of B. cereus s.l. sequences from NYC worm intestines was the impetus for developing three new methods to generate input DNA for metagenomic library construction. Here, the goal was to select for bacterial or viral subsets of interest. First, the ability of Bacillus and Clostridium species to form dormant endospores that resist temperatures which destroy other organisms was exploited. Although the resulting heat treatment method was only partially successful, the spore-enhanced metagenomic library was heavily biased toward B. cereus s.l. (based on analysis of 25 sequences). B. cereus s.l. was therefore the dominant spore-former in NYC worms. This method should provide an excellent means to access the diversity of spore-formers in any particular environment.

The absence of B. anthracis-like sequences in the spore fraction of NYC worms led us to develop a second method for generating input DNA for metagenomic library construction. Here, the focus was only on vegetative B. anthracis-like organisms sensitive to lysis by the PlyG lysin (32). PlyG is a cell wall hydrolase that selectively lyses B. anthracis and some B. cereus strains, even in complex samples containing a vast excess of lysis-resistant organisms. Treatment of washed worm gut extracts with PlyG was, therefore, expected to release only B. anthracis DNA and the DNA of other sensitive B. cereus s.l. organisms present in a vegetative state in worm guts. Of the 48 sequences examined from the “PlyG-released” metagenomic DNA library, >90% corresponded to B. cereus s.l. organisms, and 20% of the inserts were 100% identical to the B. anthracis chromosome. These findings indicate the presence of organisms very similar to B. anthracis (like some B. cereus strains) in the NYC worm intestinal tract niche. These findings also suggest lysins are useful for the construction of species-specific metagenomic libraries.

Given the presence of B. anthracis-like sequences in NYC worms, B. anthracis-infective phages should have been detected as well. To identify such phages, a B. anthracis phage-enhanced metagenomic library was created from the total viral pool of extracted worm guts. This “enhancement” was essentially attained after a 24-h cycle of phage amplification in the presence of a B. anthracis host. Based on sequences from 24 library members, the resulting “phage” DNA pool yielded a variety of mobile DNA elements, including phages, plasmids, and transposons. Significantly, ca. 40% of the sequences corresponded to a variety of inserts from the W family of B. anthracis-infective phages, including Fah, β, d’Herelle, Cherry, LSU, Porton, and USAMRIID (5, 11, 24, 29). The W family of phages are diagnostic for B. anthracis and have evolved from a common ancestor over decades of use in clinical labs (22). The identification of W sequences in a B. anthracis-phage enhanced metagenomic library is consistent with the presence W-sensitive B. cereus organisms, such as ATCC 4342 and CDC32805, in NYC worms.

Our findings suggest that B. cereus strains with limited B. anthracis-like phenotypes are present as spore and vegetative forms in earthworm intestines. To continue with our investigation of the complex ecology of B. cereus s.l. the next step will be to screen for such organisms and phages in worms from areas where anthrax is endemic and from anthrax carcass burial sites. In this manner, the question of whether worm intestines provide a reservoir for bona fide B. anthracis may be answered. Several observations from environmental surveys in areas of anthrax endemicity already infer a role for earthworms in the ecology of B. anthracis (9, 16, 26, 35). Most notably, increases in earthworm abundance and diversity are associated with soil and climatic conditions considered to be predictive of enzootic risks for anthrax. As such, slightly alkaline soils with elevated calcium levels and rich in organic matter favor both anthrax persistence (16, 35) and high earthworm densities (9, 26). In addition, anthrax outbreaks are also known to occur after rainy seasons or periods of flooding, when earthworms are driven to the soil surface. To establish a link between B. anthracis (including its vegetative and spore forms), earthworms and the occurrence of anthrax outbreaks will again require direct sampling at enzootic areas.
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REFERENCES


