Infection with Bacteroides Phage BV01 Alters the Host Transcriptome and Bile Acid Metabolism in a Common Human Gut Microbe

Graphical Abstract

Highlights

- Integration of BV01 in the tspO promoter causes genome-wide transcriptome change
- *B. vulgatus* bile salt hydrolase (BSH) activity is repressed by BV01 integration
- Transduction of BV01 appears to be host dependent because it is only observed in vivo
- Phage BV01 is a member of the Salyersviridae phage family

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In Brief
Integrative phages can have myriad effects on bacterial hosts. Campbell et al. find that integration of Bacteroides phage BV01 into *Bacteroides vulgatus* disrupts expression of TspO, reducing bile acid deconjugation. This is predicted to affect gut colonization and interactions with the human host. BV01 belongs to the Salyersviridae family.
Infection with Bacteroides Phage BV01 Alters the Host Transcriptome and Bile Acid Metabolism in a Common Human Gut Microbe

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SUMMARY

Gut-associated phages are hypothesized to alter the abundance and activity of their bacterial hosts, contributing to human health and disease. Although temperate phages constitute a significant fraction of the gut virome, the effects of lysogenic infection are underexplored. We report that the temperate phage, Bacteroides phage BV01, broadly alters its host’s transcriptome, the prominent human gut symbiont Bacteroides vulgatus. This alteration occurs through phage-induced repression of a tryptophan-rich sensory protein (TspO) and represses bile acid deconjugation. Because microbially modified bile acids are important signals for the mammalian host, this is a mechanism by which a phage may influence mammalian phenotypes. Furthermore, BV01 and its relatives in the proposed phage family Salyersviridae are ubiquitous in human gut metagenomes, infecting a broad range of Bacteroides hosts. These results demonstrate the complexity of phage-bacteria-mammal relationships and emphasize a need to better understand the role of temperate phages in the gut microbiome.

INTRODUCTION

The human gut is colonized by a dense and diverse microbial community comprised of bacterial, archaeal, and fungal cells as well as the viruses that infect them. This gut microbiome is vital for human health and development and is linked to an increasingly long list of disease states (Hills et al., 2019; Schmidt et al., 2018). Recent work has specifically implicated the gut phagome in disease, including inflammatory bowel disease (Norman et al., 2015), malnutrition (Reyes et al., 2015), AIDS (Monaco et al., 2016), colorectal cancers (Hannigan et al., 2016), and hypertension (Han et al., 2018). Broadly, gut phages act as important modulators of bacterial community structure (Hannigan et al., 2018; Khan Mirzaei et al., 2020; Moreno-Gallego et al., 2019) and metabolism (Hsu et al., 2019). Despite their apparent importance, little is known about how most gut-associated phages interact with their bacterial hosts (Mirzaei and Maurice, 2017).

Bacteroides is one of the most common and abundant bacterial genera in the distal human gut. The genus is known to degrade a diversity of complex carbohydrates (El Kaoutari et al., 2013; Salyers et al., 1977) and interact with host immune cells (Hickey et al., 2015; Shen et al., 2012). In a single human host, many Bacteroides species and strains coexist, competing for nutrients under changing environmental conditions caused by host diet (Tuncil et al., 2017), host metabolites (Ridlon et al., 2016), host immune system activities (Cullen et al., 2015; Planer et al., 2016), and phage predation (Porter et al., 2020). Moreover, horizontal gene transfer plays an important role in shaping the evolution and function of Bacteroides genomes (Coyne et al., 2014; Lange et al., 2016). How the diversity of Bacteroides strains in the human gut persists over time in such a dense, dynamic, and competitive environment is likely to have many reasons and is perhaps afforded by their highly plastic genomes.

Phage diversity and phage-host interactions in most commensal gut-associated bacteria, including Bacteroides species, is underexplored, but bioinformatics predictions of Bacteroides phages have been made (Krupovic and Forterre, 2011; Lange et al., 2016). Currently, the most abundant gut-associated phages are crAssphages (Dutilh et al., 2014; Koonin and Yutin, 2020; Yutin et al., 2018), a group of related lytic phages that infect Bacteroides intestinalis and potentially other species. CrAssphages demonstrate how traditional phage techniques (e.g., agar overlay plaque assays) are not reliable for Bacteroides hosts (Shkoporov et al., 2018), likely because of heterogeneity in capsular polysaccharide composition in isogenic cultures.
ATCC 8482

Bacteroides Phage BV01 Is a Prophage in B. vulgatus ATCC 8482

Bacteroides phage BV01 was partially predicted previously with the computational tool ProPhinder in the genome of B. vulgatus ATCC 8482 and deposited in the CLAssification of Mobile genetic Elements (ACLAME) database (Leplae et al., 2010; Lima-Mendez et al., 2008). Through comparative genomics and re-annotation of the host genome, we extended the predicted BV01 prophage to 58.9 kb (NCBI RefSeq: NC_009614.1; 3,579,765–3,638,687), which comprises 75 predicted open reading frames (ORFs) (Table S1). This larger predicted region is consistent with an earlier report of a prophage induced from this bacterial strain colonizing gnotobiotic mice (Reyes et al., 2013). BV01 encodes genes suggesting a temperate lifestyle with a putative phage repressor and anti-repressor as well as a holin-lysin-spanin operon for lytic release of phage progeny (Table S1; Figure 1A; Kongari et al., 2018; Young, 2014).

BV01 is detectable outside of host cells in the supernatants of in vitro cultures as a DNase-protected, double-stranded DNA (dsDNA) genome by shotgun sequencing (Figure 1A). Assembly of sequencing reads from free BV01 phage DNA results in a circular contig that spans the phage attachment site (attP). BV01 attT is identical to the left and right attachment sites (attL and attR, respectively), a pair of 25-bp direct repeat sequences (5’-GTCTAGTGGTTTGTGTTTGA-3’), suggesting that BV01 enters a circular intermediate before replication.

Routine detection of phage BV01 in DNase-treated, cell-free supernatants by PCR suggests that it is spontaneously induced. Furthermore, examination of the phage-to-host ratio by qPCR revealed that it remained approximately 1 throughout log and stationary phase host growth (Figure 1B), which is expected for phages that are spontaneously induced (Baugher et al., 2014; Bonanno et al., 2016). We have yet to identify in vitro induction conditions that increase phage BV01 production above this background level detectable by qPCR. Subinhibitory concentrations of the antibiotics norfloxacin, mitomycin C, tetracycline, and erythromycin; UV irradiation; or co-culture with gut-associated microbes do not increase BV01 production. Furthermore, attempts to visualize BV01 by transmission electron microscopy were not successful, likely because of the small number of phage particles produced.

An isogenic cured lysogen (ΔBV01) strain was constructed by allelic exchange with the corresponding chromosomal attB region of an uninfected B. vulgatus strain (Figure 1A). No apparent fitness effect of BV01 infection was detected in vitro because growth of the lysogen (wild-type [WT]) and cured lysogen strains (ΔBV01) are not significantly different (Figures 1C and 1D). This further supports the conclusion that spontaneous induction of BV01 occurs at a low rate and that sporadic cell lysis events do not have an effect on cell density.

Despite sequence evidence of BV01 phage particles in the culture supernatants, BV01-containing supernatants were never found to produce new infection. No lytic infection was detected using plaque assays (Kropinski et al., 2009) with 10 B. vulgatus isolates (Table S4), the B. vulgatus cured lysogen (ΔBV01), B. thetaiotaomicron, and 4 B. dorei isolates on four different growth media: tryptone-yeast extract-glucose broth (TYG), brain heart infusion (BHI) agar supplemented with 10% defibrinated horse blood (HB), BHI supplemented with hemin and menadione (BHI-HM) (Thornton et al., 2012), and supplemented TYG (TYG3a) (Goodman et al., 2009). Enrichment of infectious BV01 particles was attempted by passaging BV01-containing supernatants on B. vulgatus cured lysogen (ΔBV01) every day for 10 days (Shkoporov et al., 2018). However, enrichment was unsuccessful because subsequent plaque assays on naive cured lysogen did not yield any plaques. The same panel of 15 Bacteroides strains was tested for formation of lysogenic infection using non-concentrated or 200× concentrated BV01-containing supernatants. However, no novel infection was detected by PCR with primers designed to span the attachment site. To further test for new lyogenic infection, BV01 was tagged with a copy of tetQ, conferring tetracycline resistance, using allelic exchange. Five recipient strains, 4 B. vulgatus hosts and 1...
B. dorei host, were tagged with erythromycin resistance with pNBU2-\textit{bla} (Table S4). Double antibiotic selection found no new lysogens after attempted infection assays using BV01-\textit{tetQ} supernatants or 200x concentrated BV01-\textit{tetQ} supernatants or after 24-h co-culture of donor and recipient strains. We conclude that BV01 is a latent prophage \textit{in vitro}, which is further supported by transcriptional data showing that BV01 exists in a largely repressed state (Figure S1A), a state likely maintained by the predicted phage repressor (BVU_RS14475), the most highly transcribed gene in the BV01 prophage (Figure S1A).

The only infectious conditions that were identified for BV01 are in a gnotobiotic mouse model (Figure 2A). Within 1 day of gavage, BV01-\textit{tetQ} transductants were identified on doubly selective medium from mouse pellets from 4 of the 7 mice. Over the course of the 11-day experiment, transductants were
eventually observed in all animals in all cages. The average frequency of transduction ranged from 1.9 \( \times 10^{-3} \) to 3.6 \( \times 10^{-9} \) per animal. Of 26 transductant isolates that were colony purified, all were confirmed by PCR to have gained BV01 integrated at the known attB. Further, a representative transductant isolate was confirmed by whole-genome sequencing (Tables S2 and S3). These results support our hypothesis that an unknown mammalian host factor is required for novel BV01 infection.

To confirm that the release of DNase-protected BV01 genomes from host cells is a phage-encoded process, we sought to identify the phage integrase. BV01 encodes three genes with integrase domains (PF00589). We hypothesized that the gene BVU_RS14130, adjacent to the phage attachment site, was the most likely candidate for catalyzing integration and excision of BV01, as is common in phages (Casjens, 2003). A BVU_RS14130 deletion mutant (\( \Delta int \)) was constructed (Figure 1A), and its activity was assayed by PCR of paired cell pellets and culture supernatants (Figure 2B). Phage DNA was not detected in the supernatants of the \( \Delta int \) strain. Furthermore, the \( \Delta int \) mutant did not yield an amplicon for circularization of BV01. Expression of the int gene in trans complements the circularization and release phenotypes (Figure 2B). These results demonstrate that the integrase encoded by BVU_RS14130 is necessary for phage excision, circularization, and release from the host. Furthermore, they suggest that BV01 is an intact prophage that directs its own mobilization.

**Lysogeny with BV01 Alters the Host Transcriptome**

We hypothesized that lysogeny with a prophage such as BV01 could alter the activities of the *B. vulgatus* host. RNA sequencing (RNA-seq) of the *B. vulgatus* WT lysogen and the isogenic cured lysogen was performed to identify transcripts differentially regulated in response to lysogeny (Figure 3). Our expectation was that if prophage BV01 modulates bacterial activities, then some of those changes would be apparent as changes in host gene expression.

Analysis of the RNA-seq data revealed 115 host transcripts differentially regulated in response to lysogeny with BV01 (Figure 3A), 103 of which (89%) were upregulated in the cured lysogen (Table S5). Resequeencing of mutant genomes did not
reveal any secondary mutations likely to contribute to this global transcriptional response (Tables S2 and S3). These transcriptional changes occur across the host genome (Figure 3B). Functional analysis of these transcripts revealed that most function in metabolism and cellular processes and signaling (Figure 3C). However, pathway analysis using the Kyoto Encyclopedia of Genes and Genomes Pathway Database (Kanehisa and Goto, 2000) failed to yield pathway-level differences, which may be reflective of the level of annotation of the *B. vulgatus* genome. Taken together, these results indicate broad repression of a diverse array of host metabolic activities in response to prophage BV01, suggesting that it acts through one or more transcriptional regulators.

One possible explanation for the widespread transcriptomic response to BV01 lysogeny (Figure 3B) is that a phage product directly alters the transcriptional activity of host genes. BV01
encodes three candidate genes that might act in this way: a predicted phage repressor (BVU_RS14475), anti-repressor (BVU_RS14135), and sigma factor-like protein (BVU_RS14235) (Figure S1A). The putative phage repressor encoded by BVU_RS14475 is the most highly transcribed gene in the BV01 prophage, which might also interact with host promoters. Transcription of BVU_RS14135 and BVU_RS14235 is very low, so they are less likely to play a major role in the observed transcriptional response (Figure S1A). It is also possible that the observed transcriptional response to BV01 is the result of a host response to infection. A host-encoded, universal stress protein (uspA) homolog (BVU_RS16570) is upregulated in the BV01 lysogen (Table S5), but it is unknown whether that is a direct or indirect effect of infection.

**BV01 Disruption of the tspO Promoter Alters Bile Acid Metabolism**

Notably, integration of BV01 at attB is associated with a 23-fold downregulation of the adjacent downstream transcript (BVU_RS14490) encoding a predicted tryptophan-rich-sensory protein (TspO). We hypothesized that this downregulation is caused by disruption of the tspO promoter because BV01 integrates between the predicted promoter and the ORF (Figure 4A). A low level of expression of this gene is observed in the WT lysogen, perhaps a result of readthrough from phage transcripts. TspO is an intramembrane protein whose endogenous ligand is unknown but that is broadly implicated in metabolic regulation and stress response in other bacteria (Baisemão-Pires et al., 2011; Davey and de Bruijn, 2000; Yeliseev and Kaplan, 1999). Sequence conservation specifically in the unstructured loop (LP1) and the cholesterol recognition/interaction amino acid consensus (CRAC) sequence of TspO suggest that the B. vulgatus copy may function similar to other bacterial copies (Guo et al., 2015; Li et al., 2019a; Zeno et al., 2012; Figure 4B). Although TspO is conserved in many bacteria, archaea, and eukaryotes, it is considered an accessory protein. Indeed, not all gut-associated members of the family Bacteroidaives or the genus Bacteroides encode tspO (Figure 4C). Within Bacteroidaives, tspO is restricted to the clade that includes B. vulgatus, and among B. vulgatus strains, TspO is highly conserved (Figure 4D). Together, this suggests that TspO might play a specialized role in the lifestyle of B. vulgatus.

Given TspO’s important role in regulating cellular activities in other bacterial systems, we hypothesized that it may be responsible for some of the differential regulation observed in response to BV01 lysogeny. A tspO deletion mutant was constructed in the cured lysogen background (ΔBV01ΔtspO), and its transcriptome was sequenced alongside that of the WT and cured lysogen strains (Figure 3). The predicted tspO region extends beyond the differential expression observed in the cured lysogen (Figures 3A and 3B), suggesting that the small amount of tspO transcription in the BV01 lysogen exerts effects on the rest of the transcriptome (Table S5).

Transcripts differentially regulated between the BV01 WT lysogen and cured lysogen that returned to WT-like levels upon further deletion of tspO were classified as tspO-dependent transcripts (Figure 3C). TspO-dependent transcripts are not significantly different in a ΔBV01ΔtspO/WT comparison because tspO is absent or minimally active under both of these conditions. Of the 115 transcripts differentially regulated in response to BV01, 69 (60%) are tspO-dependent. Of these 69 tspO-dependent genes, 67 are downregulated by BV01 integration (i.e., positive fold change for ΔBV01/WT) and downregulated by tspO deletion (i.e., negative fold change for ΔBV01ΔtspO/ΔBV01), suggesting that tspO is largely involved in activation of downstream transcripts.

How TspO, a membrane protein, affects downstream transcription is unknown (Davey and de Bruijn, 2000; Liu et al., 2006; Yeliseev and Kaplan, 1999). However, we hypothesize that, in B. vulgatus, like in other bacterial models, TspO is involved in a signaling cascade, possibly by transporting unknown ligands across the membrane or by interacting with other membrane proteins (Davey and de Bruijn, 2000; Liu et al., 2006; Yeliseev and Kaplan, 1999). Consistent with TspO’s role in regulating stress, many tspO-dependent transcripts fall into the Clusters of Orthologous Groups (COG) category for post-translational modification, protein turnover, and chaperones, including several thioredoxins, peroxidasies, and protein chaperones. TspO-dependent transcripts also account for the majority of metabolic genes differentially regulated in response to BV01 (Table S5).

Two tspO-dependent transcripts that are downregulated in response to lysogeny with BV01 encode putative bile salt hydrolyses (BSHs; BVU_RS13575, 5.38-fold change, q < 10^{-100}, BVU_RS20010, 6.58-fold change, q < 10^{-200}) (Table S5). It was hypothesized that these transcriptional differences would reflect enzyme activities. To this end, B. vulgatus strains were grown in the presence of bile acids, and the deconjugation of those bile acids was measured by liquid chromatography-mass spectrometry (LC-MS) (Figure 5).

The LC-MS results show that the WT B. vulgatus lysogen does not significantly deconjugate glycocholic acid (GCA) to cholic acid (CA) and may exhibit modest deconjugation of taurocholic acid (TCA) (Figure 5; Figure S2). This agrees with a previous study that showed that B. vulgatus ATCC 8482 can deconjugate TCA but not GCA over a 48-h incubation (Yao et al., 2018). Importantly, CA is clearly detectable only in the cured lysogen background, consistent with the RNA-seq data (Figures 5A and 5B). This bile acid deconjugation phenotype is ablated with deletion of tspO, further supporting our hypothesis that tspO activates transcription of BSHs via an unknown mechanism, resulting in increased enzymatic activity (Figure 5C).

Finally, resequencing of mutant genomes revealed no additional mutations likely to cause the restoration of BSH activity (Tables S2 and S3).

To see whether tspO-disrupted lysogens occur in natural human gut microbiomes, read mapping from 246 healthy human gut metagenomes was performed. Starting with reads that mapped to tspO in the reverse orientation, read mates were checked for mapping to phage BV01 and its relatives (Figure 6). All samples had reads mapping to tspO, with an average of 0.0004% of metagenome reads mapping, indicating that the corresponding population of B. vulgatus and B. dorei encoding tspO is relatively abundant (Figure S3A). Incidence of tspO-associated with BV01 or a related phage is also common; 13.3% of samples (n = 34) contained read pairs mapping to tspO and an adjacent
prophage. Within an individual microbiome, incidence of tspO-disrupted lysogens appears to be rare, usually comprising 3% or less of the combined B. vulgatus and B. dorei population, although, for some individuals, this incidence rate can be more than 10% (Figure S3B). Together, these data suggest that BV01 and other phages’ effects on downstream phenotypes via tspO are likely quite common among humans, although these lysogens comprise the minority of the overall microbiome.

**BV01 Represents the Proposed Virus Family Salyersviridae**

While searching for potential new hosts for BV01, its predicted 25-bp attB was queried against 154 gut-associated Bacteroidales genomes. It was found that all Paraprevotella and Bacteroides genomes had at least the first 21 bases of the attachment site conserved (Figure S3B). In 20 genomes, two copies of the att site were found, and all were associated with putative prophages. In all instances, the putative attL and attR sites are direct repeats, as is true for BV01. Importantly, only B. vulgatus lysogens encode tspO (Figure 4); other prophages and attB sites occur in alternate genomic contexts. Alignment of these putative prophage-associated att sites finds that the first 22 bp are always conserved and that an additional 3 bp are variable among lysogen genomes (Figure 6B).

Integration at the same att site suggests that these prophages are genetically related. To assess relatedness, the Virus Classification and Tree Building Online Resource (VICTOR) (Meier-Kolthoff and Göker, 2017) was used to build a genome tree from all of the identified prophages (Figure 6C), and OPTSIL clustering (Göker et al., 2009) was used to predict taxonomic groups, which we named Salyersviridae, Salyersvirinae, and Salyersvirus (Table 1). Taxonomic clustering also defined phage species, four of which have more than one member. The species containing BV01 also contains BV02, a prophage 99.8%
nucleotide identical to BV01. BV02 occurs in *B. vulgatus* VPI-4245, a strain cross-listed as ATCC 8482, suggesting that the BV01 prophage is ancestral to the strain lineage. Interestingly, the phage species BX01 has representatives in two distantly related hosts, *Bacteroides xylanisolvens* XB1A (phage BX01) and *B. thetaiotaomicron* 1_1_6 (phage BT01), suggesting that at least some salyersviruses have broad host ranges. Phage BC01 was placed outside of the subfamily Salyersvirinae, which is consistent with its considerable sequence divergence (Figure S4). Further examination of the BC01 attL and attR sites found that the 22-bp consensus sequence is only a portion of the full 69-bp repeat flanking the BC01 prophage, which further supports placement of BC01 outside of Salyersvirinae. Also included in the analysis were three known *Bacteroides*-infecting lytic phages: B124-14, B40-8, and crAssphage. OPT-SIL taxonomic clustering placed phages B124-14 and B40-8 in the family Salyersviridae. This is consistent with an independent network analysis using vConTACT 2.0 (Bin Jang et al., 2019) and existing NCBI RefSeq viral genomes, which creates a cluster consisting of only BV01, B124-14, and B40-8. Similarities between Salyersviridae prophages and the two lytic phages are not detectable at the nucleotide sequence level (Figure S4A) but is likely driven by similarities in several proteins, including homologs to the predicted lysin, excisionase, and single-stranded DNA (ssDNA) binding protein from BV01 (Figure S4B).

To check that the proposed Salyersviridae clade was comprised of active phages, we searched for evidence of activity in culture and in wastewater samples. Sequencing and PCR were used to test the activity of six additional salyersviruses (Figure S5). Phages BV02, BV05, and BV06 were found to be released from host cells in culture, whereas BV03 is likely inactivated by genomic rearrangement (Figure S5D). We did not observe activity for BV04 or BO03 from *B. ovatus* ATCC 8483, both of which appear complete based on synteny with intact salyersviruses. This may be the result of inactivating mutations or not being induced under the growth conditions tested (Figure S5D).

Furthermore, we used read mapping to search for evidence of all 20 Salyersviridae phages in a wastewater metavirome.
(Figure S5E). Reads mapped to all Salyersviridae genomes; however, it may be the result of sequence conservation within the family because reads often accumulate at the most conserved regions of each phage genome. Further, very little read mapping occurred in the distal portion of the inactivated prophage BV04, which resembles the host chromosome more than the phage sequence, indicating that virome processing removed most cellular DNA prior to sequencing (Figure S5E).

De novo assembly of the individual wastewater viromes finds contigs that align with high identity but imperfectly to each Salyersviridae phage, supporting the findings seen by read mapping and suggesting that the real diversity of the family Salyersviridae is far greater than what has been observed to be integrated in cultured host genomes so far. This analysis found that phages infecting *B. vulgatus* are more abundant than other Salyersviridae phages based on maximum normalized read coverages. A similar comparison concludes that most individual temperate Salyersviridae phages are active prophages, confirmed by sequencing and/or PCR, indicated by black dots; prophages confirmed to have been inactivated by genome rearrangement are indicated by an X; prophages that were tested for activity with inconclusive results are indicated by asterisks (Figure S4).
Repression of bile acid deconjugation as a consequence of BV01 integration is particularly relevant in the context of the mammalian gut. Mammals secrete conjugated primary bile acids into the small intestine, where they reach concentrations as high as 1 mM (Northfield and McColl, 1973); although the majority of bile acids secreted into the small intestine are reabsorbed, they can still accumulate to concentrations of 0.2–1 mM in the colon (Hamilton et al., 2007). Although bile acids are broadly capable of damaging lipid membranes, Bacteroides species are generally considered bile resistant (Citron et al., 1990), the mechanism of which is unknown. Bile acid deconjugation is a common activity encoded by gut-associated microbes, although its direct benefit to those microbes is unclear. Microbial modification of the bile acid pool can be linked to beneficial changes in the human host metabolism (Joyce et al., 2014; Yao et al., 2018) and varied epithelial susceptibility to viral pathogens (Grau et al., 2020). The link between BV01 and bile acid metabolism suggests a so far undescribed mechanism by which gut phages might influence mammalian host phenotypes.

Here, bile acid deconjugation in B. vulgatus is dependent on a putative TspO. Bacterial TspOs are important for regulating metabolic switches and stress regulation in at least three diverse systems (Balsemão-Pires et al., 2011; Davey and de Bruijn, 2000; Yeliseev and Kaplan, 1999), but the mechanism of action for the protein is unknown. The crystal structure of TspO shows a periplasm-facing binding pocket distinct from the intramembrane cholesterol recognition consensus sequence, which may bind or degrade porphyrins (Guo et al., 2015; Li et al., 2015a;...
Zeno et al., 2012). The porphyrin degrading and cholesterol transporting functions of TspO, however, have been disputed (Li et al., 2015b; Rone et al., 2012). Despite this, it is notable that cholesterol is structurally similar to bile acids, being their biosynthetic precursor. In at least one other gut-associated microbe, tspO is upregulated by bile acids, suggesting that TspO may be involved more broadly in bile acid metabolism in gut microbes (Devendran et al., 2019). The regulatory link described here between bile acid hydrolysis and TspO suggests a hypothesis where the B. vulgatus TspO might be a sensor and regulator of bile acid interactions.

Induction of BV01 from its integrated state and infection of new hosts remains enigmatic. Prophage induction is canonically linked to stress-dependent pathways, as in lambdoid phages that respond to DNA damage via RecA-dependent cleavage of the CI repressor protein (Roberts et al., 1978). It is possible that prophages in Bacteroides hosts respond to alternative stimuli, as is the case for CtnDOT, a well-studied Bacteroides conjugative transposon whose excision is inducible only by tetracycline (Moon et al., 2005). Neither DNA damage nor antibiotics induce prophage BV01 in vitro, so all experiments here relied on an apparently low rate of spontaneous prophage induction. Similarly, no infection conditions or susceptible hosts have been identified for BV01 in vitro. We demonstrate that BV01 is transducible in a gnotobiotic mouse model, suggesting that an unknown mammalian host factor is required for novel BV01 infection. Enigmatic infection dynamics may be a result of the phase-variable polysaccharide capsule; recent work suggests that heterogeneity in capsule composition hinders phage infection on population scales (Porter et al., 2020). Indeed, it has long been observed that finding phages in Bacteroides using traditional techniques is difficult or impossible for most host strains (Puig and Girone’s, 1999; Salyers et al., 2000), making traditional techniques is difficult or impossible for most host strains (Puig and Girone’s, 1999; Salyers et al., 2000), making traditional techniques difficult or impossible for most host strains (Puig and Girone’s, 1999; Salyers et al., 2000), making traditional techniques difficult or impossible for most host strains (Puig and Girone’s, 1999; Salyers et al., 2000), making traditional techniques difficult or impossible for most host strains (Puig and Girone’s, 1999; Salyers et al., 2000), making

Finally, phage BV01 is the first explored representative of the broad family Salyersviridae, which spans an entire host genus and includes lytic and temperate members. Salyersviridae is common and diverse among natural human samples but rare within individuals, suggesting that lysogenization may confer frequency-dependent advantages to the bacterial host. The genetic context of non-B. vulgatus Salyersviridae lysogens remains unexplored and likely underlies different phage-host interactions. The absence of tspO in these other host systems may provide the ideal background for studying more direct effects on the bacterial host. Certainly, other interactions between BV01 and its host remain to be studied, although they were overshadowed here by the enormous effects of tspO. Future studies should also examine the role of Salyersviridae phages on bacterial host fitness and evolution (Oh et al., 2019; Zhao et al., 2019) because these phages likely have important ripple effects throughout the microbiome and on the mammalian host that remain to be elucidated.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:
SUPPORTING CITATIONS

The following references appear in the Supplemental Information: Caporaso et al. (2011), Eggert and Gagnon (1933), Johnson (1978), Shoemaker et al. (2001), Simon et al. (1983), Whittle et al. (2001).

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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Bacterial and Virus Strains** | | |
| Bacteroides vulgatus ATCC 8482 | ATCC | ATCC 8482 |
| B. vulgatus Δtdk | This paper | N/A |
| B. vulgatus Δint | This paper | N/A |
| B. vulgatus ΔBV01 | This paper | N/A |
| B. vulgatus ΔBV01ΔtspO | This paper | N/A |
| B. vulgatus ΔBV01, attN::pNBU2-bla-ermGb | This paper | N/A |
| B. vulgatus BV01-tetQ, attN::pNBU2-bla-cfx | This paper | N/A |
| B. vulgatus ΔBV01, attB::BV01-tetQ, attN::pNBU2-bla-ermGb | This paper | N/A |
| B. vulgatus VPI-4245 | Abigail Salyers | N/A |
| B. vulgatus VPI-2365 | Abigail Salyers | N/A |
| B. vulgatus VPI-6186 | Abigail Salyers | N/A |
| B. vulgatus VPI-5710 | Abigail Salyers | N/A |
| B. vulgatus DH4096S | Abigail Salyers | N/A |
| B. ovatus ATCC 8483 | ATCC | ATCC 8483 |
| **Chemicals, Peptides, and Recombinant Proteins** | | |
| Taurocholic acid sodium salt hydrate, ≥ 95% | Sigma-Aldrich | CAS #345909-26-4 |
| Glycocholic acid hydrate, ≥ 97% | Sigma-Aldrich | CAS #1192657-83-2 |
| **Critical Commercial Assays** | | |
| Nextera XT Library Preparation Kit | Illumina | Cat. #FC-131-1096 |
| Nextera XT Index Kit | Illumina | Cat. #FC-131-1002 |
| NEBNext Ultra II FS Kit for Illumina | New England Biolabs | E7805L |
| RNeasy Kit | QIAGEN | Cat. #74104 |
| **Deposited Data** | | |
| Mutant genome DNA sequencing, see Table S3 | This paper | PRJNA655911 |
| RNA sequencing, see Table S3 | This paper | PRJNA622597 |
| Wastewater virome DNA sequencing, see Table S3 | This paper | PRJNA622299 |
| Bacterial genomes, see Table S3 | This paper | PRJNA622758 |
| Phage DNA sequencing, see Table S3 | This paper | PRJNA655697 |
| **Experimental Models: Organisms/Strains** | | |
| Germfree C57BL/6J mice | UCR Gnotobiotic Facility | N/A |
| **Oligonucleotides** | | |
| Primers for strain construction, see Table S4 | This paper | N/A |
| Primers for integrase activity assays, see Table S4 | This paper | N/A |
| Primers for detection of Salyersviridae phages, see Table S4 | This paper | N/A |
| **Recombinant DNA** | | |
| pKNOCK-bla-ermGb | Koropatkin et al., 2008 | N/A |
| pExchange-tdkBV | This paper | N/A |
| pNBU2-bla-ermGb | Koropatkin et al., 2008 | N/A |
| pNBU2-int | This paper | N/A |
| pNBU2-bla-cfx | This paper | N/A |

(Continued on next page)
RESOURCE AVAILABILITY

Lead Contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Dr. Patrick Degnan (patrick.degnan@ucr.edu).

Materials Availability
Bacterial strains and plasmids generated in this study are available upon request to the Lead Contact, Dr. Patrick Degnan (patrick.degnan@ucr.edu).

Data and Code Availability
Trimmed DNA sequencing reads from mutant genomes generated in this study are deposited in the NCBI SRA under PRJNA655911. Trimmed RNA sequencing reads from this study are deposited in the NCBI SRA under PRJNA622597. Trimmed wastewater virome DNA sequencing reads are deposited in the NCBI SRA under PRJNA622299. Ten new assembled B. vulgatus genomes are deposited in NCBI GenBank under PRJNA622758. Untrimmed phage DNA sequencing reads from culture supernatants are deposited in the NCBI SRA under PRJNA655697. All NCBI BioSample Accession numbers are listed in Table S3. The code for annotation of bacterial genomes is available at https://github.com/phdegnan/MICROBIALOMICS.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Bacterial Strains
A list of bacterial strains used in this work can be found in the Key Resources Table and Table S4. Escherichia coli S17-1 λ pir was grown aerobically in Lysogeny Broth (LB; 10 g/L tryptone, 5 g/L yeast extract, 5 g/L NaCl) at 37°C under agitation. Bacteroides strains were cultured anaerobically in a vinyl anaerobic chamber using 70% N2, 20% CO2, and 10% H2 gas mixture (Coy Laboratory Products, Grass Lake, MI). All Bacteroides cultures were grown on Difco Brain Heart Infusion (BHI) agar supplemented with 10% defibrinated horse blood (BHI-HB; Quad Five, Ryegate, MT), or in tryptone-yeast-extract-glucose broth (TYG; 10 g/L tryptone, 5 g/L yeast extract, 2 g/L glucose, 500 mg/L cysteine free base, 100 mM pH 7.2 potassium phosphate, 20 mg/L MgSO4 · 7H2O, 400 mg/L NaHCO3, 80 mg/L NaCl, 100 mg/L resazurin, 0.0008% CaCl2, 0.4 μg/L FeSO4, 1 μg/L menadione, 0.2 mM histidine, 1.9 μM hematin) at 37°C. When necessary, ampicillin (100 μg/mL), gentamicin (200 μg/mL), erythromycin (25 μg/mL), 5’-flourodeoxyuridine (F UdR; 20 μg/mL), or tetracycline (2 μg/mL) were supplemented in the media.

Mice
Germfree C57BL/6J mice were used in this study. With no a priori reason to expect age to influence transduction rates, animals ranged from 7 weeks to nearly 12 months old. Animals were maintained in flexible plastic gnotobiotic isolators with a 12-hr light/dark cycle. Animals caged individually (n = 1, female) or in pairs of litter mates (n = 6, males) were provided with standard, autoclaved mouse chow (5K67 LabDiet, Purina) ad libitum. All experiments using mice were performed using protocols approved by the University of California Riverside Institutional Animal Care and Use Committee.
METHOD DETAILS

Cloning and Mutagenesis

All primers used to construct genetic mutants are listed in Table S4. Markerless deletion mutants in *B. vulgatus* were achieved by allelic exchange using a system analogous to that developed in *B. thetaiotaomicron* (Koropatkin et al., 2008) and confirmed by PCR and whole genome sequencing. The *tdk* gene (BVU_RS09305), encoding thymidine kinase, was deleted from *B. vulgatus* ATCC 8482 by allelic exchange, conferring resistance to the toxic nucleotide analog FUdR. Cloning was performed as described by (Degnan et al., 2014). Briefly, the 3.5 Kb regions flanking either side of *tdk* were amplified with Kapa HiFi Taq MasterMix (Kapa Biosystems, Wilmington, MA) and joined by splicing overlap exchange (SOE) PCR. The SOE product was purified, restriction digested, and ligated into the suicide vector pKNOCK-*bla*-ermG₄ in *E. coli*, and conjugated into *B. vulgatus*. Single recombinant merodiploids were selected for on BHI-HB supplemented with gentamicin and erythromycin, and double recombinant deletion mutants subsequently selected for on BHI-HB with FudR. The counterselectable suicide vector pExchange-tdkBV was constructed by amplifying *tdk* from *B. vulgatus* and cloning the *tdk* amplicon into pKNOCK-*bla*-ermG₄ by the same methods used to clone the SOE product above.

Subsequent deletions were accomplished similarly as described for *tdk*, except using pExchange-tdkBV and flanking regions of ~1 Kb to create the deletion alleles (*tspQ, int*). For deletion of the entire BV01 prophage, an empty attachment site (*attB* and the flanking 800 bp were cloned from *B. vulgatus* VPI-4506, which is 99.9% nucleotide identical to the analogous regions flanking BV01 in *B. vulgatus* ATCC 8482.

Complementation of the BV01 integrase (BVU_RS14130) was accomplished by cloning the gene and its native promoter into the integrative plasmid pNBU2-*bla*-ermGb, which has a single integration site in the *B. vulgatus* genome (*attN*; RefSeq: NC_009614.1; 2,710,334-2,710,346). This construct was conjugated into *B. vulgatus* and transconjugants selected for on BHI-HB supplemented with gentamicin and erythromycin (Degnan et al., 2014).

The BV01-*tetQ* strain was constructed by inserting *tetQ* from pNBU2-*bla*-tetQ immediately downstream of the stop codon of BVU_RS14265, upstream of a predicted transcriptional terminator. As was done for deletion constructs, the desired region was constructed on the pExchange-tdkBV plasmid and moved into the wild-type *B. vulgatus* strain by allelic exchange. First, a ~2 Kb region surrounding the BVU_RS14265 stop codon was amplified in two pieces with SOE primers designed to insert adjacent SpeI and BamHI cut sites downstream of the stop codon and ligated into pExchange-tdkBV. This construct was confirmed by Sanger sequencing before *tetQ* and its promoter were amplified from CtnDOT, and ligated into the SpeI and BamHI cut sites. Tetracycline was used to select for mutants, and release of BV01-*tetQ* phages confirmed by PCR.

Select mutant strains were confirmed by whole genome sequencing and analyzed with Breseq (Deatherage and Barrick, 2014) aligned to the wild-type *B. vulgatus* ATCC 8482 genome (RefSeq: NC_009614.1), and summarized in Table S2.

Bacterial Growth Rate

Triplicate 200 μL cultures of *B. vulgatus* strains were grown in TYG medium and their optical densities (OD) monitored at a wavelength of 600 nm at 30 min intervals. Doubling times were calculated from cells in early to mid-log phase using the least-squares fitting method on http://www.doubling-time.com/compute.php.

Genome Sequencing

Cells were pelleted from 5 mL overnight culture in TYG by centrifugation at 4,000 g for 5 min at 4°C, resuspended in 0.5 mL TE buffer (10 mM Tris, 1 mM EDTA), and lysed by adding sodium dodecyl sulfate (SDS) and proteinase K (GoldBio, Olivette, MO) to final concentrations of 0.07% and 300 μg/mL, respectively, and incubating for 2 hr at 55°C. Cellular material was removed by washing twice in an equal volume of buffered phenol, phenol-chloroform-isoamyl alcohol (VWR, Radnor, PA), and DNA precipitated with SDS and 1 mg/mL proteinase K for 2 hr at 55°C. DNA was further isolated using the same phenol-chloroform-isoamyl alcohol extraction and ethanol precipitation procedures as for cellular DNA.

DNA libraries were constructed with the Nextera XT Library Preparation Kit and Index Kit (Illumina, San Diego, CA) or the NEBNext Ultra II FS kit for Illumina (New England Biolabs, Ipswich, MA). DNA libraries were pooled and sequenced on an Illumina HiSeq5000, HiSeq2500 and MiSeq instruments. Fastq files were generated from demultiplexed reads with bc2fastq Conversion Software (Illumina, San Diego, CA). Reads were trimmed and assembled using the A5ud pipeline (Riley et al., 2017). Sequencing methods and assembly data are summarized in Table S3. Reads were mapped to the *B. vulgatus* ATCC 8482 reference genome (RefSeq: NC_009614.1) using bwa in paired end mode (Li and Durbin, 2009) with default parameters to generate Figure 1A.

Genome Annotation
Annotation of cellular genomes was accomplished with a custom Perl script (https://github.com/phdegnan/MICROBIALOMICS) that calls protein coding genes with Prodigal (Hyatt et al., 2010) and RNA coding genes with trNAScan-SE (Lowe and Eddy, 1997), Rnammer (Lagesen et al., 2007) and Infernal (Nawrocki and Eddy, 2013). Functional predictions are assigned by searching against Kyudo Encyclopedia of Genes and Genomes (Kanehisa and Goto, 2000), Cluster of Orthologous Genes (Tatusov et al., 2003), Pfam (Bateman et al., 2004), and TIGRFAM (Haft et al., 2003) databases, and subCELlular LOcalization predictor (Yu et al., 2014) is used to predict cellular localization.

For phage genomes, open reading frames were called by Prodigal and the gene calling tool within Artemis (Carver et al., 2012). Functional predictions were made as above except with relaxed search parameters (cut_nc in hmmscan), plus using Basic Local Search Alignment Tool (Mount, 2007) with the GenBank virus database (Benson et al., 2009), Phyre2 (Kelley et al., 2015) to identify conserved protein folds, SignalP 5.0 (Almagro Armenteros et al., 2019) to predict signal peptides, and iVireons (Seguritan et al., 2012) to predict structural proteins, and manually combining and comparing results.

In Vivo Transduction
Individually antibiotic resistance marked bacterial strains were grown for ∼20h in TYG medium with appropriate antibiotics and frozen at −80°C in anaerobic cryovials. Cell viability was tested by plating and viable CFU counts were used to combine equal parts of the B. vulgatus cured lysogen tagged with pNBu2-bla-ermGb and B. vulgatus BV01-tetQ tagged with pNBu2-bla-cfx. Approximately 4 × 10^6 CFUs of the combined strains were administered to each animal by oral gavage. Fecal samples were collected on days 1, 3, 7 and 11 from each animal. Fecal pellets were processed by adding 500μl of TYG+20% glycerol to each tube and vigorously shaking in a beadbeater without beads for 1 m 30 s. Fecal slurries were spun down at 2,000 x g for 1 s, followed by serial dilution on selective media (BHI+Tet+Gn, BHI+Erm+Gn, BHI+Erm+Tet+Gn) to determine CFUs. Animals were sacrificed on day 11 following final fecal collection.

Integrate Activity Assays
Integrate activity was assayed through PCR of DNase-treated supernatant DNA. Briefly, free phage DNA was prepared as for DNA sequencing, and amplified with Kapa HiFi Taq MasterMix with primers specific to BV01 (BVU_RS14350) or spanning the circularized attP (Table S4). Free phage DNAs were checked for the presence of contaminating cellular DNA by amplifying the 16S rRNA gene with universal primers. Amplicons were cleaned with a QIAGEN PCR Cleanup kit (Hilden, Germany) and run on an agarose gel in 0.5X Tris-borate-EDTA buffer at 70V alongside 1 Kb ladder (New England BioLabs, Ipswich, MA) or GeneRuler Express DNA Ladder (Thermo Scientific, Waltham, MA) and stained with GelRed (VWR, Radnor, PA). Amplicons generated with attP-flanking primers were sequence confirmed by Sanger sequencing performed by ACGT, Inc (Wheeling, IL).

Transcriptomics
B. vulgatus was grown overnight in 5 mL TYG medium. Each culture was pelleted (4,000 x g for 5 min at 4°C), supernatant decanted, and washed in an equal volume of TYG three times. Cells were normalized to an OD600 of ∼0.3 (∼2.9x10^8 CFU/mL) and used to inoculate cultures in 10 mL TYG at a final dilution of 1:1000 in biological triplicate. Cell growth was monitored and cells were harvested at an OD600 of ∼0.4 (3.9x10^6 CFU/mL). Total RNA was prepared with a QIAIEN PCR Cleanup kit (Hilden, Germany) and treated on-column with RNase-free DNase (QIAGEN, Hilden, Germany). RNA was quantitated with a Qubit 2.0 fluorometer (Thermo Fisher, Waltham, MA) and stored at −80°C.

RNA was submitted to the W. M. Keck Center for Comparative and Functional Genomics at the University of Illinois at Urbana-Champaign for quality analysis, rRNA depletion with the RiboZero Bacteria kit (Illumina, San Diego, CA), library construction with the TruSeq Stranded mRNASeq Sample Prep kit (Illumina, San Diego, CA), and sequencing on an Illumina NovaSeq 6000 with the NovaSeq S4 reagent kit. Fastq files of multiplexed reads were prepared with the bcl2fastq v2.20 Conversion Software (Illumina, San Diego, CA).

RNaseq reads were quality filtered and trimmed with trim_galore v0.4.4 (https://www.bioinformatics.babraham.ac.uk/projects/trim_galore). Rockhopper (McClure et al., 2013) was used to calculate individual transcript Expression and pairwise q-values. Fold-change was calculated in Excel as Expression A/Expression B if A > B, or as -(Expression B/Expression A) if B > A. Differentially expressed transcripts between the isogenic mutants were defined as those ≥ 2-fold change and q ≤ 0.01. Trimmed reads were mapped to the B. vulgatus ATCC 8482 reference genome (RefSeq: NC_009614.1) using bwa in single end mode (Li and Durbin, 2009) with default parameters to generate coverage curves.

Bile Salt Deconjugation Assay and LC-MS
B. vulgatus strains were inoculated in TYG liquid supplemented with 50 μM glycocholic acid (GCA) or 50 μM taurocholic acid (TCA) and allowed to grow for 16 or 48 hr, respectively. B. vulgatus ATCC 8482 had previously been shown to completely hydrolyze TCA in 48 hr (Yao et al., 2018); the 16 hr time point was chosen to ensure incomplete hydrolysis to better detect increased activity. B. vulgatus ATCC 8482 was also unable to hydrolyze GCA in 48 hr (Yao et al., 2018); the 48 hr time point was chosen to ensure that any increase in hydrolysis would be apparent. Growth cultures were brought to a pH 2.0-3.0 with 10 N hydrochloric acid, centrifuged for 5 min at 4,000 x g, and the pellets discarded. Bile acids were isolated by solid phase extraction over Sep-Pak
tC18 500 mg cartridges (Waters Corp., Milford, MA). Cartridges were preconditioned by serial washes with 6 mL hexane, 3 mL acetone, 6 mL methanol, and 6 mL water (pH = 3.0). Acidified supernatants were loaded before washing with 3 mL 40% methanol. The column was allowed to dry, then bile acids eluted in 3 mL methanol. Samples were evaporated under nitrogen, resuspended in 20 μL methanol, and centrifuged before analysis by liquid chromatography-mass spectroscopy (LC-MS).

LC-MS for all samples was performed on a Waters Aquity UPLC coupled with a Waters Synapt G2-Si ESI MS. Chromatography was performed using a Waters Cortecs UPLC C18 column (1.6 μm particle size) (2.5 mm x 50 mm) with a column temperature of 40°C. Samples were injected at 1 μL. Solvent A consisted of 95% water, 5% acetonitrile, and 0.1% formic acid. Solvent B consisted of 95% acetonitrile, 5% water, and 0.1% formic acid. The initial mobile phase was 90% Solvent A, 10% Solvent B and increased linearly until the gradient reached 50% Solvent A and 50% Solvent B at 7.5 min. Solvent B was increased linearly again until it was briefly 100% at 8.0 min until returning to the initial mobile phase (90% Solvent A, 10% Solvent B) over the next 2 min. The total run was 10 min with a flow rate of 10 μL/min. MS was performed in negative ion mode. Nebulizer gas pressure was maintained at 400°C and gas flow was 800 L/hour. The capillary voltage was set at 2,000 V in negative mode. MassLynx was used to visualize and analyze chromatographs, including calculating areas under the curve (AUC). Ratios of AUCs are presented in Figure 5C.

Taxonomic nomenclature
The family, subfamily, and generic viral taxa names were chosen to honor the microbiologist Abigail A. Salyers, who made significant contributions to understanding the functions and genetics of human gut anaerobes and the importance of their mobile genetic elements.

Wastewater collection, processing, and viromics
From the Urbana and Champaign Sanitary District Northeast Plant (Urbana, IL), 1 L of unprocessed wastewater was collected at each of three time points: May 25, 2016, June 23, 2016, and October 3, 2016.

Wastewater samples were transported on ice, immediately centrifuged at 2,500 × g for 10 min at 4°C and filtered through a 0.4 μm polyethersulfone filter to remove large particulate and cellular matter. The sample was split into three aliquots and processed three ways. One aliquot was not processed further (F). Another aliquot was filtered a second time through a 0.22 μm polyethersulfone filter (DF). The last aliquot was washed three times with an equal volume of chloroform (FC). All aliquots were concentrated 100-fold and virome DNA was isolated from each as described for genome sequencing of phages.

DNA libraries of virome DNA were prepared using the same methods as described for phage genome sequencing and were sequenced on an Illumina HiSeq 2500 sequencer with a HiSeq v4 SBS sequencing kit (Illumina, San Diego, CA) producing 2 × 160-bp paired-end reads. Fastq files of demultiplexed reads were generated with the bcl2fastq v2.17.1.14 Conversion Software (Illumina, San Diego, CA). Reads were trimmed and quality filtered using Trimmomatic 0.38 (Bolger et al., 2014) and assembled with metaSPAdes v3.13.0 using default parameters (Nurk et al., 2017). Sequencing and assembly data for wastewater viromes is summarized in Table S4. Read mapping to phage genomes was performed with bwa in paired end mode (Li and Durbin, 2009).

Human Microbiome Project Healthy Human Subjects Study samples were downloaded with portal_client (Table S6). Read mapping was performed with bwa (Li and Durbin, 2009).

QUANTIFICATION AND STATISTICAL ANALYSIS
Statistical tests, number of events quantified, standard deviation of the mean, and statistical significance is reported in figure legends. ANOVA and Tukey HSD tests were run on http://vassarstats.net/anova1u.html.
Supplemental Information

Infection with Bacteroides Phage BV01 Alters the Host Transcriptome and Bile Acid Metabolism in a Common Human Gut Microbe

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Figure S1. Transcriptional activity at BV01 attachment site and confirmation of free BV01 integrase amplicons.
Related to Figures 1, 2, and 3. (A) RNAseq reads from wild-type lysogen (WT) and cured lysogen (ΔBV01) strains were mapped to the region, and coverage was normalized to the total number of reads mapping to the genome. Locations of the phage marker gene (BVU_RS14350) and three phage transcriptional regulators are indicated: the putative phage repressor (BVU_RS14475), anti-repressor (BVU_RS14135), and an RNA polymerase sigma factor (BVU_RS14235). Three replicates shown. The average normalized read coverage for each genome is displayed as the grey line. Maximum read coverage for the region is indicated on the y-axis. Forward transcription (red) and reverse transcription (blue) were plotted separately. (B) Nucleotide alignment of BVU_RS14350 with trimmed Sanger sequencing data generated from DNase-treated cell-free BV01 supernatant. Despite appearing larger in size by gel electrophoresis (Fig. 2B), no large insertions were detected. Locations of primers used to amplify BVU_RS14350 and for Sanger sequencing denoted by underlined sequence.
Figure S2. Representative extracted ion count (EIC) chromatograms for the bile acid species of interest. Related to Figure 5. EIC are shown for (A) the internal control nordeoxycholic acid (NorDCA), (B) the product of bile acid deconjugation, cholic acid (CA), taurocholic acid (TCA), and glycocholic acid (GCA). Times of flight indicated in grey.
Figure S3. Prevalence of prophage insertion adjacent to tspO in human gut samples. Related to Figures 4 and 6. Reads from 246 healthy human gut metagenomes were obtained from the Human Microbiome Project Healthy Human Subjects Study. (A) Reads were first mapped to representative sequences of tspO from B. vulgatus and B. dorei. Percent abundance tspO reads was calculated on a per sample basis as the number of reads mapping to tspO divided by the total number of reads. Histogram shows counts of samples. (B) Reads mapping to tspO were filtered to only include reads antisense to tspO, predicted to point toward the attB based on the known genomic architecture. Mates to those reads were subsequently mapped to either the empty B. vulgatus attB (To host) or to BV01 and its Salyersvirus relatives (To Salyersvirus) (Fig. 6). Percent tspO reads mated was calculated as the read pairs bridging tspO and the sequence of interest, sequence divided by the total number of reads mapping antisense to tspO. Histogram shows counts of samples.
Figure S4. Whole genome tree, nucleotide alignment and proteomic similarity of Salyersviridae phages. Related to Figure 6. (A) Phylogenomic Genome-BLAST Distance Phylogeny implemented with the VCTOR online tool (Meier-Kolthoff and Göker, 2017) using amino acid data from all phage ORFs. For consistency, all phage genomes were annotated with MetaGeneAnnotator (Noguchi et al., 2008) implemented via VirSorter (Roux et al., 2015). Support values above branches are GBDP pseudo-bootstrap values from 100 replications. Genome alignment of all phages made with MAUVE. One locally collinear block (LCB) connects phages B124-14 and B40-8 to the Salyersviridae at the nucleotide level (green). Other LCB connecting lines removed for clarity. (B) Gene similarity matrix for BV01-encoded genes among the Salyersviridae. Genes for non-BV01 predicted phages were predicted with Prodigal (Hyatt et al., 2010). Similarity was computed as amino acid percent identity for protein coding genes and nucleotide percent identity for tRNA genes.
Figure S5. Confirmation of activity of Salyersviridae phages in vitro and in human-associated samples. Related to Figure 6. DNase-treated culture supernatants for predicted Salyersviruses BV02 (A), BV04 (B), and BV05 (C) were sequenced. Assembly resulted in contigs corresponding to the free form of phages BV02 and BV05; BV04 did not yield any contigs corresponding to the putative prophage region, suggesting it is inactivated. Assembled free phage contigs were aligned to their integrated prophage region with Mauve (A-C). Sequence reads were mapped back to their free and integrated forms and represented as coverage curves (A-C). Vertical orange lines indicate the location of att sequences; vertical dashed blue lines indicate the location of contig breaks. (D) PCR amplification with phage-specific primers tests for phage presence in pellet and supernatant fractions for 7 predicted Salyersviruses. Supernatant fractions were treated with DNase, eliminating all contaminating host genomic DNA, as demonstrated by the amplification of a host marker gene (16S rRNA). BV04 is not detectable in supernatant, supporting the conclusion that it is an inactivated prophage. PCR amplicons were visualized by agarose gel electrophoresis alongside GeneRuler Express DNA ladder (16S rRNA); ladder band sizes shown in Kb. (E) Wastewater viromes were collected, and processed in three ways prior to sequencing (see Methods). Resulting reads were trimmed, pooled, and mapped to all Salyersviridae genomes and crAssphage. Only Salyersvirinae genomes shown in alignment to better demonstrate conservation, constructed with Mauve. Read coverages normalized to total number of reads in the metavirome. The
maximum normalized read coverage for BC01 is 4.1, B40-8 and B124-14 is 1.13, and crAssphage is 7.33. (F) Reads from 246 healthy human gut metagenomes were obtained from the Human Microbiome Project Healthy Human Subjects Study. Reads were mapped to all 20 temperate Salyersviridae phages and crAssphage. Percent reads mapping was calculated on a per sample basis as the number of reads mapping to any virus divided by the total number of reads. Histogram shows counts of samples.