

Loyola University Chicago Loyola eCommons

Bioinformatics Faculty Publications

Faculty Publications

6-11-2018

Detecting Viral Genomes in the Female Urinary Microbiome

Andrea Garretto Loyola University Chicago

Krystal Thomas-White Loyola University Chicago

Alan J. Wolfe Loyola University Chicago

Catherine Putonti Loyola University Chicago, cputonti@luc.edu

Follow this and additional works at: https://ecommons.luc.edu/bioinformatics_facpub



Part of the Bioinformatics Commons, and the Biology Commons

Recommended Citation

Garretto, Andrea; Thomas-White, Krystal; Wolfe, Alan J.; and Putonti, Catherine. Detecting Viral Genomes in the Female Urinary Microbiome. Journal of General Virology, 99, : 1141-1146, 2018. Retrieved from Loyola eCommons, Bioinformatics Faculty Publications, http://dx.doi.org/10.1099/jgv.0.001097

This Article is brought to you for free and open access by the Faculty Publications at Loyola eCommons. It has been accepted for inclusion in Bioinformatics Faculty Publications by an authorized administrator of Loyola eCommons. For more information, please contact ecommons@luc.edu.



This work is licensed under a Creative Commons Attribution-Noncommercial-No Derivative Works 3.0 License. © The Authors 2018

1	Detecting Viral Genomes in the Female Urinary Microbiome
2	
3	Andrea Garretto ¹ , Krystal Thomas-White ^{2,*} , Alan J. Wolfe ² , Catherine Putonti ^{1,2,3,4}
4	1 Bioinformatics Program, Loyola University Chicago, Chicago, IL, USA
5	2. Department of Microbiology and Immunology, Loyola University Chicago, Maywood, IL, USA
6	3 Department of Biology, Loyola University Chicago, Chicago, IL, USA
7	4 Department of Computer Science, Loyola University Chicago, Chicago, IL, USA
8	
9	*Current address: 325 Sharon Park Dr, Suite 522, Menlo Park, CA, USA
10	
11	Corresponding Author Details: Catherine Putonti. cputonti@luc.edu 1-773-508-3277
12	Keywords: urinary microbiome; urinary virome; bacteriophage; JC polyomavirus
13	Subject Category: Prokaryotic Viruses and Animal Viruses
14	
15 16 17	The GenBank SRA accession numbers for the 30 metagenomes produced here are as follows: ERR926109 through ERR926123 and ERR926139 through ERR926153, under the BioProject Accession PRJEB8104.
18	

#

Α	bs	tr	a	ct
_	\sim	••	u	·

Viruses are the most abundant component of the human microbiota. Recent evidence has uncovered a rich diversity of viruses within the female bladder, including both bacteriophages and eukaryotic viruses. We conducted whole genome sequencing of the bladder microbiome of 30 women: 10 asymptomatic "healthy" women and 20 women with overactive bladder. These metagenomes include sequences representative of human, bacterial, and viral DNA. This analysis, however, focused specifically on viral sequences. Using the bioinformatic tool virMine, we discovered sequence fragments, as well as complete genomes, of bacteriophages and the eukaryotic virus JC polyomavirus. The method employed here is a critical proof-of-concept: the genomes of viral populations within the low biomass bladder microbiota can be reconstructed through whole genome sequencing of the entire microbial community.

The old paradigm that the bladder is sterile results from the use of standard urine culture-dependent methods that are optimized for *E. coli* (1,2). However, there is definitive evidence that communities of bacteria exist within the bladder (3-6), as well as for associations between these bladder microbiota and urinary symptom levels, treatment response, and UTI risk (7-15). Furthermore, the bladder microbiota of individuals both with and without urinary symptoms includes viral species. The viruses isolated from urine include several viruses that infect eukaryotes (16-22), as well as those that infect bacteria (bacteriophages [phages]) (23-25). Metagenomic sequencing of the urinary virome, which detects eukaryotic viruses and phages in the lytic cycle, revealed an abundance of phages (26,27).

Because the bladder microbiota exist at a substantially lower biomass (1,5,6) than many other human niches (e.g. the gut (28)), sequencing the bladder's virome presents unique technical difficulties. From the gut, the viral biomass can be separated and the extracted DNA can be sequenced directly (29,30). In contrast, previous urine virome metagenomic studies have relied on DNA amplification prior to sequencing to increase DNA concentrations (26). These amplification methods, however, have well documented biases (31). As such, the complete diversity of the virome may not be captured. Alternatively, we hypothesized that the challenges of sequencing the bladder virome could be overcome bioinformatically. Bioinformatic approaches have successfully identified complete viral genomes from

47 bacterial metagenomes (e.g. 32). Moreover, complete viral genomes have been reconstructed from viral 48 metagenomes containing significant quantities of non-viral (bacterial and eukaryotic) DNA (e.g. 33). 49 Thus, we conducted whole genome sequencing of the bladder microbiota and examined the sequence 50 data specifically for viral sequences. This approach has the potential to capture both lytic and lysogenic 51 phage sequences present in the community. 52 In a previously published study (10), urine was collected aseptically via transurethral catheter from 10 53 women without urinary symptoms (control) and 20 women with reported overactive bladder symptoms 54 (OAB) and stored with the DNA preservative AssayAssure (Sierra Molecular) at -80°C. In the current 55 study, 5 ml of each urine sample was thawed and the DNA was extracted, as described previously 56 (10,34). Briefly, the urine was incubated in a lysis solution containing mutanolysin and lysozyme and the 57 DNA extracted from the sample using the DNeasy blood and tissue kit (Qiagen, Valencia, CA), according 58 to the manufacturer's instructions. The Illumina Nextera kit was used for whole genome library 59 preparation with fragment sizes of 200-300 bp. Sequencing was conducted on the Illumina HiSeq 2500 60 platform, producing paired-end 100bp x 2 reads. Human contaminating reads were filtered out by 61 mapping to the Human reference genome (hg19) with bowtie2 (35). Supplementary Table 1 lists the 62 number of raw reads and filtered reads for each patient sample sequenced. Most reads produced 63 represent bacterial and viral species; on average, only 5.3% of the reads mapped to the human 64 reference genome sequence. Raw sequencing data are available from NCBI's SRA database, BioProject 65 Accession # PRJEB8104. The accession numbers for each sample are listed in Supplementary Table 1. 66 Supplemental Figure S1 outlines the analytic process. Each individual metagenome data set was 67 assembled separately. Raw reads were first trimmed for quality using the tool sickle (36) and then 68 assembled by SPAdes (v3.10.1) with the "meta" (metagenomic) option (37). There was only a weak 69 correlation between the number of reads produced for a given sample and the number of contigs 70 assembled from those reads (r=0.23). Next, the virMine (38) tool was used to classify the contigs 71 produced. Briefly, virMine first filtered out contigs less than 1000 bp in length; this length is a user-72 defined parameter and was selected to eliminate partial gene sequences and repetitive elements from 73 downstream analyses. For the remaining contigs, open reading frames were predicted, translated, and 74 compared to virMine's bacterial and viral protein sequence databases (RefSeq protein sequences). 75 These comparisons enabled us to classify each contig as bacterial, viral, or unknown (exhibiting no 76 similarity to bacterial or viral). Genome assembly and virMine statistics are listed in Supplementary

Table 1. The microbiomes were dominated by bacterial contigs (90% on average). The contigs classified as "unknown" were queried via megablast to the NCBI nr/nt database finding that the overwhelming majority were human in origin (results not shown). Thus, here we will focus on the 252 contigs from the 30 metagenomes that were predicted to be viral.

Twenty-seven of the 30 bladder metagenomes examined included contigs predicted to be viral. To further evaluate these contigs, each was queried against the nr/nt database via the NCBI web interface using the megablast algorithm (**Supplementary Table 2**). In comparing the contigs to this database, eight samples were identified as containing sequences of human origin. The virMine software characterized these contigs as viral, as they did not resemble bacterial sequences and had moderate sequence similarity to a sequence in the viral database. The contigs within another seven samples were uniformly short (~1 kbp) and only exhibited sequence similarity to annotated transposases. Transposases, along with integrases, can be encoded by a phage to allow that phage to enter its lysogenic (latent) life cycle by inserting itself into the bacterial genome (the inserted phage genome is now called the "prophage") (39). Thus, while these contigs suggest the presence of lysogenic phages within the bladder microbiota, they do not provide insight into the phage species. The remaining 12 metagenomes, however, had recognizable phage and/or eukaryotic virus sequences.

Two patient samples – OAB045 and OAB052 – contained numerous contigs with homologies to annotated phage genes, including genes annotated as encoding tail proteins, phage tail tape measure proteins, phage DNA packaging proteins, phage portal proteins, terminases, and capsid proteins. Furthermore, these contigs represented phage genome fragments, including several coding regions. For instance, in the OAB052 sample, a 4898 bp contig was identified, containing annotated regions for a phage terminase, phage portal protein, endopeptidase Clp, major capsid protein, phage DNA packaging protein, and two hypothetical proteins. This contig is homologous to a region within the 18.3 kbp putative prophage (determined via PHAST (40)) in the *Gardnerella vaginalis* HMP9231 genome. As such, it is unlikely that the contig identified here represents a complete, intact phage genome. Nevertheless, it may represent a *Gardnerella* prophage, which we previously showed to be prevalent within *Gardnerella* strains of the bladder (41). We next examined the contigs that were classified as bacterial by the virMine tool. Blast queries found significant homology (e-score=0) between the larger contigs within the OAB052 metagenome and *G. vaginalis* genome records in GenBank. Thus, we hypothesize that the larger viral contigs detected within the OAB052 patient sample represent lysogenic phages. While here we have

presented the analysis of just one of these contigs, similar observations were made: viral sequences exhibited homologies to annotated prophages within bacterial species that were also found within the sample's metagenome.

107

108

109

110

111

112

113

114

115

116

117

118

119

120

121

122

123

124

125

126

127

128

129

130

131

132

133

134135

136

Larger phage sequences were identified in three patient samples – OAB010, OAB018, and OAB039. Table 1 lists the contigs identified in each of these samples. While many of these larger phage sequences include novel genic content (i.e. low or no sequence homology to records in GenBank), each exhibited some homology to recognized prophage sequences within bacterial genomes (per PHAST (40)). The most similar phage species are listed in Table 1. Based upon the size of the assembled genome and the presence of "hallmark" viral genes (42), we were able to confidently predict the completeness of several of these assembled sequences. The phage sequences listed in Table 1 were then annotated using the RAST server (43) (Supplementary Table 3). The genome map for the putative complete phage genome sequence within the OAB018 patient sample is shown in Fig. 1 (generated using Geneious, Auckland, NZ). Phage sequences identified here are not necessarily unique to the microbiota of the urinary tract (Supplementary Table 2). For instance, the sequence of contig 28 from the OAB010 sample is 99% identical to a prophage found within a Streptococcus agalactiae strain isolated from a patient's blood sample (44), as well as from a strain isolated from a diseased tilapia (GenBank record CP016501). These larger sequences are informative both of the bioinformatic approach employed here and the samples themselves. First, complete (or near-complete) phage genomes can be reconstructed by sequencing bladder microbiome samples. Second, because we sequenced the bacterial and viral fractions together, it is possible to associate phages and their bacterial host. Last, we found evidence of related phages present in the bladder microbiota of different patients. For instance, the OAB018 and OAB039 patient samples both contain phage sequences similar to the Lactobacillus-infecting phages PLE2 and phi adh. These phages were first detected as prophages within the genomes of the probiotic strains L. casei BL23 (45) and L. gasseri ADH, respectively. Further sequencing of the bladder microbiota is necessary to ascertain if these phage families are common constituents of the bladder virome.

Five patient samples, OAB021, OAB026, OAB032, OAB042, and OAB045, contained recognizable complete genomes for the human polyomavirus JC (JCV). Furthermore, a partial genome sequence, 1023 bp, was retrieved from patient sample OAB025. JCV is a circular double-stranded DNA virus (~5130 bp) and naturally occurs in the urine of healthy individuals. A previous study found that up to 80% of adults excreted JCV in their urine (46). Furthermore, JCV quasispecies have been detected in healthy

individuals (47). JCV, however, was not detected within any of the 10 asymptomatic "healthy" individuals (controls) included in this study. While JCV infection has been associated with progressive multifocal leukoencephalopathy, a fatal neurological disorder (48), JCV within individuals with overactive bladder has yet to be studied. The prevalence of JCV within these five samples varied. Raw reads were mapped to the RefSeq for the species (GenBank Accession: NC_001699) using Bowtie 2 (v. 2.2.6) (35) revealing coverage of the JCV genome ranging from 12x to 726.9x. Coverage correlated with the % reads in the sample corresponding to the JCV genome (r^2 =0.9570). JCV was most abundant in patient samples OAB042 and OAB045, in which 4.4% and 3.2%, respectively, of the total reads generated were classified as JCV.

Sample	Contig #	Length	Coverage	Bacterial Blast Homology	Most Similar Phage
		(kbp)		(sequence ID/ query coverage)	(length)
	28	17.5	16.56	S. agalactiae (99%/ 100%)	phiCT453B (36.7 kbp)
OAB010	31	8.1	11.61	S. agalactiae (95%/ 99%)	phiCT453B (36.7 kbp)
	39	3.4	14.21	S. agalactiae (100%/ 100%)	phiARI0923 (33.5 kbp)
	28	37.1	9.54	L. helveticus (87%/ 71%)	phig1e (42.3 kbp)
040010	49	26.8	10.89	L. helveticus (85%/ 15%)	phig1e (42.3 kbp)
OAB018	66	17.8	7.30	L. allii (72%/ 3%)	PLE2 (35.1 kbp)
	148	7.6	6.96	L. helveticus (76%/ 25%)	phi adh (43.8kbp)
048020	55	13.6	18.08	L. allii (72%/ 4%)	PLE2 (35.1 kbp)
OAB039	79	8.5	23.09	L. gasseri (67%/ 57%)	phi adh (43.8kbp)

Table 1. Putative complete/near-complete phage genomes identified within bladder microbiome samples. Most similar phage sequences were determined using PHAST (40).

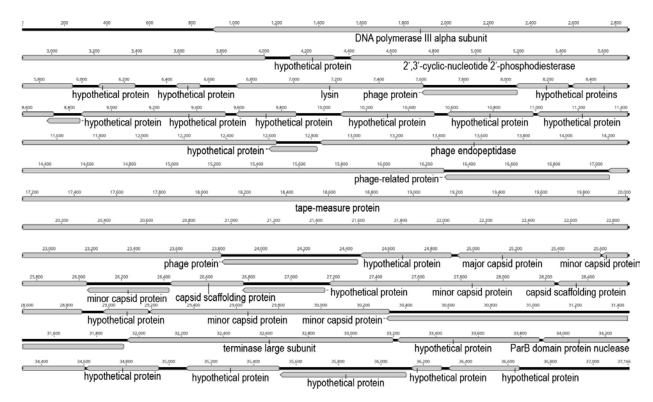


Fig. 1. Genome map for the 37.1 kbp contig 28 from the OAB018 patient sample.

Previous research has identified subtypes of JCV and found that these subtypes can correspond with different human population groups (49). Thus, we next determined the subtypes of the five JCV complete genomes from the bladder microbiome samples by comparing their genomes to 605 publicly available genomes representative of the diversity of the species (**Supplementary Table 2**). The sequences were aligned using MUSCLE through Geneious; the alignments were trimmed, removing the tandem repeats (as their placement at the 5' or 3' end of the genome sequence varied among the genome sequence records), and a phylogenetic tree was inferred using FastTree (50) (**Fig. 2**). Clades were labeled according to their documented genotype, determined from the literature (49) and from GenBank records. Genotype classifications rely on coding sequence variation, most notably the VP1 capsid coding sequence (51). This tree aids in gaining greater insight into the JCV genomes detected within the patient samples. The JCV strains identified in patient samples OAB026 and OAB045 are representative of subtype 1, genotype 1B (exhibiting greatest sequence similarity to isolates from individuals of German heritage (49)). The JCV virus from patient sample OAB042 is also categorized as subtype 1 (genotype 1A) via sequence homology (50). Subtype 1 is relatively common in the United States and Europe (52) and these three patients self-reported as "White/ Non-Hispanic." The JCV strains

identified in patient samples OAB032 and OAB021 are classified as belonging to genotype 3A (prevalent in Africa and southwestern Asia) and 2A (prevalent amongst individuals of Japanese and Native American decent), respectively, based upon their nearest neighbors and placement within the phylogenetic tree (Fig. 2) (49,53). However, the self-reported ethnicities of these patients are incongruent with the ethnicities typically associated with these subtypes; patient OAB032 self-reported as "White/ Hispanic" and patient OAB021 self-reported as "Black/ Non-Hispanic." As the majority of sequencing and genotyping studies of JCV have been largely restricted to individuals with or without neurological diseases, our findings here prompt further investigation of the presence and genotypes of JCV in individuals with and without lower urinary tract symptoms to ascertain if JCV plays any role in urinary tract symptoms or disease.

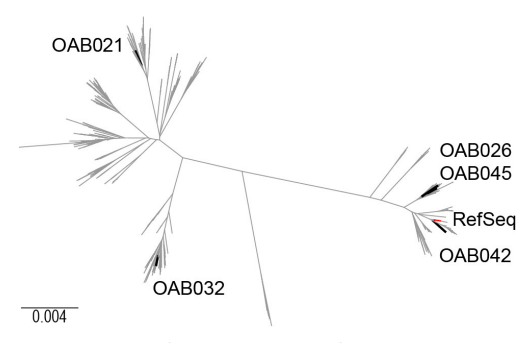


Fig. 2. Phylogenetic tree for 610 complete genomes of JCV, including strains isolated in this study (tree branches shown in black and labeled) and the reference sequence (NC_001699) for the species (shown in red).

Here, we have shown that challenges in isolating viral species from the low biomass bladder microbiome can be circumvented via bioinformatic classification tools; whole genome, as well as partial genome, sequences can be reconstructed from complex samples. While the sheer size of bacterial genomes lends to greater representation in whole genome sequencing data, viral genomes were detected without

- amplification within 27 of the 30 urinary samples examined here. This further supports prior estimates
- of the abundance of viruses within the bladder microbiota (25,26). Moreover, as our results show, our
- strategy can detect both lysogenic and lytic phages, as well as eukaryotic viruses.
- 190 **Abbreviations**:
- 191 OAB=overactive bladder
- 192 JCV=Human polyomavirus JC
- 193 References
- 194 1. Price TK, Dune T, Hilt EE, Thomas-White KJ, Kliethermes S et al. The clinical urine culture: enhanced
- techniques improve detection of clinically relevant microorganisms. *J Clin Microbiol* 2016;54:1216–1222.
- 196 2. **Kass EH.** Pyelonephritis and bacteriuria. A major problem in preventive medicine. *Ann. Intern Med*
- 197 1962;56:46-53.
- 198 3. Wolfe AJ, Toh E, Shibata N, Rong R, Kenton K et al. Evidence of uncultivated bacteria in the adult
- 199 female bladder. *J Clin Microbiol* 2012;50:1376–1383.
- 4. Fouts DE, Pieper R, Szpakowski S, Pohl H, Knoblach S et al. Integrated next-generation sequencing of
- 201 16S rDNA and metaproteomics differentiate the healthy urine microbiome from asymptomatic
- bacteriuria in neuropathic bladder associated with spinal cord injury. *J Transl Med* 2012;10:174.
- 5. Khasriya R, Sathiananthamoorthy S, Ismail S, Kelsey M, Wilson M et al. Spectrum of bacterial
- colonization associated with urothelial cells from patients with chronic lower urinary tract symptoms. J
- 205 *Clin Microbiol* 2013;51:2054–2062.
- 206 6. Hilt EE, McKinley K, Pearce MM, Rosenfeld AB, Zilliox MJ et al. Urine is not sterile: use of enhanced
- 207 urine culture techniques to detect resident bacterial flora in the adult female bladder. J Clin Microbiol
- 208 2014;52:871-876.
- 7. Thomas-White KJ, Kliethermes S, Rickey L, Lukacz ES, Richter HE et al. Evaluation of the urinary
- 210 microbiota of women with uncomplicated stress urinary incontinence. Am J Obstet Gynecol
- 211 2017;216:55.e1-55.e16.
- 8. Brubaker L, Nager CW, Richter HE, Visco A, Nygaard I et al. Urinary bacteria in adult women with
- 213 urgency urinary incontinence. *Int Urogynecology J* 2014;25,1179–1184.
- 9. Pearce MM, Zilliox MJ, Rosenfeld AB, Thomas-White KJ, Richter HE et al. The female urinary
- 215 microbiome in urgency urinary incontinence. *Am J Obstet Gynecol* 2015;213:347.e1-11.
- 10. Pearce MM, Hilt EE, Rosenfeld AB, Zilliox MJ, Thomas-White K et al. The female urinary
- 217 microbiome: a comparison of women with and without urgency urinary incontinence. mBio
- 218 2014;5:e01283-1214.
- 11. Thomas-White KJ, Hilt EE, Fok C, Pearce MM, Mueller ER et al. Incontinence medication response
- relates to the female urinary microbiota. *Int Urogynecology J* 2016;27:723–733 (2016).

- 12. Nienhouse V, Gao X, Dong Q, Nelson DE, Toh E et al. Interplay between bladder microbiota and
- 222 urinary antimicrobial peptides: mechanisms for human urinary tract infection risk and symptom severity.
- 223 PloS One 2014;9:e114185.
- 13. Karstens L, Asquith M, Davin S, Stauffer P, Fair D et al. Does the urinary microbiome play a role in
- 225 urgency urinary incontinence and its severity? Front Cell Infect Microbiol 2016;6:78.
- 14. Thomas-White KJ, Lin H, Gao X, Fok C, Mueller ER et al. Urinary symptoms and associated urinary
- microbes in urogynecologic surgical patients. 2018a. In review.
- 228 15. Thomas-White KJ, Gao X, Lin H, Fok C, Ghanayem K et al. Urinary microbes and post-operative
- urinary tract infection risk in urogynecologic surgical patients. 2018b. In review.
- 230 16. **Iwasawa A, Kumamoto Y, Maruta H, Fukushima M, Tsukamoto T** *et al.* Presence of human
- papillomavirus 6/11 DNA in condyloma acuminatum of the urinary bladder. *Urol Int* 1992;48:235–8.
- 17. Echavarria M, Forman M, Ticehurst J, Dumler JS, Charache P. PCR method for detection of
- adenovirus in urine of healthy and human immunodeficiency virus-infected individuals. J Clin Microbiol
- 234 1998;36:3323-6.
- 18. Karim RZ, Rose BR, Brammah S, Scolyer RA. Condylomata acuminata of the urinary bladder with
- 236 HPV 11. Pathology 2005;37:176-8.
- 19. **Burián Z, Szabó H, Székely G, Gyurkovits K, Pankovics P et al.** Detection and follow-up of torque
- teno midi virus ("small anelloviruses") in nasopharyngeal aspirates and three other human body fluids in
- 239 children. Arch Virol 2011;156:1537-41.
- 240 20. Hirsch HH, Kardas P, Kranz D, Leboeuf C. The human JC polyomavirus (JCPyV): virological
- background and clinical implications. *APMIS* 2013;121:685–727.
- 242 21. Rinaldo CH, Tylden GD, Sharma BN. The human polyomavirus BK (BKPyV): virological background
- and clinical implications. APMIS 2013;121:728–45.
- 22. **Assetta B, Atwood WJ.** The biology of JC polyomavirus. *Biol Chem* 2017;398:839–55.
- 245 23. Brown-Jaque M, Muniesa M, Navarro F. Bacteriophages in clinical samples can interfere with
- 246 microbiological diagnostic tools. *Sci Rep* 2016;6:33000.
- 24. Malki K, Sible E, Cooper A, Garretto A, Bruder K et al. Seven bacteriophages isolated from the
- female urinary microbiota. *Genome Announc* 2016;4:e01003-16.
- 25. Miller-Ensminger T, Garretto A, Brenner J, Thomas-White K, Zambom A et al. Bacteriophages of the
- 250 urinary microbiome. *J Bacteriol* 2018;200:e00738-17.
- 251 26. Santiago-Rodriguez TM, Ly M, Bonilla N, Pride DT. The human urine virome in association with
- urinary tract infections. Front Microbiol 2015;6:14.
- 27. Rani A, Ranjan R, McGee HS, Metwally A, Hajjiri Z et al. A diverse virome in kidney transplant
- patients contains multiple viral subtypes with distinct polymorphisms. Sci Rep 2016;6:33327.

- 255 28. Sender R, Fuchs S, Milo R. Revised estimates for the number of human and bacteria cells in the
- 256 body. *PLoS Biol* 2016;14:e1002533.
- 29. Minot S, Sinha R, Chen J, Li H, Keilbaugh SA et al. The human gut virome: Inter-individual variation
- and dynamic response to diet. Genom Res 2011;21:1616-1625.
- 30. Marique P, Bolduc B, Walk ST, van der Oost J, de Vos WM et al. Healthy human gut phageome. Proc
- 260 Natl Acad Sci USA 2016;113:10400-10405.
- 31. Yilmaz S, Allgaier M, Hugenholtz P. Multiple displacement amplification compromises quantitative
- analysis of metagenomes. *Nat Methods* 2010;7:943-4.
- 32. **Dutilh B, Cassman N, McNair K, Sanchez S, Silva G et al.** A highly abundant bacteriophage
- discovered in the unknown sequences of human faecal metagenomes. *Nature Commun* 2014;5:4498.
- 33. Mokili JL, Dutilh BE, Lim YW, Schneider BS, Taylor T et al. Identification of a novel human
- 266 papillomavirus by metagenomic analysis of samples from patients with febrile respiratory illness. PLoS
- 267 One 2013;8:e58404.
- 34. Yuan S, Cohen DB, Ravel J, Abdo Z, Forney LJ. Evaluation of methods for the extraction and
- purification of DNA from the human microbiome. *PLoS One* 2012;7:e33865.
- 270 35. Langmead B, Salzberg SL. Fast gapped-read alignment with Bowtie 2. Nature Methods 2012;9:357-
- 271 359.
- 36. **Joshi NA, Fass JN.** Sickle: A sliding-window, adaptive, quality-based trimming tool for FastQ files.
- 273 2011. https://github.com/najoshi/sickle.
- 37. Bankevich A, Nurk S, Antipov D, Gurevich AA, Dvorkin M et al. SPAdes: A new genome assembly
- algorithm and its applications to single-cell sequencing. *J Comput Biol* 2012;19:455–477.
- 38. Garretto A, Hatzopoulos T, Kalesinskas L, Putonti C. virMine: Automated detection and annotation
- of complete viral genomes from complex metagenomic samples. 2017.
- 278 https://github.com/thatzopoulos/virMine.
- 39. **Krupovic M, Forterre P.** Single-stranded DNA viruses employ a variety of mechanisms for integration
- into host genomes. *Ann NY Acad Sci* 2015;1341:41–53.
- 40. **Zhou Y, Liang Y, Lynch KH, Dennis JJ, Wishart DS.** PHAST: A fast phage search tool. *Nucl Acids Res*
- 282 2011;39:W347-52.
- 41. Malki K, Shapiro JW, Price TK, Hilt EE, Thomas-White K et al. Genomes of Gardnerella strains reveal
- an abundance of prophages within the bladder microbiome. *PLoS One* 2016;11:e0166757.
- 42. Koonin EV, Senkevich TG, Dolja VV. The ancient Virus World and evolution of cells. Biol Direct
- 286 2006;1:29.
- 43. Aziz RK, Bartels D, Best AA, DeJongh M, Disz T et al. The RAST Server: Rapid annotations using
- subsystems technology. *BMC Genomics* 2008;9:75.

- 289 44. Teatero S, McGeer A, Li A, Gomes J, Seah C et al. Population structure and antimicrobial resistance 290 of invasive serotype IV group B Streptococcus, Toronto, Ontario, Canada. Emerg Infect Dis 2015;21:585-291 91. 292 45. Dieterle ME, Martin JF, Durán R, Nemirovsky SI, Rivas CS et al. Characterization of prophages 293 containing "evolved" Dit/Tal modules in the genome of Lactobacillus casei BL23. Appl Microbiol 294 Biotechnol 2016;100:9201-9215. 295 46. Chang H, Wang M, Tsai RT, Lin HS, Huan JS et al. High incidence of JC viruria in JC-seropositive older 296 individuals. J Neurovirol 2002;8:447-451. 297 47. Van Loy T, Thys K, Tritsmans L, Stuyver LJ. Quasispecies analysis of JC virus DNA present in urine of 298 healthy subjects. PLoS One 2013;8:e70950. 299 48. Khalili K, Gordon J, White MK. The polyomavirus, JCV and its involvement in human disease. Adv Exp 300 Med Biol 2006;577:274-87. 301 49. Shackelton LA, Rambaut A, Pybus OG, Holmes EC. JC virus evolution and its association with human 302 populations. J Virol 2006;80:9928-33. 303 50. Price M, Dehal P, Arkin A. FastTree 2 - approximately maximum-likelihood trees for large 304 alignments. PLoS One 2010;5:e9490. 305 51. Agostini HT, Ryschkewitsch CF, Stoner GL. Genotype profile of human polyomavirus JC excreted in 306 urine of immunocompetent individuals. J Clin Microbiol 1996;34:159-64. 307 52. Reid CE, Li H, Sur G, Carmillo P, Bushnell S et al. Sequencing and analysis of JC virus DNA from 308 natalizumab-treated PML patients. J Infect Dis 2011;204:237-44.
- 309 53. **Saruwatari L, Sugimoto C, Kitamura T, Ohno N, Sakai E** *et al.* Asian domains of four major genotypes
- 310 of JC virus, Af2, B1-b, CY and SC. *Arch Virol* 2002;147:1-10.

312 Funding Information

- 313 This work was supported by the NIH (R01 DK104718 to A.J.W) and NSF (ABI 1149387 and 1661357 to
- 314 C.P.). A.G. is supported by the Carbon Research Fellowship at Loyola University Chicago and the CREU
- 315 research fellowship (CRA).

316

317

311

Acknowledgements

- For prior patient recruitment, we want to acknowledge the Loyola Urinary Education and Research
- 319 Collaborative (LUEREC), specifically Mary Tulke RN and Drs. Linda Brubaker, Elizabeth Mueller, Cynthia
- 320 Brincat, Susanne Taege, and Tanaka Dune and the patients who provided the samples for this study.

321

322

323	Conflicts of Interest
324	The authors declare that there are no conflicts of interest.
325	
326	Ethical Statement
327	N/A
328	
329	
330	SUPPLEMENTARY LEGENDS:
331 332	Supplementary Table 1: GenBank Accession number, genome assembly statistics, and virMine analysis statistics for the 30 metagenomes examined.
333 334	Supplementary Table 2: Results for contigs predicted to be viral queried via BLAST against the nr/nt database via the NCBI web interface.
335 336	Supplementary Table 3: Annotations for putative complete/near-complete phage genomes listed in Table 1.
337	Supplementary Figure 1: Workflow for bioinformatic analysis.
338	