Longitudinal Evaluation of Gut Bacteriomes and Viromes after Fecal Microbiota Transplantation for Eradication of Carbapenem-Resistant Enterobacteriaceae

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ABSTRACT Understanding the role of fecal microbiota transplantation (FMT) in the decolonization of multidrug-resistant organisms (MDRO) is critical. Specifically, little is known about virome changes in MDRO-infected subjects treated with FMT. Using shotgun metagenomic sequencing, we characterized longitudinal dynamics of the gut virome and bacteriome in three recipients who successfully decolonized carbapenem-resistant Enterobacteriaceae (CRE), including Klebsiella spp. and Escherichia coli, after FMT. We observed large shifts of the fecal bacterial microbiota resembling a donor-like community after transfer of a fecal microbiota dominated by the genus Ruminococcus. We found a substantial expansion of Klebsiella phages after FMT with a concordant decrease of Klebsiella spp. and striking increase of Escherichia phages in CRE E. coli carriers after FMT. We also observed the CRE elimination and similar evolution of Klebsiella phage in mice, which may play a role in the collapse of the Klebsiella population after FMT. In summary, our pilot study documented bacteriome and virome alterations after FMT which mediate many of the effects of FMT on the gut microbiome community.

IMPORTANCE Fecal microbiota transplantation (FMT) is an effective treatment for multidrug-resistant organisms; however, introducing a complex mixture of microbes also has unknown consequences for landscape features of gut microbiome. We sought to understand bacteriome and virome alterations in patients undergoing FMT to treat infection with carbapenem-resistant Enterobacteriaceae. This finding indicates that transkingdom interactions between the virome and bacteriome communities may have evolved in part to support efficient FMT for treating CRE.

KEYWORDS CRE, FMT, bacteriome, virome

Drug-resistant enteric bacteria, including carbapenem-resistant Enterobacteriaceae (CRE) and vancomycin-resistant enterococci (VRE), are emerging worldwide (1, 2). Intestinal colonization by multidrug-resistant organisms (MDRO) may predispose at-risk individuals to invasive infection, which is associated with high mortality and can serve as a reservoir for spreading the bacteria to other patients (3). Within-patient transfer of high-risk carbapenemase-encoding plasmids has been reported via intestinal microbiota of colonized patients and may contribute to the overall dissemination of carbapenem resistance in the clinical setting (4), giving rise to an urgent need for effective and safe strategies to prevent and eradicate intestinal MDRO colonization (5).
Fecal microbiota transplantation (FMT) is highly effective in the treatment of recurrent*Clostridioides difficile* infections (CDI) (6) and has recently emerged as a promising therapy for decolonization of intestinal multidrug-resistant microorganisms (7). In four small case series with various study protocols, FMT resulted in 33 to 50% decolonization of CRE (7–10). Previous studies revealed compositional alterations of the intestinal bacteriome and the donor-derived engraftment as assessed by fecal sampling before and after FMT (11). FMT would alter recipient microbial ecology in carriers of carbapenem-resistant *Enterobacteriaceae* (12). However, the fate of native and introduced microbes may not be clearly associated with the success of FMT, and the mechanisms by which species are enriched or cleared after FMT in recipients remain unclear (9). Apart from the bacterial community, accumulating evidence suggest that gut viruses are also associated with FMT treatment outcomes in FMT (13, 14). To date, there are limited data on how FMT affects CRE carriage in relation to the gut microbiome after FMT. Using shotgun metagenomic sequencing, we report for the first time comprehensive longitudinal dynamics of the gut bacteriome and virome in three CRE-positive patients who successfully decolonized CRE following FMT.

**RESULTS**

**Intestinal CRE was eradicated after FMT.** Three CRE-positive patients showed persistently positive isolation of carbapenem-resistant *Klebsiella* species and *Escherichia coli* from two consecutive rectal swabs at least 1 week apart. They received two FMTs and successfully cleared the CRE (Fig. 1A). Recipient 1 (female, 90 years old) underwent two FMTs 5 days apart. Tests for CRE performed at 11, 18, and 35 days after the first FMT were negative. Recipient 1 then developed infection of a foot ulcer and received antibiotic therapy (four courses of amoxicillin-clavulanic acid [Augmentin]) from day 36 to 134 (week 6 to week 19) after the first FMT and tested positive for CRE on days 96 and 134. However, the patient became CRE negative on days 152 and 170 after completion of antibiotic therapy. Recipient 2 (male, 70 years old) and recipient 3 (male, 74 years old) each received two FMTs on consecutive days and tested negative for CRE on days 3 and 33, respectively, after FMT.

**FMT induced significant changes in bacteriome diversity and composition.** We next assessed alteration in the fecal bacteriome in CRE-infected subjects after FMT. Fecal samples were collected serially every 4 weeks up to 24 weeks (recipients 1 and 2) or 12 weeks (recipient 3). Although alpha diversity of the recipients’ fecal bacteriomes fluctuated over time, all recipients exhibited increased fecal bacterial diversity and richness at the last follow-up after FMT compared to their corresponding baseline levels (Fig. 2A; Fig. S1A). Following FMT, the relative abundance of *Klebsiella* spp. (CRE species) detected in all recipients and *E. coli* in recipient 2 (CRE *E. coli* carrier)
substantially decreased (Fig. 2B; Fig. S1B), which was further corroborated by qPCR quantification. Next, we compared species-level dissimilarity (using Bray-Curtis dissimilarity) of stool bacterial community composition in the three CRE subjects before and after FMT with that of their respective donors. Principal-coordinate analysis (PCoA) with Bray-Curtis dissimilarity demonstrated that the gut bacteriome in CRE subjects resembled that of the donor immediately after FMT and remained significantly different from the CRE subjects’ own compositions before FMT (Fig. 2C).

Recipient 1 developed infection of a foot ulcer, received antibiotic therapy (four courses of amoxicillin-clavulanic acid [Augmentin]) from day 36 to 134 (week 6 to week 19) after the first FMT, and tested positive for CRE on days 96 and 134. This may have led to the observation that R1_post5 and R1_post6 were more different from the donor than R1_post1.
To explore transfer of donor-derived microbial taxa to recipients, we applied Bayesian source tracking (15) to compare bacterial composition between FMT recipients and their corresponding donors. We observed distinct microbial engraftment ratios in the three recipients. Engraftment of donor-derived bacteria was more pronounced when FMT was performed on consecutive days (in recipient 2 and recipient 3, mean engraftment ratios were 37.2% and 13.3%, respectively) than when FMTs were performed 5 days apart (in recipient 1, the mean engraftment ratio was 2.8%) (Fig. S1C). These data suggest that consecutive FMT infusions may facilitate better engraftment of donor-derived microorganisms in the recipient.

We found a marked shift in bacterial community composition toward that of the donor, with increased *Firmicutes* and decreased *Actinobacteria*, in recipients 2 weeks after FMT (Fig. 2D and E; Fig. S1D and S2). At the species level, members of the *Clostridiales*, such as *Ruminococcus* sp. strain 5_1_39BFAA, were obviously engrafted in recipients (Fig. 2E), as demonstrated by their predominance in donors’ and recipients’ bacteriomes after FMT. The proportions of *Ruminococcus* increased significantly after FMT ($P < 0.05$, Wilcoxon test) (Fig. S1F). Differential analysis by linear discriminant analysis (LDA) effect size (LEfSe) between pre-FMT and post-FMT samples pinpointed the enrichment of *Ruminococcus* sp. strain 5_1_39BFAA after FMT in CRE recipients (Fig. S1F) (false discovery rate [FDR]-corrected $P$ value $< 0.05$; LDA effect size $= 4.3$). Recipients and donors shared a large number of bacterial species (142 species) in their fecal bacteriomes before FMT. After FMT, recipients shared an additional 27 species with the donors (Fig. S1E). These donor-like assemblages and features of donor-transferred taxa after FMT suggest engraftment of donor microbiomes in recipients and a potential role of bacterium-bacterium competitive exclusion dominated by *Ruminococcus* for CRE decolonization.

**Functional alterations of the bacteriome after FMT.** We next investigated the functionality of the bacteriomes of recipients following treatment with FMT. Total bulk DNA sequencing reads from the metagenomic data set were evaluated against reference databases (MetaCyc) to interrogate the abundance of various metabolic pathways. We found that two prominent pathways in donors, the pentose phosphate pathway (nonoxidative branch) and the UDP-N-acetylglucosamine biosynthesis pathway (Fig. S3A and B), were significantly more abundant than in pre-FMT samples ($P < 0.05$). Notably, the differential presence of these two pathways between CRE carriers and donors was primarily contributed by the bacterial genera *Faecalibacterium* and *Ruminococcus* (Fig. S3C and D), both of which are members of the family *Ruminococcaceae*, implying a bacteriome function-level alteration along with taxonomic-level alteration in CRE. We identified marked differences in overall predicted microbiome functionality between the donors and pre-FMT recipients; however, these differences were not reflected in recipients post-FMT (Fig. 3A). In recipients, there were no significant changes in representative functionality before and after FMT. Though the bacteriome functionality profiles of the two donors were similar, the post-FMT functionalities of the recipients were considerably different from each other (Fig. 3B). For instance, recipient 2 showed greater abundances of tricarboxylic acid cycle (TCA)-related functions and lower abundances of lysine and isoleucine biosyntheses than recipients 1 and 3. The core functionality remained stable with respect to each study subject over the course of post-FMT follow-up. However, the persistently salient discrepancies in microbiome functionality between these recipients suggest a significant interindividual variation among individuals after FMT treatment.

**Characterization of the gut virome in patients after FMT.** To explore the links between gut bacteriome and the virome, we carried out interaction correlation analysis of the relative abundance of gut microbiomes. Tight connections were identified in both donor and post-FMT recipient microbiomes (Fig. 4A and C). However, the correlation network was clearly different from that of donor and post-FMT microbiomes, as only ~35% of correlations were common compared to donors. Nevertheless, correlations between bacteriome and virome were enriched in the two groups, as regards direction (positive or negative), supporting the finding that FMT impacts the transkingdom interactions of gut microbiome components. We next analyzed the bacteriophages in the donors’ and the recipients’ fecal samples. Before FMT, the proportions of *Myoviridae* and *Microviridae* were high in terms of...
Functionality alterations following FMT in donor and recipients. (A) Metagenomic functional predictions for donor, pre-FMT, and post-FMT samples. Mean relative gene pathway abundances for pre- and post-FMT samples were not significantly different (Mann-Whitney test; \( P > 0.05 \)). Gene pathway abundances were calculated using the HUMAnN pipeline and grouped by major functional categories. (B) Abundance distribution of bacteria function pathways in donors and in recipients following FMT.
relative abundance (Fig. 5A). Concomitantly, we found a striking increase in *Klebsiella* phages in all recipients (Fig. S4B), including Kp32virus (T7-like phage), Kp34virus (ΦKMV-like phage), Kp36virus (Siphoviridae phage), and Kp15virus (Myoviridae phages) after FMT (Fig. 5C) by metagenomic data analysis. Among these species, Kp34virus was previously reported as a novel virus belonging to the subfamily *Autographivirinae* and lytic for extended-spectrum-β-lactamase-producing *Klebsiella pneumoniae* strains (16). These findings indicate a potential mechanism through which FMT could reduce *Klebsiella* spp. in the gut via transfer or expansion of bacteriophages. Recipient 2 also carried CRE *E. coli* apart from CRE *Klebsiella* spp. before FMT (Fig. 1). By examining the relative abundance of *Escherichia* phages in all three recipients, we found a marked increase of *Escherichia* phage only in recipient 2 (CRE *E. coli* carrier) after FMT (Fig. 5E), supporting targeted bacteriophage expansion for CRE causative bacteria in FMT.

Using virus-like particle (VLP) sequencing, we found that most *Klebsiella* phages and *Escherichia* phages were present at low abundance or absent in stools of either donors or recipients prior to FMT (Fig. S4C and D). As metagenomic analysis on the VLP enriched virome data set provide information only on the presence of actively replicating phage particles at the time of sampling (17), we harnessed our whole-community metagenomic data set, which contained significant fractions of sequence data, to identify the origin of these expanded phages predating CRE bacteria. We found the presence of donor-derived Kp34virus and *Klebsiella* virus KP27 in recipients’ bulk metagenomes (Fig. 5D). Furthermore, we assembled bulk metagenomic sequence reads into contigs and screened with the gene enrichment-based method VirSorter (18). The viral contigs predicted as category 1 and 2 (intact and incomplete virus, respectively) were mapped back onto randomly subsampled (approximately 6 million reads per sample) sequences. The predicted open reading frames from these viral contigs were mapped to the viral protein database in viral RefSeq protein (v84). We identified 86 *Klebsiella* phage contigs with a mean of 59,422 ± 173,404 counts per sample. Numbers of mapped sequences that could be assigned to *Klebsiella* phage were 7,146, 1.5, and 27,425 (mean value) in donor, pre-FMT, and post-FMT microbiotas of all subsampled reads (Fig. 5F). The marked higher abundance level in donors also implied that the bloom of bacteriophages after FMT were more likely of donor origin.

**FMT decolonizes carbapenem-resistant *Klebsiella pneumoniae* and reconstitutes the microbiota in mice.** The data above indicated that FMT effectively eradicates CRE in recipients; hence, we performed experiments with CRE-challenged mice, focusing on virome and bacteriome alteration after FMT. We tested whether the FMT or fecal virome fraction transplantation alone (FVT; actively replicating phage particles, lytic phage) could clear CRE from the gut of densely colonized mice (Fig. 6A). CRE
FIG 5 Analysis of viral composition of donors and CRE recipients, and alterations of bacteriophages following FMT. (A) Relative abundance of gut virome at the order level of donor and recipient before and after FMT. (B) Alpha diversity (Shannon's diversity) of viromes in stools of donors and recipients. (Continued on next page)
eradication was observed in a parallel in vivo study of mice specifically treated by gavage with carbapenem-resistant \textit{Klebsiella pneumoniae} isolated from CRE patients. Carbapenem-resistant \textit{K. pneumoniae} was introduced into antibiotic-treated mice, and the mice were treated with fecal microbiota/virome fractions from healthy mice twice by oral gavage. All mice were densely colonized with CRE prior to the treatment. We observed three distinct clusters in mice treated by FMT and FVT, indicating significantly distinct microbiota community compositions (Fig. 6B) \((P < 0.05)\). We confirmed the clearance by the culture method (stool suspension plated on chromID CS; bioMérieux, France). By the culture method, we found that both FMT and FVT appear to clear CRE: FMT resulted in clearance on day 3, while FVT resulted in clearance on day 10, whereas controls had persistently positive isolation of CRE by the end of experiment, at least 1 month after challenge.

Using metagenomic sequencing, we first characterized the fecal microbiota taxonomic composition. Taxonomic analysis demonstrated that \textit{Klebsiella} and \textit{Escherichia} were highly represented on day 0, approximately 75% and 20%, respectively (Fig. 6C to E). While the microbiota composition of control mice was \textit{Akkermansia} dominated, FMT/FVT resulted in a highly diverse population on day 5 (Fig. 6D and E). Administration of healthy fecal microbiota to \textit{K. pneumoniae}-challenged mice resulted in engraftment and restoration of normal gut microbial community structure. Importantly, the FMT resulted in progressive reduction in \textit{Klebsiella} sp. levels and achieved clearance on day 3 (Fig. 6D). Conversely, at day 5, the genus \textit{Klebsiella} still represented approximately 4.18% and 0.78% of bacterial taxa in control and fecal virome recipient mice (Fig. 6C and E). We found that FMT-treated mice had bacterial compositions more similar to that of the donor than FVT-treated mice at day 5. Despite these differences, fecal microbial communities after FMT or FVT were similarly diverse, as quantified by the Shannon index (Fig. 6F). Bacterial diversity (Shannon index) and richness after FMT were significantly greater than those in control mice \((P < 0.05)\) (Fig. 6F and G). A heat map displays the trajectory of gut virus derived from mouse stool VLP metagenomes (Fig. 7A). At day 1, \textit{Enterobacteria} phage phiP27, \textit{Klebsiella} phage phiKO2, \textit{Salmonella} virus SJ46, and \textit{Escherichia} virus RCS47 were more prominent. Notably, we also observed higher relative abundances of 10 \textit{Klebsiella} phages in samples from FMT-treated mice than in control samples (Fig. 7B).

DISCUSSION

Emerging evidence suggests that FMT has the potential to reduce colonization of multidrug-resistant organisms (MDRO) in the human gut, but the underlying mechanism remains unknown, although it is likely to be similar to that of eradication of \textit{C. difficile}, i.e., outcompition of MDRO and repopulation with a normal gut microflora (19). Competition between microorganisms is not limited to access for nutritional resources. Production of bacteriocins such as nisin and lactacin 3147, which are antimicrobial peptides with bactericidal or bacteriostatic activity, is a more active strategy against competitors (20, 21). FMT outcome would be a synergistic effect of bacterium-virome interaction and outcompition by bacteria introduced by FMT. In this proof-of-concept study, we observed substantial decreases in the relative abundances of \textit{Klebsiella} spp. accompanied by a marked increase in \textit{Klebsiella} phages in three recipients who successfully cleared CRE after FMT. These data highlight for the first time the intimate interplay of the predator-prey relationship between phage and the bacteria in the human gut during FMT treatment for multidrug-resistant microorganisms. We also report an increase of \textit{Escherichia} phage in a CRE \textit{E. coli} carrier after FMT, which further supports the importance of bacteriophages for clearing specific CRE bacteria.

Unlike targeted microbial therapies and specific bacteriophage therapy, which have a target in the host, FMT is “nonselective” and alters the entire microbiota. Until recently, it
has been unclear which components of donor stool drive FMT efficacy, i.e., whether it is bacteria, viruses, fungi, archaea, proteins, chemical mediators, or the interactions between these entities. Our data provide new and novel evidence that increased level of *Klebsiella* and *Escherichia* phages brought by FMT may play a key role in CRE decolonization. All patients were confirmed to have clearance of CRE following treatment with two FMTs. To date, there are only three reports on one-time FMT resulting in successful CRE decolonization; however, they had lower clearance success rates of 33% (9) (two of six patients) after 1 month and 50% (10) (four of eight patients) after 3 months, and 69.2% of participants who received consecutive days of FMT achieved eradication of CRE after 1 month (22). The successful outcome of CRE eradication in all subjects could be due to the protocol of two consecutive FMTs (10), indicating a need to optimize FMT strategies. Furthermore, we observed a greater donor-derived proportion of bacterial community in recipients given two consecutive-day FMTs than in those given a single FMT. Given the small sample size

FIG 6 FMT decolonizes carbapenem-resistant *Klebsiella pneumoniae* and reconstitutes the microbiota in mice. (A) Experimental scheme for *Klebsiella pneumoniae* challenge and treatment with FMT and with FVT alone. (B) PCoA of gut microbiota composition of mice stool identifies separation between control (gray), FMT (yellow), and FVT (blue) samples. (C to E) Fecal microbiota composition at the genus level of treated mice (5 mice per group). (F and G) Diversity (Shannon index) and richness of mouse gut bacteriomes.
FIG 7 Alteration of mice gut virome and impact of FMT in CRE-challenged mice. (A) Abundance distribution of virus in mouse stool following FMT. (B) The relative abundance of 10 predominant *Klebsiella* phages increased after FMT.
of our study, future large-scale studies on MDRO are warranted to investigate the mechanism of FMT in MDRO decolonization.

In addition, our serial sampling allows us to study the longitudinal dynamics of gut bacterial and viral communities and their interactions after FMT. Engraftment of microbiome after FMT, especially by members of the phylum *Firmicutes* and the family *Ruminococcaceae*, supports a more favorable microbiome profile and MDRO resolution in CRE recipients by FMT. Findings derived from previous studies (23, 24) also revealed that FMT patients had an enrichment of *Ruminococcaceae* upon immunotherapy treatment for recurrent hepatic encephalopathy (rHE) and anti-programmed cell death 1 protein (PD-1). In this study, we found interindividual variability of gut microbiome composition after FMT. Recipient factors such as genetics, immunity status, inflammatory status (25), and microbiota have crucial roles in gut microbiome restoration after FMT. Following FMT, donor bacterial strains cohabit with those of the recipient. The colonization success for a given strain may be greater if another representative of the same species is also present in the recipient before FMT (26).

Bacteriophages are of biomedical importance because of their ability to infect bacteria and transmit genes to their bacterial hosts, resulting in altered microbiota compositions, antibiotic resistance, and novel metabolic capacity in distinct microbial lineages (27, 28). In this study, although the overall virome composition fluctuated in the recipients after FMT, alterations in the transkingdom interactions between the virome and bacteriome communities revealed a role of FMT in the recipients’ gut microbiome. Reset of the bacteria-phage ecology after FMT indicated a “kill-the-winner” dynamic (29), whereby upregulation of *Klebsiella* phage was accompanied by decrease in abundance of *Klebsiella* spp. after FMT. These data highlight the importance of restoration of a balanced relationship between the bacterial microbiome and virome after FMT. As bacteriophages propagate via lytic or lysogenic infection of bacteria (30), bacteriophages play an important role in restricting and eliminating antibiotic-resistant organisms. The increased levels of *Klebsiella* phages and *Escherichia* phages introduced by FMT support a new mechanism of FMT and its potential therapeutic use to target decolonization of MDRO. Phages are ubiquitous in bacterium-rich environments, including the gut, and phages are highly strain specific, which potentially makes the direct modulation of the gut microbiota feasible (31). As such, predation by phages and bacterium-bacteriophage coevolution could contribute to the effective decolonization of CRE *Klebsiella* species and CRE *E. coli* in recipients by FMT.

Given the alarming increase in the frequency of multidrug-resistant bacterial infections globally, phage therapy is gaining increasing interest among medical researchers. However, phages infect bacterial hosts with a very narrow range, and bacteria can rapidly develop resistance to phages, suppressing their effectiveness (32, 33). Development of resistance is less likely when multiple different phages are used simultaneously in a cocktail. Moreover, it is challenging to obtain a set of phages that is effective against all variants of a given causative pathogen. FMT is akin to an overall community containing an “active ingredient” of bacteriophages, and thus, it can be used as a means to overcome the disadvantages of unknown target bacteria as well as the inevitable development of phage resistance. Our data highlight the possibility and importance of viral transfer for causative CRE bacteria species. This pilot study was not able to define the origin of all bacteriophages and the specific host-bacteriophage relationship due to existing challenges in the analysis of viral metagenomes (34). Nonetheless, our data will serve as a reference in FMT practice and research for eradication of other antibiotic-resistant microorganisms.

The limitations of this study are in keeping with the nature of the proof-of-concept study design. It remains an exploratory study with a small sample size, and validation is needed in an independent cohort. For this pilot analysis we included all consecutive cases of subjects who received FMT for CRE, and the first three cases had successful FMT. To date, we do not have any cases of CRE infection with unsuccessful FMT, but we believe that this is an important consideration for future analysis and comparison.
With emerging evidence that the virome is associated with FMT outcome, longitudinal studies are needed to provide further information on mechanisms of the roles of gut microorganisms in the resolution of MDRO.

From our pilot study, we cannot clearly define the exact mechanism of FMT action. In particular, it cannot be determined whether the observed taxonomic shifts in the microbiota promoted the clearance of MDRO or rather reflect the resolution of disease with re-establishment of a health-associated microbiome after transfer of the active agents from the donor. However, our findings support one plausible explanation, that phage blooms induced by FMT may act on the community dynamics of gut microbiota, thereby leading to a resolution of CRE. This is supported by the findings that (i) CRE decolonization and bloom of phage targeting CRE were observed in both human recipients and a mouse model and (ii) annotation of quality-controlled viral contigs revealed that phage might have originally derived from sequenced donor genomes.

In conclusion, this study showed successful decolonization of CRE by FMT. Analysis of multikingdom fecal microbiomes created new research opportunities for understanding the feature of MDRO clearance in CRE carriage patients by FMT. This study is unique, as it demonstrates direct evidence of bacterial repopulation, particularly Ruminococcaceae, and an inverse relationship between bacteriophages and CRE in recipients. The decolonization of CRE species, accompanied by an expansion of Klebsiella phages and Escherichia phages, after FMT supports the idea that phage-directed predation of bacteria may lead to CRE eradication. Our results suggest that the gut virome represents an important yet understudied component of the gut microbiome. Evaluating the virome composition of donors and bacteriophage regulations during FMT may be crucial for enhancing the efficacy of FMT in the treatment of various diseases.

MATERIALS AND METHODS

Clinical study of FMT for eradication of CRE in humans. Patients who were ≥18 years old, had two or more stools or rectal swabs positive for CRE (see the supplemental material) at least 1 week apart, and had not received antimicrobial therapy for at least 48 h prior to infusion of FMT were recruited to a clinical trial (NCT03479710). Study approval was provided by The Joint Chinese University of Hong Kong, New Territories East Cluster Clinical Research Ethics Committee (the Joint CUHK-NTEC CREC; CREC reference no. 2017.553). All relevant ethical regulations were followed. Patients consented to participating in the study and agreed to the findings of the research being published. Patients who had active infection with CRE or vancomycin-resistant Enterococcus (VRE) requiring antimicrobial therapy, pregnancy, active gastrointestinal tract infection, or inflammatory disorders, had undergone recent intra-abdominal surgery, had short gut syndrome, used medications which alter gastrointestinal motility were excluded. CRE colonization was defined as the presence of any Enterobacteriaceae with resistance to any of the carbapenems. CRE colonization was confirmed by specific selective medium (chromID Carba Smart; bioMérieux, France).

In this study, patients received two FMTs using frozen donor stool samples. FMT solution (100 mL; raw stool, 50 g) in 0.9% sterile saline was infused over 2 to 3 min into the distal duodenum or jejunum via esophagogastroduodenoscopy (OGD). Stool samples were collected from patients before and after FMT prospectively. Each recipient received FMT from the same single donor for the two FMTs. Donor 4 was the donor source for recipients 1 and 2, and donor 8 was the donor source for recipient 3. Stools for FMT infusion were obtained from donors recruited to Stool Biobank for the Faculty of Medicine, The Chinese University of Hong Kong. Donors were volunteers from the general population, including spouses or partners, first-degree relatives, other relatives, friends, and others who were known or unknown to the potential patients. Donors were required to fulfill a set of eligibility criteria and passed screening laboratory tests for infectious diseases, including infection with CRE and VRE. A total of four controls were recruited from the community through advertisement and from the endoscopy center at the Prince of Wales Hospital from among subjects who had a normal colonoscopy (stools collected before bowel preparation), and two donors served as healthy controls. Additional stool samples that were CRE positive from two patients who spontaneously recovered were collected for gut microbiome comparison. All samples from patients with CRE, donors, and controls were processed simultaneously.

Microbiome analysis. Microbiome analysis was performed on stool samples from three CRE recipients before and after FMT and on samples collected from donors. Approximately 100 mg fecal sample was preswashed with 1 mL double-distilled water (ddH2O) and pelleted by centrifugation at 13,000 × g for 1 min. The fecal pellet was resuspended in 800 μL Tris-EDTA (TE) buffer (pH 7.5), supplemented with 1.6 μL 2-mercaptoethanol and 500 U lytase (Sigma), and incubated at 37°C for 60 min. The sample was then centrifuged at 13,000 × g for 2 min, and the supernatant was discarded. Fecal DNA was subsequently extracted from the pellet using a Maxwell RSC PureFood GMO and authentication kit (Promega) following the manufacturer’s instructions. Briefly, the fecal pellet was added to 1 mL of CTAB (cetyltrimethylammonium bromide) buffer and vortexed for 30 s, and then the sample was heated at 95°C for 5
min. After that, the samples were vortexed thoroughly with beads at maximum speed for 15 min. Then, 40 μL of protease K and 20 μL of RNase A were added to the samples, and the mixture was incubated at 70°C for 10 min. The supernatant was then obtained by centrifuging at 13,000 x g for 5 min and added to a Maxwell RSC machine for DNA extraction. The total extracted fecal DNA was used for metagenomics sequencing.

**VLP enrichment.** Virus-like particles (VLPs) were enriched by using a protocol described in a previous study (13). Two hundred milligrams of stool sample was added to 400 μL saline-magnesium buffer (0.1 M NaCl, 0.002% gelatin, 0.008 M MgSO₄·H₂O, 0.05 M Tris [pH 7.5]) and vortexed for 10 min. The sample then was centrifuged at 2,000 x g, and a suspension was obtained. To remove the bacterial cells and residual host cells, the suspension was further filtered by one 0.45-mm and two 0.22-mm filters. The cleared suspension was incubated with lysozyme (1 mg/mL at 37°C for 30 min) and chloroform (0.2 x volume at room temperature [RT] for 10 min) in turn to degrade any remaining bacterial and host cell membranes. A DNase cocktail including 1 U Baseline Zero DNase (Epicenter) and 10 U Turbo DNase I (Ambion) was added to the sample, and the mixture was incubated at 65°C for 10 min to eliminate non-virus-protected DNA. VLPs were lysed (4% SDS plus 38 mg/mL protease K at 56°C for 20 min) and treated with CTAB (2.5% CTAB plus 0.5 M NaCl at 65°C for 10 min), and nucelic acid was extracted with phenol-chloroform (pH 8.0) (Invitrogen). The aqueous fraction was washed once with an equal volume of chloroform, purified, and concentrated on a column (DNA Clean & Concentrator 89-5; Zymo Research). VLP DNA was amplified for 2 h using Phi29 polymerase (GenomiPhi V2 kit; GE Healthcare) prior to sequencing. Four independent reactions were performed for each sample and amplified DNA was pooled to reduce.

**Metagenomics sequencing and analysis.** Qualified fecal DNA and VLP DNA were cut into fragments, and the sequencing libraries were prepared through the processes of end repair, adding A tails, purification, and PCR amplification. The fecal and VLP DNA libraries were sequenced on an Illumina NovaSeq 6000 system with a PE150 sequencing strategy by Novogene, Beijing, China, and yielded an average number of 48 ± 5.3 million reads (12G data) and 25 ± 3.3 million reads (6G data) per sample, respectively. Raw sequence reads were filtered and quality trimmed using Trimomatic v0.36 (35). Human reads were filtered out using Kneaddata (https://huttenhower.sph.harvard.edu/kneaddata/; reference database, GRCh38 p12) with the default argument to generate clean reads. Bacterial taxonomic and functional profiling were implemented in HUMANn2 v0.11.1 (36). This work includes taxonomic identification by MetaPhlAn2 using clade-specific marker genes (37), annotation of species pangenomes through Bowtie2 (38) with the ChocoPhlAn database, a translated search of unmapped reads with DIAMOND (39) against the UniRef90 universal protein reference database (40), and pathway connection obtained from the gene list with reference to the MetaCyc database (41). The gene families and pathway abundance files of all samples were joined and normalized according to relative abundances for comparison between samples. ABRicate v.1.0.0 (https://github.com/tseemann/abricate) was used to screen for acquired antimicrobial resistance genes against CARD (42) databases (last updated November 2020). Genomic DNA was extracted from CRE isolates, and paired-end reads (150 bp) were generated on the illumina HiSeq platforms and assembled with SPAdes v3.15.0.

**Viral classification, read assembly, and annotation.** Taxonomic profiles of viruses were first determined from the fecal DNA metagenomic data set, using Kraken2 v2.0.7-beta (43). The complete NCBI viral RefSeq database (accessed 18 January 2021) was downloaded (https://www.ncbi.nlm.nih.gov/refseq). Each query was thereafter classified to a taxon with the highest total k-mer hits matched by pruning the general taxonomic trees affiliated with mapped genomes, as described previously (11). To improve the tentative classification of the virome, we additionally used assembly contigs for virome analysis. The quality-controlled reads were assembled into contigs using MEGAHIT (36) for each individual. To quantify contigs in each sample, the quality-controlled reads were mapped back to the contigs using Bowtie2 (37), and the number of mapped reads was calculated by processing SAM files using custom code. To remove differences in sequencing depth, reads per million total reads (RPM) were calculated for each contig. Assembled contigs from virome libraries with lengths greater than 3,000 bp and predicted as category 1 and 2 with VirSorter (22) were selected to predict open reading frames (ORFs) using Prodigal in “meta” mode (38). To annotate the predicted ORFs, the amino acid sequences of the ORFs were queried by Diamond (39) against the viral RefSeq protein (v84) with an E value of <10⁻²⁰ and a bit score of >50. The viral RefSeq proteins with the closest homologies (E value of <10⁻³⁰ and bit score of >50) were considered for each ORF, analogous to a previously reported method (44).

**Animal experiment.** C57BL/6J male mice were used at 6 to 8 weeks of age and were randomly assigned to experimental and control groups in all experiments, age- and gender-matched mice were used. All mice were kept at a strict 24-h light-dark cycle, with lights on from 6 a.m. to 6 p.m. For antibiotic treatment, mice were given vancomycin (0.125 g)-neomycin (0.25 g)-metronidazole (0.25 g)-ampicillin (0.25 g) (combined in 250 mL water) in their drinking water for 2 weeks as previously described (45). On the day of FMT, a fresh FMT was prepared by harvesting stools from normal healthy mice. The stool pellets were then suspended in 100 μL sterile PBS, and mice were subsequently given a 100-μL suspension by oral gavage. Fecal virome fraction transplantation (FVT) was carried out with a VLP preparation. A stool pellet from the untreated healthy mice were suspended in 300 μL sterile PBS and centrifuged at 2,500 x g for 10 min. Then, bacteria were removed in the VLP-containing supernatant using a 0.45-μm filter, followed by a 0.22-μm filter. A 100-kDa centrifugal filter was used at 3,220 x g for 5 min to capture VLPs in the fecal filtrate and then washed 3 times with PBS under the same conditions. Afterward, VLPs on the filter were suspended in 100 μL PBS, and mice received a 100-μL suspension by oral gavage. Control mice received the same volume of PBS as their FMT/FVT group counterparts. All experimental procedures were approved by the Animal Ethics Committee of the Chinese University of Hong Kong.
Culture-based analyses and molecular identification of carbapenemase. Stool samples (200 mg) were plated, transferred into 1 mL phosphate-buffered saline buffer, and agitated to release the microorganisms. An inoculum volume of 100 μL was directly plated onto chromID Carba agar (bioMérieux, France), which consists of a nutrient base combining different peptones, three chromogenic substrates enabling the detection of activities of specific metabolic enzymes for Escherichia coli, Klebsiella/Enterobacter/Serratia/Citrobacter, and Proteaceae, and a proprietary mixture of antibiotics favoring the selective growth of carbapenemase-producing Enterobacteriaceae. Species identification was further performed using matrix-assisted laser desorption ionization–time-of-flight mass spectrometry (MALDI-TOF MS) (BrukerDaltonics, Bremen, Germany) (Table S1). Carbapenemase genes (blaKPC, blaVIM, blaIMP, blablaNDM and blablaOXA-48-like) were detected by multiplex PCR (46–48). Antibiotic susceptibility test results for CRE isolates are shown in Table S2. The distribution of the genetic determinants of resistance detected by whole-genome sequencing (WGS) in each CRE isolate is shown in Fig. S5.

Quantification of abundance of Klebsiella pneumoniae. For quantification of K. pneumoniae loads in human stools, quantitative PCR amplifications were performed using the K. pneumoniae-specific primers (49) K. pneumoniae-F (5′-ATTGAAAGGTTGCAAACGAT-3′) and K. pneumoniae-R (5′-TTCACTCTGAAATTTCTGTGTC-3′) and the universal 16S rRNA primers 16s-1114_F (5′-CGGCAACGACGC AACC-3′) and 16s-1275_R (5′-CCATTGAGACGTGTGAGCC-3′) (50). The primers were validated by determining qPCR efficiency using isolated bacterial DNA from a reference K. pneumoniae strain. Replicates were averaged, and bacterial load was quantified using the 2−ΔΔCT method per the following formula: 2−ΔΔCT of experimental sample − CT of control bacteria 16S)−CT of control sample−CT of control bacteria 16S). The assay was performed using Power SYBR green PCR master mix (TaKaRa BioMedicals, Japan) with the StepOne real-time PCR system.

Quantification and statistics. The abundance data for bacteria and viruses were imported into R 3.3.5. Diversity and rarefaction calculations were performed using the Phylsoq package. Alpha diversity was analyzed by calculating the Shannon index (a proxy for diversity, taking into account both richness and evenness) and Chao1 (a proxy for community richness). Principal-coordinate analysis (PCoA) based on the Bray-Curtis dissimilarity matrix of the microbial community structure was calculated with the vegan R package. Linear discriminant analysis (LDA) effect size (LEfSe) (37) was performed using the online tool available at http://huttenhower.sph.harvard.edu/galaxy/. LDA identifies taxa based on their contribution to the overall observed differences between groups, that is, taxa that are significantly increased in abundance. Statistical significance was verified through analysis of variance with post hoc multiple-comparison testing between groups or the nonparametric Wilcoxon rank-sum analysis, as reported above. Differences in bacterial and viral abundance were calculated using LEfSe (37). Plots were generated with RStudio (using ggplot2 R package) and edited with Illustrator CC (V21.0.0). Statistical analysis was performed using RStudio.

Data availability. Raw data have been deposited in a BioProject in the NCBI Sequence Read Archive: PRJNA556087. Data are available on reasonable request.

SUPPLEMENTAL MATERIAL
Supplemental material is available online only.

FIG S1, JPG file, 0.4 MB.
FIG S2, JPG file, 0.1 MB.
FIG S3, JPG file, 0.1 MB.
FIG S4, JPG file, 0.2 MB.
FIG S5, JPG file, 0.1 MB.
TABLE S1, DOCX file, 0.01 MB.
TABLE S2, DOCX file, 0.01 MB.

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Q.L. performed the experiment, data analysis and drafted the manuscript. T.Z. provided technical support for data analysis and critical comments on the manuscript. F.Z. and C.P.C. contributed to data analysis and DNA sample preparation. K.Y., W.T., Q.S., and Z.X. contributed to data analysis. W.L. and W.Z. contributed to animal work. L.H.S.L., R.N.S.L., M.L.C., and R.W. contributed to clinical FMT practice, management, and the clinical database. Y.K.Y., P.K.S.C., F.K.L.C., and G.C.L. provided critical comments on the manuscript. S.C.N. designed and supervised the study and wrote the manuscript.

We declare no competing interests.
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