Midbiotics: conjugative plasmids for genetic engineering of natural gut flora

Pilvi Ruotsalainen, Reetta Penttinen, Sari Mattila, and Matti Jalasvuori

University of Jyväskylä, Department of Biological and Environmental Science, Nanoscience Center, Jyväskylä, Finland; Department of Biological Sciences, University of Helsinki, Helsinki, Finland; Department of Genetics, University of Cambridge, Cambridge, UK

ABSTRACT
The possibility to modify gut bacterial flora has become an important goal, and various approaches are used to achieve desirable communities. However, the genetic engineering of existing microbes in the gut, which are already compatible with the rest of the community and host immune system, has not received much attention. Here, we discuss and experimentally evaluate the possibility to use modified and mobilizable CRISPR-Cas9-endcoding plasmid as a tool to induce changes in bacterial communities. This plasmid system (briefly midbiotic) is delivered from bacterial vector into target bacteria via conjugation. Compared to, for example, bacteriophage-based applications, the benefits of conjugative plasmids include their independence of any particular receptor(s) on host bacteria and their relative immunity to bacterial defense mechanisms such as restriction-modification systems due to the synthesis of the complementary strand with host-specific epigenetic modifications. We show that conjugative plasmid in association with a mobilizable antibiotic resistance gene targeting CRISPR-plasmid efficiently causes ESBL-positive transconjugants to lose their resistance, and multiple gene types can be targeted simultaneously by introducing several CRISPR RNA encoding segments into the transferred plasmids. In the rare cases where the midbiotic plasmids failed to resensitize bacteria to antibiotics, the CRISPR spacer(s) and their adjacent repeats or larger regions were found to be lost. Results also revealed potential caveats in the design of conjugative engineering systems as well as workarounds to minimize these risks.

Introduction
The possibility to engineer gut microbiome has become a notable avenue of research. Restoration of microbial balance in the gut can provide a cure to a multitude of complex diseases. Nonetheless, stable installation of foreign beneficial microbes in the gut is problematic. Studies have shown that dietary supplement bacteria (probiotics) disappear from the community soon after their ingestion ceases. This has led many teams to compile bacterial cocktails that would establish a more stable population within the gut. Also, the near-complete replacement of gut flora has been used to revert dysbiosis. This so-called bacterial transplantation is an effective approach to cure especially recurrent diarrhea caused by Clostridium difficile, but could also be used to improve various other conditions. The composition of gut flora is also sensitive to diet, and, for example, increase of fiber can result in notable shifts in the community composition. In some circumstances, however, the possibility to modify the genomes of existing bacteria in the gut could provide an alternative to remodel the system.

So far, the genetic engineering of bacterial communities in situ has mainly focused on bacteriophage-based applications. Conjugative plasmids offer an alternative route with differing engineering qualities. They are circular antagonistic genetic elements that can mediate their own transfer from one bacterium to another. In addition, these self-transmissible plasmids can co-transfer non-conjugative plasmids with appropriate oriT site. The relaxosome of the conjugative plasmid recognizes the similar oriT site in non-conjugative plasmid and mobilizes it through conjugation. The exact conjugation mechanisms vary between plasmids, but they all form a channel between the cells through which the plasmid is usually transported as...
a single-stranded DNA molecule to the recipient bacterium. Plasmids can be readily modified with various molecular biology methods, thus providing a relatively simple platform for carrying out \textit{in situ} genetic engineering of bacterial cells. Given that the diversity of gut microbiota varies even between genetically identical twins, the attempts to colonize maladapted (engineered) bacteria within an already established community can be a challenging if not an impossible task. In this respect, the introduction of an engineered mobile element into the existing community instead of relying on the establishment of an entire bacterium provides a potential workaround for deploying desired functionalities within the system. Given the established concepts of probiotics (health-promoting bacteria) and prebiotics (nutrients that promote the growth of beneficial bacteria), “midbiotics” (plasmid-probiotics in a sense) provide yet an alternative form of biotic substances that can be used to acquire beneficial changes in the gut flora. Naturally, such plasmids have only limited use, albeit, in certain instances, they may be even a preferable choice over probiotics such as when only particular genes need to be removed from the community.

Plasmids are divided into incompatibility groups (Inc) based on their potential to stably coexist in a bacterial cell. In other words, two plasmids that share the same Inc-group cannot be maintained in a single cell indefinitely. Conjugative plasmids also often encode entry-exclusion mechanisms that prevent related plasmids from entering the cell. Due to these natural features, the plasmids used for engineering should be uncommon in the targeted flora. Naturally, determining the existence of certain plasmid types routinely from a heterogenous community is a laborious task. Yet, certain antibiotic resistance-conferring plasmids of Enterobacteriaceae, for instance, are relatively rare in patients. Indeed, in a metastudy, Carattoli reviewed the prevalence of different resistance plasmid families in Extended Spectrum Beta-Lactamase (ESBL) strains. Among the rarest were IncP-type plasmids. Despite this, conjugative IncP-plasmids are well-studied, they have a robust conjugation machinery and a broad host range. As such, they provide an example of potential backbones that could be utilized for engineering purposes.

In bacteriophage-based applications, the genetic material within the phage is replaced, and as it infects a cell after the attachment to a specific receptor on the host cell surface, it delivers the genomic cargo into the bacterial host. Phage-based tools have acquired notable attention and are currently under development toward drugs. The advantage of phages is that they have a narrow host range, and thus, they target specifically only the desired fraction of the bacterial community. Yet, bacteria rapidly become resistant to phages, and phages cannot be easily used to exert activity against even all variants of certain species. In this regard, conjugative plasmids provide qualities that could be useful for alternative and more generally applicable engineering purposes. As noted above, conjugative plasmids are usually delivered as a single DNA strand to the recipient cell. The complementing strand is synthesized in the recipient bacterium and thus it contains all the host-specific modifications in the nucleic acids. This way the host does not recognize the incoming plasmid as foreign genetic material, which, in turn, allows the plasmid to establish itself into a natural community without prior knowledge of the features of bacteria therein. Additionally, unlike phages, conjugative plasmids are not dependent on specific receptors on host cells as plasmids require only cell-cell contact. And once plasmid gets into natural bacteria, it can further disseminate itself into the next host. The transfer rate from one bacterium to another is, of course, slower and less-precise than phage-mediated delivery of DNA. This sets certain boundaries for the utilization of plasmids. Nevertheless, conjugative plasmids can provide a broad host range for introducing genetic material into the gut flora.

The advent of CRISPR-Cas9 editing has provoked numerous studies where specific target sequences within various host organisms are modified, even enabling strain-specific elimination of bacteria from heterologous communities. Introduction of CRISPR-Cas9 editing components into conjugative plasmids provides a potential mean to remove unwanted genes such as those conferring antibiotic resistance from diverse bacterial systems. ESBL carriage refers to nonsymptomatic colonization of the gut by bacteria
which are resistant to a wide range of different beta-lactam antibiotics.\textsuperscript{17,18} This is of major concern, as beta-lactams are the most commonly used class of antimicrobials, owing to their broad spectrum and minimal side effects. They are abundantly administered to treat and prevent bacterial infections during various medical procedures. Over the past few decades, ESBL carriage has become increasingly more common among long-term hospitalized patients as well as in the community.\textsuperscript{19} ESBL carriage serves as a reservoir of resistance genes and significantly increases the risk of clinical infections.\textsuperscript{20-22} As such, we here set to evaluate the possibility to use conjugatively transferred plasmids to induce the loss of ESBL genes (located either in plasmids or in the chromosome) from a bacterial community.

\textbf{Results and discussion}

We constructed a midbiotic system consisting of a conjugative IncP plasmid RP4\textsuperscript{23} and a mobilizable pCas9 plasmid containing \textit{Streptococcus pyogenes}-derived CRISPR/Cas9\textsuperscript{24} that targets conserved sites in two different beta-lactamase genes via plasmid-encoded CRISPR RNA (crRNA). Part of RP4 origin-of-transfer (oriT) site was cloned into pCas9 plasmid in order to make it horizontally transferrable by the RP4-encoded relaxosome complex. Further, 543 bp region, including the target site of the CRISPR/Cas9 system, was deleted from the beta-lactamase gene \textit{blaTEM-2} of RP4 to prevent the system from self-targeting. From now on, the RP4\textsuperscript{blaTEM-2\Delta172-714} plasmid is referred to as delivery plasmid and the modified pCas9 as pCRISPR plasmid, crRNA/multi-crRNA referring to spacer(s) targeting the beta-lactamase gene(s).

A donor bacterium (\textit{Escherichia coli} HMS174) harboring midbiotic plasmids (delivery and pCRISPR-crRNA plasmids) was cocultured together with recipient \textit{E. coli} strain (HB101) carrying a conjugative ESBL-plasmid pEC15 that encodes \textit{blaTEM-52b} target gene.\textsuperscript{25} The transfer of these plasmids to ESBL-positive bacteria and the subsequent coexpression of endonuclease Cas9 and crRNA should induce the loss of resistance by guiding the Cas9 complex to ESBL gene and create a double-stranded nick within the target site (Figure 1a). Nicking linearizes the plasmid and prevents its replication. Indeed, after 24 h, only approximately 1:10 000 transconjugants retained the resistance in comparison to a control treatment lacking the crRNA (Figure 1b). To rule out the possibility that this might result from the unequal conjugation rates between pCRISPR-crRNA and pCRISPR-control plasmid, both were conjugated independently to a recipient HB101 lacking the target plasmid (Figure 2a). Altogether, this suggests that in principle the dispersal of such midbiotics in the bacterial flora would relatively efficiently resensitize the ESBL-harbouring recipients to beta-lactams. Yet, while this approach appears promising in accelerating ESBL loss, there are still potential obstacles to be taken into account when specific genes are targeted with Cas9. These obstacles would be relevant to most \textit{in situ} applications that seek to delete specific functions from the community (and sometimes in applications that attempt to introduce them); hence, we decided to take a closer look at the caveats and the realistic prospects of midbiotic engineering.

In many cases, there can be multiple variants of the genes that encode undesired phenotypes. For example, there is no single guiding crRNA sequence that would direct Cas9 to all possible ESBL variants. However, all classes of beta-lactamase genes share sequences that are usually conserved within the class (Figure 3). Targeting these sites would provide a broad activity against the class regardless of specific knowledge of the variant in any particular case. When various crRNAs are combined into the same plasmid similarly to spacer arrays of natural CRISPR systems, several targets could be abolished with a single pCRISPR plasmid. We tested this by adding two crRNA coding sites separated by a repeat into the pCRISPR plasmid. This pCRISPR-multi-crRNA plasmid was then transferred into two bacterial strains each harboring a different type of an ESBL gene (\textit{blaTEM-52b} and \textit{blaCTX-M-14}). The plasmid exhibited the activity against both ESBL types, leading to a nearly 500-fold decrease in cell density in treated bacteria compared to control, suggesting that combination of crRNA sites could indeed be utilized to achieve broad activity (Figure 1c).
We further studied the individual bacteria that appeared to have avoided the anti-ESBL effect despite having been introduced with the midbiotic system. In other words, some bacteria which had received the pCRISPR-crRNA/multicrRNA plasmid still retained the resistance to beta-lactams (Figure 1b–c). Sequencing of CRISPR spacer locus of these plasmids (8 escape colonies/replicate/experiment) revealed that the observed tolerance to the midbiotic treatment after 24
h was mainly due to loss of the beta-lactamase-targeting spacer(s) and their adjacent repeat (see the graphic illustration of spacer deletions in Figure 1b-c). In some cases, however, we did not succeed to amplify the crRNA coding region at all, suggesting that a larger deletion might have occurred within the region. On the other hand, sometimes the crRNA site was unaltered, indicating potential changes elsewhere, such as mutations in tracrRNA, Cas9 or PAM sequence.

Nevertheless, the emergence of mutants may be difficult to prevent, but in principle several copies of the crRNA regions, for example, could be included in the plasmid, hence allowing it to retain its activity even if one of the sites is lost.

Another potential concern derives from the separation of the midbiotic into two or more plasmids. It is possible that the delivery plasmid mobilizing the pCRISPR plasmid goes ‘rogue’ and spreads alone in the community, thus attenuating the desired effect. We investigated this possibility by cultivating midbiotic bacteria (harboring pCRISPR control plasmid) together with ESBL-positive strain for 72 h during which the culture was refreshed once a day. All of the studied clones (90 colonies) with the mobilizable pCRISPR plasmid also contained the delivery plasmid. In contrast to this, all bacteria harboring the delivery plasmid had lost the pCRISPR plasmid (Figure 2b). This indicates that the mobilizable pCRISPR plasmid is not always delivered together with the conjugative plasmid, thus requiring countermeasures to minimize the probability of such events. There are at least two possibilities to achieve this: either the pCRISPR plasmid and delivery plasmid could be combined into a single plasmid or the toxin–antitoxin system could be separated so that the pCRISPR plasmid carries the gene for antitoxin and the delivery plasmid encodes the toxin. In the latter case, the dispersal of the delivery plasmid alone would lead to cytotoxic response and death of the recipient cell.

Conjugative plasmids are agents in natural microbial communities, albeit not an inherent part of any particular strain or species. In the recent bloom in microbiota research, they have so far been a seldom utilized tool for inducing genetic changes in existing bacterial communities. Plasmids could be used both to introduce desired genes or remove existing ones. Whether they have applications beyond laboratories is yet to be demonstrated, and the possible spread of malevolent traits via horizontal gene transfer may be a deterrent against using plasmids for engineering purposes. Indeed, the obvious risk in introducing a conjugative plasmid into a bacterial community is that the element may pick up an unwanted gene and disperse it further into other hosts. Before introduction into clinical applications, the resistance genes of delivery plasmid should be deleted to prevent dispersal of new resistance genes. However, it must be noted that the communities aimed to be engineered will nevertheless harbor various types of mobile genetic elements, and, thus, if there is notable selection within the population for acquiring a particular gene, it is likely to disperse anyway. In any event, if the plasmid used for midbiotic-like engineering must be removed.
from the community, the plasmid-dependent bacteriophages could provide a way to induce direct selection against the plasmid. However, while in vitro experiments suggest that this would result in plasmid loss, it is yet to be determined whether this occurs also in vivo.

Overall, the fraction of the community that can be engineered with conjugative plasmids is equal to the fraction of the flora that receives them. Studies suggest that plasmid dynamics and persistence in a community is a complicated matter where trophic levels and various characteristics of plasmids, their hosts and the environment play an indispensable role. Without extensive selection for the midbiotic plasmid, it is unlikely to spread to even all possible hosts. Therefore, as in the case of ESBL carriage, the midbiotic system could be considered as a booster which accelerates ESBL curing rather than an outright treatment. Sometimes, however, even a small fraction of engineered bacteria may be enough, such as in the case of making the midbiotics encode externally secreted bacteriocins against unwanted bacterial species. Yet, the overall improved understanding of the survival conditions of plasmids can help us find ways both to get rid of conjugative plasmids and, if necessary, to facilitate their dispersal.

Nevertheless, while caution is necessary, the ability to introduce or remove genes within natural bacterial communities is a real possibility that could be considered as a potential tool for genetic engineering of existing bacterial systems or, for example, modification of gut microbe transplants prior to their implementation.

Materials and methods

Plasmids, bacterial strains and culture conditions

In this study, the so-called midbiotic system consists of the conjugative RP4blaTEM-2Δ172–714 plasmid (delivery plasmid) and mobilizable pCas9 plasmid (pCRISPR plasmid, a gift from Luciano Marraffini, Addgene plasmid # 42876) encoding the S. pyogenes CRISPR/Cas9 system with crRNA(s) targeting conservative sites of different beta-lactamase resistance genes in ESBL plasmids (Table 1). pCas9 was made mobilizable by cloning RP4 oriT site into pCas9 digested with Sall (ThermoScientific; Waltham, Massachusetts, United States) into region spanning 7377–7486 bps. The phosphorylated ESBL-gene-targeting crRNA oligonucleotides (2 µM each) were first annealed together in 50 µl reaction with 1x of T4 ligase buffer (New England Biolabs; Ipswich, Massachusetts, United States) and 0.05 M NaCl by heating first at 95°C for 5 min and then cooling it down gradually (1°C/35 sec) to 20°C. Then, crRNA insert was ligated into Bsal (ThermoScientific) digested pCas9 plasmid by T4 ligase in T4 ligase buffer (New England Biolabs; Ipswich, Massachusetts, United States). In order to prepare the pCRISPR-multi-crRNA plasmid, the multicrRNA insert was multiplied by PCR from a synthetic plasmid (GenScript; Nanjing, China) with primers spacer-multi-crRNA-F and spacer-multi-crRNA-R (Supplementary Table 1). PCR product was purified according to instructions of Qiagen PCR purification kit before being ligated (similarly as above) into the plasmid. The pCRISPR-control plasmid was otherwise similar but lacked the crRNA (Table 1). If not mentioned otherwise, all the PCRs were done according to instructions of Phusion Hot Start II High-Fidelity PCR mastermix (ThermoScientific), except for an extended initial denaturation (from 5 min to 7 min 30 s), using C1000 Thermal Cycler (Bio-Rad Laboratories Inc.; Hercules, California, United States). Both ESBL plasmids, pEC13 and pEC15, in recipient strains, originate from nosocomial isolates, and the conserved sites of their respective beta-lactamase genes (Table 1) were selected as targets for the CRISPR/Cas9 system of pCRISPR plasmids.

All the bacterial cultures were grown at +37°C in Luria Bertani Lennox-broth (LB) and, as necessary, plated on LB-agar (1%) plates. When appropriate, the following antibiotic concentrations were used: rifampicin (50 µg/ml), streptomycin (25 µg/ml), kanamycin (25 µg/ml), chloramphenicol (25 µg/ml) and ampicillin (150 µg/ml). Liquid cultures were shaken at 220 rpm.

Partial deletion of blaTEM-2 in RP4

The part of blaTEM-2 gene (172–714 bp) containing the crRNA target site was deleted from RP4 to prevent the midbiotic system from self-targeting the
Table 1. Bacterial strains and plasmids used in the experiments and the spacer sequences of pCRISPR plasmid. Only the resistance genes relevant to the experiments are mentioned here.

| Strain features | Plasmid | Relevant characteristics | Resistance genes |
|-----------------|---------|--------------------------|-------------------|
| **DONOR**       | E. coli K-12, chromosomal rifampicin-resistance | | aph(3')-Ib, tet, blaTEM-Δ172-714 |
| HMS174          | | | |
| | RP4*blaTEM-Δ172-714 | IncP plasmid | |
| | pCRISPR-crRNA | oriT site of RP4 (50 980-51 793 bp) | A spacer targeting conservative site of blaTEM genes |
| | pCRISPR-multi-crRNA | 3 spacers targeting conservative sites of blaTEM, blaCTX-M, blaSHV genes, respectively |
| | pCRISPR-control | Without crRNA |
| **RECIPIENT**   | E. coli K-12, chromosomal streptomycin resistance | | |
| HB101           | | | |
| | pEC13 | Target of pCRISPR-multi-crRNA | blaCTX-M-14 |
| | pEC15 | Target of pCRISPR-crRNA/multi-crRNA | blaTEM-52b |
| **RECIPIENT**   | E. coli B, chromosomal tetracyclin resistance | | |
| BL21 Gold       | | | |
| | pCRISPR-crRNA | A spacer targeting conservative site of blaTEM genes | cat |

**Sequence of the crRNA (5'→3')**

| crRNA | multi-crRNA | crRNAa | multi-crRNAb | crRNAC |
|-------|-------------|--------|--------------|--------|
|       |             | AAACCTACGTCGACAGAAAAGCATCCTTAG | AAACCTACGTCGACAGAAAAGCATCCTTAGTTTTGAGCTATGCTGTTTTGAATGGTCCCAAAACAAATAGGTCACCAGAACCAGGT TTATA | GAGCTATGCTGTTTTGAATGGTCCCAAAACAACTGAATGAGGCGCTTCCCG |

a Sequence of crRNA of pCRISPR-crRNA
b Sequence of crRNAs of pCRISPR-multi-crRNA
The plasmid was isolated from the DH5α strain.
delivery plasmid. The deletion was first created in the RP4 
blaTEM-2 gene cloned in pET24 plasmid by polymerase chain reaction (PCR) using 0.2 µM 
of primers deletion-F and deletion-R (Supplementary Table 1) with elongation time (5 
min 15 s) adjusted so that the plasmid without the unwanted sequence was amplified with extension 
rate 0.5 kb/min. As the deletion was confirmed with agarose gel electrophoresis, pET24-blaTEM-2Δ172–714 
PCR-product was recombined back to circular plasmid by Red/ET recombination in recombineering-proficient E. coli strain GB08-red 
RifR (Gene Bridges; Heidelberg, Germany) according to the manufacturer’s instructions, with the exception of using 10 ng of DNA for transformation. 
Briefly, the truncated blaTEM-2Δ172–714 gene was amplified by PCR with primers delblaTEM2-F and delblaTEM2-R (Supplementary Table 1). Template of the PCR was removed by DpnI treatment (ThermoScientific), and blaTEM-2Δ172–714 PCR product was purified from the gel according to instructions of Qiagen’s Gel purification kit (Hilden, Germany). RP4 plasmid containing the deletion was obtained by recombining blaTEM-2Δ172–714 PCR product into RP4 in GB08-red 
RifR strain. This RP4 
blaTEM-2Δ172–714 was then conjugated from GB08-red 
RifR to BL21 Gold(pCRISPR-crRNA) with a donor to recipient ratio of 2:1 in 
3 ml and then cultivated at +37°C, 220 rpm, for 16 h. Transconjugant bacteria were selected on LB agar plates with chloramphenicol-kanamycin selection. 
The colonies were picked and transferred into LB medium with same antibiotic selection as above and 
cultivated overnight without shaking. The colonies containing the deletion (RP4 
blaTEM-2Δ172–714) were identified by negative selection by plating on LB agar plates with and without ampicillin selection. This RP4 
blaTEM-2Δ172–714 plasmid was conjugated to HMS174 by incubating donor and recipient in ratio 1:1 in 5 ml cultivation for 2 h at +37°C, 220 rpm. 
Transconjugants were selected by plating on LB agar plates with rifampicin-kanamycin selection.

**Midbiotic conjugation**

The efficiency of the midbiotic plasmids in inducing ESBL loss from the transconjugants was investigated with the following setup. Before the experiments, donors 
HMS174(RP4 
blaTEM-2Δ172–714) (pCRISPR-crRNA) and control HMS174(RP4 
blaTEM-2Δ172–714) (pCRISPR-control) and the recipient strain HB101 (pEC15) were cultivated overnight with antibiotic selection. To mix the equal number of cells for 
conjugation experiments, the cell density (colony forming units; cfu/ml) of all the cultures were determined either by plating or by measuring the optical density at OD950 (Multiskan FC, ThermoScientific; Waltham, Massachusetts, United States). The experiments were performed in two sets with slightly different concentrations of bacteria (see below), and each conjugation setup was replicated six times in total. The recipient strain (~3.0 × 10^5 cfu/ml in the first experiment and 
~7.0 × 10^5 cfu/ml in the second) was mixed with the donor (~1.0 × 10^6 cfu/ml in the first and ~2.0 × 10^6 cfu/ml in the second) and cultivated 24 h in 5 ml LB 
without antibiotics. After the experiment, plating was used to measure the cell density of transconjugants (streptomycin-ampicillin-chloramphenicol), recipients (streptomycin-ampicillin), donors (rifampicin-kanamycin-chloramphenicol in the first experiment and rifampicin-chloramphenicol in the second) and 
of the community (no antibiotics). Donor and recipient strains were distinguished by differing resistance for rifampicin and streptomycin, respectively. The presence of RP4 
blaTEM-2Δ172–714 was controlled by kanamycin and the pCRISPR by chloramphenicol selection. In order to observe the potential for different midbiotic plasmids to disperse separately, RP4 
blaTEM-2Δ172–714 and pCRISPR-control plasmid were cultivated for 72 h during which the culture 
was renewed daily by transferring 50 µl of culture into fresh 5 ml LB medium. After 72 h, the density of bacteria carrying either RP4 
blaTEM-2Δ172–714 or pCRISPR-control plasmid and the total cell density was determined by plating with appropriate antibiotics. From the total of 90 colonies (30 colonies/replicate), we determined whether RP4 
blaTEM-2Δ172–714 or pCRISPR-control plasmid containing colonies also accommodated the other midbiotic plasmid.

The conjugation efficiencies of the pCRISPR-crRNA and pCRISPR-control plasmid were determined to be equal by conjugating the plasmids into HB101 without target ESBL plasmid. The donors 
were mixed with the recipient in ratio 1:1, 1.65:100 in 5 ml LB and cultivated overnight in the absence of antibiotics. The cell density (cfu/ml) of transconjugants with pCRISPR plasmids (streptomycin-chloramphenicol), recipients (streptomycin) and
donors (rifampicin-chloramphenicol) as well as the total cell density (no antibiotics) were determined by plating.

**pCRISPR plasmid with multiple ESBL targets**

To test the activity of multi-crRNA, electroporation was used to transform the pCRISPR-multi-crRNA plasmid to ESBL-plasmid harboring strains. Electroporation was performed according to the protocol in a manual of recombineering-proficient *E. coli* strain GB08-red (Gene Bridges; Heidelberg, Germany). The optimal density was measured with UV-mini-1240 UV-VIS Spectrophotometer (Shimadzu; Kyoto, Japan) using 1.5 ml semimicro cuvettes (Brand; Germany). A $6.5 \times 10^5$ cfu/ml of HB101(pEC13) and $4.0 \times 10^6$ cfu/ml of HB101 (pEC15) strain were used for each plasmid transformation. Every plasmid transformation (pCRISPR-multi-crRNA and pCRISPR-control) was conducted in triplicates by using 20 ng of plasmid DNA. One negative control, transformed with 1 µl of water, per bacterial strain was done. The DNA concentration of plasmids was measured according to the protocol of Qubit™ dsDNA HS Assay Kit (Invitrogen; Carlsbad, California, United States) by using Qubit® 2.0 Fluorometer (Invitrogen; Carlsbad, California, United States). Transformants were plated on LB agar plates without antibiotics and with the combination of chloramphenicol and ampicillin. Negative control was plated with and without chloramphenicol selection. The activity of different crRNA sites was determined by counting the colonies on each plate.

**Target site selection**

The conserved regions of the *blaTEM* and *blaCTX-M* beta-lactamase genes for crRNA targets were determined by aligning sequence samples of these classes (obtained from the ResFinder 3.0 database) separately with MUSCLE algorithm with default settings by Geneious 8.1.9 (Biomatters Ltd; Auckland, New Zealand). The most conserved sites with the appropriate PAM sequence were selected for the crRNA spacer sequences.

**Escape mutants**

The survived escape mutant colonies from the conjugation and transformation were re-isolated by plating them on chloramphenicol-ampicillin. Eight colonies/replicate, except two colonies per control replicate (altogether 32 colonies/experiment), were grown in LB media with chloramphenicol-ampicillin at +37°C without shaking. CRISPR locus of pCRISPR-crRNA, pCRISPR-multi-crRNA and pCRISPR-control plasmid were amplified with PCR using one bacterial colony as a template with primers spacerseqF and spacerseqR (Supplementary Table 1). PCR product was purified from primers and nucleotides with 0.4 U of Exonuclease I (20 U/µl, ThermoScientific; Waltham, Massachusetts, United States) and 0.4 U of FastAP Thermosensitive Alkaline phosphatase (1U/µl, ThermoScientific; Waltham, Massachusetts, United States). These reactions were incubated at +37°C for 20 min and then at +80°C for 15 min in order to inactivate the enzymes. Sequencing-PCR of ExoSAP-treated DNA was performed with BigDye Terminator v3.1 Cycle Sequencing kit (Applied Biosystems; Foster City, California, United States) according to the manufacturer’s protocol. The sequencing reactions were purified using the protocol of BigDye Terminator v3.1 Cycle Sequencing kit except centrifugation was performed with 1109 x g and 100 x g and, before adding formamide, samples were dried at +37°C for 10 min. Sequencing was carried out with 3130x1 Genetic Analyzer (Applied Biosystems/HITACHI; Foster City, California, United States). The basecalling was performed with Sequencing Analysis Software v6.0 (Applied Biosystems; Foster City, California, United States), and the sequences were analyzed for deletions or mutations in CRISPR locus by mapping them against the original sequence by using Geneious 8.1.9 (Biomatters Ltd; Auckland, New Zealand).

**Acknowledgments**

Authors wish to acknowledge funding from the Academy of Finland (grant no. 252411 and no. 297049) and Emil Aaltonen Foundation. Ms Tiia Eklund, Ms Hanna Elomaa, Ms Sanni Erämies, Mr Aapo Mikkola, Mr Valtteri Nurminen
and Ms Juuli Peräkylä are acknowledged for assistance in the laboratory.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Funding

This work was supported by the Academy of Finland [grant no. 252411 and no. 297049] and Emil Aaltonen Foundation.

ORCID

Reetta Penttinen (http://orcid.org/0000-0003-2152-9889)

References

1. Kristensen NB, Bryrup T, Allin KH, Nielsen T, Hansen TH, Pedersen O. Alterations in fecal microbiota composition by probiotic supplementation in healthy adults: A systematic review of randomized controlled trials. Genome Med. 2016;8:52. doi:10.1186/s13073-016-0300-5.
2. Maldonado-Gómez MX, Martínez I, Bottacini F, O’Callaghan A, Ventura M, van Sinderen D, Ziegler B, Vangay P, Knights D, Hutkins RW, et al. Stable engraftment of bifidobacterium longum AH1206 in the gut microbiota of healthy adults: a proof of principle study. BMC Res Notes. 2013;6:190. doi:10.1186/1756-0500-6-190.
3. Petrof E, Glooor G, Vanner S, Weese SJ, Carter D, Daigneault MC, Brown EM, Schroeter K, Allen-Vercoe E. Stool substitute transplant therapy for the eradication of clostridium difficile infection: ‘RePOOPulating’ the gut. Microbiome. 2013;1:3. doi:10.1186/2049-2618-1-3.
4. Mattila E, Uusitalo-Seppala R, Wuorela M, Lehtola L, Nurmi H, Ristikankaari M, Moilanen V, Salminen K, Seppälä M, Mattila PS, et al. Fecal transplantation, through colonoscopy, is effective therapy for recurrent clostridium difficile infection. Gastroenterology. 2012;142:480–486. doi:10.1053/j.gastro.2011.11.037.
5. van Nood E, Vrieze A, Nieuwdorp M, Fuentes S, Zoetendal EG, de Vos WM, Visser CE, Kuijper EJ, Bartelsman JF, Tijsen JG, et al. Duodenal infusion of donor feces for recurrent clostridium difficile infection. N Engl J Med. 2013;368:407–415. doi:10.1056/NEJMoa1205037.
6. Singh R, de Groot PF, Geerlings SE, Hodiamont CJ, Belzer C, Berge IJM, de Vos WM, Bemelman FJ, Nieuwdorp M. Fecal microbiota transplantation against intestinal colonization by extended spectrum beta-lactamase producing enterobacteriaceae: a proof of principle study. BMC Res Notes. 2013;6:190. doi:10.1186/1756-0500-6-190.
7. Colman RJ, Rubin DT. Fecal microbiota transplantation as therapy for inflammatory bowel disease: A systematic review and meta-analysis. Journal of Crohn’s and Colitis. 2013;8:1569–1581. doi:10.1016/j.crohns.2013.08.006.
8. Wu GD, Chen J, Hoffmann C, Bittinger K, Chen YY, Keilbaugh SA, Bewtra M, Knights D, Walters WA, Knight R, et al. Linking long-term dietary patterns with gut microbial enterotypes. Science. 2011;334:105–108. doi:10.1126/science.1208344.
9. Bikard D, Euler CW, Jiang W, Nussenzweig PM, Goldberg GW, Dupontet X, Fischetti VA, Marraffini LA. Exploiting CRISPR-cas nucleases to produce sequence-specific antimicrobials. Nat Biotechnol. 2014;32:1146–1150. doi:10.1038/nbt.3043.
10. Citorik RJ, Mimeo M, Lu TK. Sequence-specific antimicrobials using efficiently delivered RNA-guided nucleases. Nat Biotechnol. 2014;32:1141–1145. doi:10.1038/nbt.3011.
11. Smillie C, Garcillán-Barcia MP, Francia MV, Rocha EPC, de la Cruz F. Mobility of plasmids. Microbiol Mol Biol Rev. 2010;74:434–452. doi:10.1128/MMBR.00020-10.
12. Pansegrau W, Ziegelin G, Lanka E. The origin of conjugative IncP plasmid transfer: interaction with plasmid-encoded products and the nucleotide sequence at the relaxation site. Biochim Biophys Acta. 1988;951:365–374.
13. Turnbaugh PJ, Quince C, Faith JJ, McHardy AC, Yatsunenko T, Niazi F, Afouritit J, Egholm M, Henriissat B, Knight R, et al. Organismal, genetic, and transcriptional variation in the deeply sequenced gut microbiomes of identical twins. Proc Natl Acad Sci USA. 2010;107:7503–7508. doi:10.1073/pnas.1002355107.
14. Carattoli A. Resistance plasmid families in enterobacteriaceae. Antimicrob Agents Chemother. 2009;53:2227–2238. doi:10.1128/AAC.01707-08.
15. Dominguez W, O’Sullivan DJ. Developing an efficient and reproducible conjugation-based gene transfer system for bifidobacteria. Microbiology (Reading, Engl). 2013;159:328–338. doi:10.1099/mic.0.061408-0.
16. Mali P, Yang L, Esvelt KM, Aach J, Guell M, DiCarlo JE, Norville JE, Church GM. RNA-guided human genome engineering via Cas9. Science. 2013;339:823–826. doi:10.1126/science.1232033.
17. Penders J, Stobberingh EE, Savelkoul PHM, Wolfs PFG. The human microbiome as a reservoir of antimicrobial resistance. Front Microbiol. 2013;4. doi:10.3389/fmicb.2013.00087.
18. Woerther P, Burdet C, Chachaty E, Andremont A. Trends in human fecal carriage of extended-spectrum β-lactamases in the community: toward the globalization of CTX-M. Clin Microbiol Rev. 2013;26:744–758. doi:10.1128/CMR.00023-13.
19. Karanika S, Karantanos T, Arvanitis M, Grigoras C, Mylonakis E. Fecal colonization with extended-spectrum beta-lactamase–producing enterobacteriaceae and risk factors among healthy individuals: A systematic review and metaanalysis. Clin Infect Dis. 2016;63:310–318. doi:10.1093/cid/ciw283.

20. de Kraker MEA, Davey PG, Grundmann H. Mortality and hospital stay associated with resistant staphylococcus aureus and escherichia coli bacteremia: estimating the burden of antibiotic resistance in europe. PLoS Med. 2011;8:e1001104. doi:10.1371/journal.pmed.1001104.

21. Giannella M, Trecarichi EM, De Rosa FG, Del Bono V, Bassetti M, Lewis RE, Losito AR, Corcione S, Saffioti C, Bartoletti M, et al. Risk factors for carbapenem-resistant Klebsiella pneumoniae bloodstream infection among rectal carriers: A prospective observational multicentre study. Clin Microbiol Infect. 2014;20:1357–1362. doi:10.1111/cmi.12747.

22. Bar-Yoseph H, Hussein K, Braun E, Paul M. Natural history and decolonization strategies for ESBL/carbapenem-resistant enterobacteriaceae carriage: systematic review and meta-analysis. J Antimicrob Chemother. 2016;71:2729–2739. doi:10.1093/jac/dkw221.

23. Pansegrau W, Lanka E, Barth PT, Figurski DH, Guiney DG, Haas D, Helinski DR, Schwab H, Stanisch VA, Thomas CM. Complete nucleotide sequence of birmingham IncP alpha plasmids compilation and comparative analysis. J Mol Biol. 1994;239:623–663. doi:10.1006/jmbi.1994.1404.

24. Jiang W, Bikard D, Cox D, Zhang F, Marraffini LA. RNA-guided editing of bacterial genomes using CRISPR-cas systems. Nat Biotechnol. 2013;31:233–239. doi:10.1038/nbt.2508.

25. Mattila S, Ruotsalainen P, Ojala V, Tuononen T, Hiltunen T, Jalasvuori M. Conjugative ESBL plasmids differ in their potential to rescue susceptible bacteria via horizontal gene transfer in lethal antibiotic concentrations. J Antibirot. 2017;70:805–808. doi:10.1038/ja.2017.41.

26. Jalasvuori M, Friman V, Nieminen A, Bamford JKH, Buckling A. Bacteriophage selection against a plasmid-encoded sex apparatus leads to the loss of antibiotic-resistance plasmids. Biol Lett. 2011;7:902–905. doi:10.1098/rsbl.2011.0384.

27. Ojala V, Mattila S, Hoikkala V, Bamford J, Hiltunen T, Jalasvuori M. Scoping the effectiveness and evolutionary obstacles in using plasmid-dependent phages to fight antibiotic resistance. Future Microbiol. 2016;11:999–1009. doi:10.2217/fmb-2016-0038.

28. Lopatkin AJ, Meredith HR, Srimani JK, Pfeiffer C, Durrett R, You L. Persistence and reversal of plasmid-mediated antibiotic resistance. Nat Commun. 2017;8:1689. doi:10.1038/s41467-017-01532-1.

29. Cairns J, Koskinen K, Penttinen R, Patinen T, Hartikainen A, Jokela R, Ruusulehto L, Viitamäki S, Mattila S, Hiltunen T, et al. Black queen evolution and trophic interactions determine plasmid survival after the disruption of the conjugation network. mSystems. 2018;3. doi:10.1128/mSystems.00104-18.

30. Bates S, Cashmore AM, Wilkins BM. IncP plasmids are unusually effective in mediating conjugation of escherichia coli and saccharomyces cerevisiae: involvement of the tra2 mating system. J Bacteriol. 1998;180:6538–6543.

31. Lennox ES. Transduction of linked genetic characters of the host by bacteriophage P1. Virology. 1955;1:190–206.

32. Hansson MD, Rzeznicka K, Rosenbäck M, Hansson M, Sirijovski N. PCR-mediated deletion of plasmid DNA. Anal Biochem. 2008;375:373–375. doi:10.1016/j.ab.2007.12.005.

33. Zankari E, Hasman H, Cosentino S, Vestergaard M, Rasmussen S, Lund O, Aarestrup FM, Larsen MV. Identification of acquired antimicrobial resistance genes. J Antimicrob Chemother. 2012;67:2640–2644. doi:10.1093/jac/dks261.