Genetic engineering of a temperate phage-based delivery system for CRISPR/Cas9 antimicrobials against Staphylococcus aureus

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Discovery of clustered, regularly interspaced, short palindromic repeats and the Cas9 RNA-guided nuclease (CRISPR/Cas9) system provides a new opportunity to create programmable gene-specific antimicrobials that are far less likely to drive resistance than conventional antibiotics. However, the practical therapeutic use of CRISPR/Cas9 is still questionable due to current shortcomings in phage-based delivery systems such as inefficient delivery, narrow host range, and potential transfer of virulence genes by generalized transduction. In this study, we demonstrate genetic engineering strategies to overcome these shortcomings by integrating CRISPR/Cas9 system into a temperate phage genome, removing major virulence genes from the host chromosome, and expanding host specificity of the phage by complementing tail fiber protein. This significantly improved the efficacy and safety of CRISPR/Cas9 antimicrobials to therapeutic levels in both in vitro and in vivo assays. The genetic engineering tools and resources established in this study are expected to provide an efficacious and safe CRISPR/Cas9 antimicrobial, broadly applicable to Staphylococcus aureus.

Invasive infections with methicillin resistant Staphylococcus aureus (MRSA) in both community and healthcare settings total approximately 80,000/year and accounted for 11,285 deaths in 2011, resulting in direct healthcare costs of more than $4.5 billion in the United States alone1,2. Moreover, the increasing occurrence of vancomycin-intermediate S. aureus (reduced efficacy of vancomycin) resulted from an accumulation of single nucleotide polymorphisms in the S. aureus chromosome by long-term exposure to vancomycin3–5. The increasing frequency of this problem underlines an urgent need for new antibiotics. However, the numbers of newly developed antibiotics and commercial interest in such drugs are decreasing, due to the high costs in development and rapidly rising resistance6. These impediments have led to an interest in the development of alternative therapeutics such as vaccines, probiotics, and phage therapy that are less likely to drive resistance.

The CRISPR (Clustered, regularly interspaced, short palindromic repeats) and CRISPR associated (Cas) genes serve as a bacterial immune system to resist foreign DNA7,8. The Cas9 present in the Type II CRISPR/Cas system of Streptococcus pyogenes is a RNA-guided endonuclease that introduces double-stranded breaks into target genes9. The specificity of Cas9 is guided by a trans-activating small RNA (tracrRNA) and CRISPR RNA (crRNA) harboring a short spacer sequence recognizing the target gene10,11. Recent studies demonstrated that a plasmid or phagemid harboring a CRISPR/Cas9 system programmed to target an antibiotic resistance gene or a specific pathogen could be delivered by a temperate phage and could successfully control antibiotic resistant Escherichia coli or MRSA with minimal effects on non-targeted bacteria12–16. These studies demonstrated the potential use of CRISPR/Cas9 system as a programmable antimicrobial to selectively control the target bacteria at the DNA level without disturbing the normal microbiome12–16. However, the efficacy of CRISPR/Cas9 antimicrobials is still far from being therapeutic, mainly due to the low efficiency in phage-based delivery systems which limited the
efficacy of CRISPR/Cas9 to reduce bacterial colony forming units (CFU) by only one or two logs in \textit{in vivo} and \textit{in vitro} assays\textsuperscript{12,14}. Furthermore, phage-based delivery systems may deliver not only a plasmid or phagemid harboring CRISPR/Cas system, but also host chromosomal segments by generalized and specialized transduction to target cells\textsuperscript{17}. This is particularly important for phage-based delivery systems using \textit{S. aureus} since many important staphylococcal virulence factors such as superantigens and cytolysins are commonly located in mobile genetic elements (MGEs) and are transferred to other \textit{S. aureus} and \textit{Listeria monocytogenes} by temperate phage-mediated generalized transduction\textsuperscript{18,19}, thereby raising the safety issues\textsuperscript{20–22}.

In this study, we demonstrate a genetic engineering strategy to overcome shortcomings in phage-based delivery systems by integrating the CRISPR/Cas9 system into the genome of temperate phage. The modifications allow for improved efficiency of delivery to target cells, expanded host specificity by complementing the tail fiber protein of the phage, and removal of virulence factor genes from the host strain to prevent contamination of harmful bacterial products in the phage lysates and subsequent spread of virulence genes by generalized transduction.

Results

Development of programmable and integrative CRISPR/Cas9 plasmid vector systems. The fact that bacteriophages can package their own genome more efficiently than host genetic elements, such as plasmids and phagemids, inspired us to develop a programmable and integrative vector system containing CRISPR/Cas9 integrated within the phage genome. This strategy was designed to improve packaging and delivery of the CRISPR system to target cells. To generate a programmable CRISPR/Cas system, synthetic oligos containing a promoter, leader sequence and direct repeat (DR) sequence flanked with the BbsI restriction sites (Supplemental Figure S1) were cloned into the pMK4 shuttle vector, resulting pKS1. Synthetic oligos specific to the target gene (spacer sequence) were cloned into the BbsI site, resulting pKS2. (B) To generate integrative CRISPR/Cas9 system, genes encoding tracr-RNA and Cas9 were amplified from chromosomal DNA of \textit{Streptococcus pyogenes} SF370 using PCR and cloned into the modified pMAD-secY temperature sensitive shuttle vector, resulting pKS3. To program CRISPR/Cas9 system specific to the target gene, the pre-crRNA (the promoter, leader sequence, DR, and spacer sequence) was amplified from pKS2 and cloned into pKS3, resulting pKS4. (C) The programmed CRISPR/Cas9 system was integrated into the non-coding region of \textit{φSaBov} genome by homologous recombinations. (D) The CRISPR/Cas9 system programmed to target \textit{S. aureus} is induced and delivered by \textit{φSaBov}. The CRISPR/Cas9 system (tracrRNA, crRNA, and Cas9) is expressed and scanned the PAM sequence and recognize the target sequence in the chromosomal DNA, leading to chromosomal DNA cleavage and bacterial death.
into pKS3, resulting pKS4 (Fig. 1B). This programmed CRISPR/Cas9 system was integrated into the genome by allelic exchange as described below.

**Integration of CRISPR/Cas9 system into the genome of φSaBov lysogenized in S. aureus strain RF122.** In order to select staphylococcal temperate phages efficiently packaging their own phage genomes, we determined the absolute copy number of phage DNA in the phage lysates using quantitative real time PCR (qRT-PCR) and standard curves generated from serially diluted plasmid templates (Supplemental Figure S3). Since a single transducing phage particle harbors a single copy of phage DNA, the copy number of phage DNA is equal to the number of transducing phage particles. We found that the φSaBov lysogenized in S. aureus strain RF122, a bovine mastitis isolate belonging to CC151 lineage, generated an exceptionally high number of transducing phage particles (10.31 Log copies/ml phage lysate) which was 3.44 and 3.98 Log magnitude higher than that of φ11 and φNM1, respectively (Fig. 2A). Based on these results, we selected the φSaBov for phage-based CRISPR/Cas9 delivery system.

To integrate the S. aureus-specific, programmed CRISPR/Cas9 system from pKS4 into the genome of φSaBov, upstream and downstream gene segments of non-coding regions between SAB1737 and SAB1738 of the φSaBov genome were used as donor templates for allelic exchange in S. aureus strain RF122. The CRISPR/Cas9 system was integrated into the genome by allelic exchange as described below.

**Figure 2.** Characterizing the φSaBov integrated with CRISPR/Cas9 system. (A) Integration of CRISPR/Cas9 system specific to S. aureus was verified by southern blot analysis using a probe specific to the leader sequence. (B) The copy number of phage DNA in the phage lysates was determined using quantitative real time PCR and standard curves presented in Supplemental Figure 3. (C) The number of transducing phage particles in the phage lysates was determined by calculating the plaque forming unit (PFU) using semi-solid agar overlay method. The bar graph indicates the average and SEM combined from triple measurements of three independent experiments (n = 9). Asterisk indicates statistical significance in student t-test, compared to the results from φ11 (p < 0.001).
Δ exchange, resulting in RF122φ SaBov (Fig. 2A). The copy number of recombinant phage genome harboring CRISPR/Cas system into the genome of φ SaBov-Cas9-nuc. Southern blot analysis using a probe specific to the pre-crRNA gene confirmed the integration of CRISPR/Cas system into the genome of φ SaBov (Fig. 2A). The copy number of recombinant phage genome harboring CRISPR/Cas9 system (φ SaBov-Cas9-nuc) was determined by quantitative real time PCR (qRT-PCR) and showed a slight decrease, compared to those of wild type φ SaBov, possibly due to the increase of genome size following integration of CRISPR/Cas9 system, but was still significantly higher than φ11 and φNM1 (Fig. 2B). The transduction efficacy of the φ SaBov-Cas9-nuc was determined by calculating the plaque forming unit (PFU) and showed that the PFU of φ SaBov-Cas9-nuc was higher than that of wild type φ SaBov, presumably due to the killing effect of CRISPR/Cas9 system in φ SaBov-Cas9-nuc which prevents lysogenic conversion events by φ SaBov (Fig. 2C).

As a control, CRISPR/Cas9 system without a spacer sequence was integrated into the genome of φ SaBov in RF122Δnuc. Phages induced from this strain were referred to as φ SaBov-Cas9-null.

The specificity and efficacy of φ SaBov-Cas9-nuc in in vitro assays. To assess the efficacy of killing by φ SaBov-Cas9-nuc, S. aureus strain CTH96, a bovine isolate susceptible to φ SaBov, was treated with various multiplicities of infection (MOIs) of φ SaBov-Cas9-nuc, and viable cells were recovered by plating on BHI agar. We defined the MOI as the number of transducing phage particles per recipient cell. When treated for 6 h, recipient cells were completely killed at MOI of 100 or above, and 5.0% and 0.08% of viable recipient cells were recovered at MOI of 10 and 50, respectively (Fig. 3A). In order to explore the natural occurrence of survival due to the spontaneous mutations in targets, a time-dependent killing was measured. Upon treatment with φ SaBov-Cas9-nuc at MOI of 50, the number of viable recipient cells gradually decreased, and no viable cells were recovered after 8 h treatment and sustained at 24 h, suggesting rare occurrence of surviving mutants (Fig. 3B). By contrast, treatment with φ SaBov-Cas9-null did not show any killing effect at MOI of 50 or less and minor killing effects at MOI of 100 or above, presumably due to the lytic cycle of φ SaBov (Fig. 3A and B).

To assess the nuc gene specific killing effect of φ SaBov-Cas9-nuc, we generated the nuc gene deletion mutant of CTH96 (CTH96Δnuc) lacking the target gene for a spacer sequence in CRISPR/Cas9 system and CTH96 carrying a plasmid expressing green fluorescent protein (CTH96pGFPuv). When treated with φ SaBov-Cas9-nuc at a MOI of 50, viable CTH96pGFPuv gradually decreased and completely lost viability at 8 h of treatment (Fig. 3C). By contrast, the number of viable CTH96Δnuc slightly decreased within 2 h of treatment, presumably due to the lytic cycle of φ SaBov, and then gradually increased thereafter (Fig. 3C). Next, the mixed cultures of CTH96Δnuc and CTH96pGFPuv were treated with φ SaBov-Cas9-nuc or φ SaBov-Cas9-null at MOI of 50 for 8 h, and viable cells recovered on BHI plates were analyzed under the UV lamp. When treated with the φ SaBov-Cas9-nuc, recipient cells expressing GFP were selectively killed (Fig. 3D). By contrast, when treated with the φ SaBov-Cas9-null, both CTH96Δnuc and CTH96pGFPuv were equally recovered (Fig. 3D). The killing effect of φ SaBov-Cas9-nuc against S. intermedius (coagulase positive staphylococci) and S. epidermidis (coagulase negative staphylococci) was also tested but was not observed (data not shown). Combined, these results clearly demonstrated the nuc gene specific killing effect of φ SaBov-Cas9-nuc.

The effect of CRISPR/Cas9 system requires biological activities including transcription and translation of the CRISPR/Cas9 system within the recipient cells which might limit the application of CRISPR/Cas9 anti-microbials under the nutritionally and metabolically-limited conditions. To simulate the nutritionally and metabolically-limited conditions, the empty antibiotic disc was inoculated with S. aureus in PBS (1 × 10⁵ CFU), followed by treatment with the φ SaBov-Cas9-nuc at MOIs of 10, 100, and 500. After 8 h treatment, viable cells were recovered by blotting the disc on to the BHI plate. Recipient cells were gradually decreased at MOIs of 10 and 100, and completely decolonized at MOIs of 500 (Fig. 3E). By contrast, the viability of recipient cells lacking the nuc gene (CTH96Δnuc) was not affected (Fig. 3E). These results suggest the potential use of φ SaBov-Cas9-nuc for sanitizing abiotic objects such as medical devices or surfaces.

We chose the nuc gene as a target for CRISPR/Cas9 system because the nuc gene is uniquely present in all S. aureus strains. Thus CRISPR/Cas9 system only kills S. aureus without affecting microbiomes. To determine whether the efficacy of CRISPR/Cas9 killing effect is affected by the spacer gene, we generated the CRISPR/Cas9 system targeting the essA (φ SaBov-Cas9-essA) which is also highly conserved in all sequenced S. aureus strains. The efficacy of φ SaBov-Cas9-essA was not significantly different from that of φ SaBov-Cas9-nuc (Supplemental Figure S4).

We tested the efficacy of CRISPR/Cas9 system targeting two proximate locations of the nuc, thereby, cleaving two chromosomal locations within the nuc gene which also showed a similar killing effect to CRISPR/Cas9 targeting a single location (data not shown). These results suggest that the efficacy of CRISPR/Cas9 killing effect is mainly determined by the efficiency of CRISPR/Cas9 system delivery to the target organisms.

The efficacy of φ SaBov-Cas9-nuc in in vivo assays. To test the efficacy of φ SaBov-Cas9-nuc in vivo, the backs of C57BL/6 mice were shaved and intradermally inoculated with recipient cells (CTH96pGFP, 1 × 10⁵ CFU). After 6 h of infection, φ SaBov-Cas9-nuc or φ SaBov-Cas9-null was injected into the infected skin at an MOI of 500. Following treatment for 24 h, infected skin regions were excised and homogenized to recover viable cells. The number of viable cells recovered from the skins treated with φ SaBov-Cas9-nuc (0.647 ± 0.128 Log CFU/g of tissue, mean ± SEM) was significantly lower than that treated with φ SaBov-Cas9-null (3.333 ± 0.131 Log CFU/g of tissue) (Fig. 4A). Next, to test the nuc gene-specific killing capacity of φ SaBov-Cas9-nuc in vivo, animals were infected with a mixture of CTH96pGFP and CTH96Δnuc (1:1, each at 5 × 10⁴ CFU) for 6 h, followed by treatment with φ SaBov-Cas9-nuc or φ SaBov-Cas9-null at MOI of 500 for 24 h. The proportion of viable cells expressing GFP in infected skin treated with φ SaBov-Cas9-nuc was 2.8 ± 2.6%, in contrast to those...
Figure 3. The efficacy and specificity of φSaBov-Cas9-nuc in *in vitro* assays. A mid exponential culture of *S. aureus* strain CTH96 (1 × 10^5 CFU) was treated (A) at various MOIs of φSaBov-Cas9-nuc or φSaBov-Cas9-null for 6h or (B) at 50 MOI of φSaBov-Cas9-nuc or φSaBov-Cas9-null up to 24h. Viable cells were recovered on BHI plates. Data point represent the average of triple measurements which were repeated in four independent experiments. (C) A mid exponential culture of *S. aureus* strain CTH96pGFP or CTH96Δnuc (1 × 10^5 CFU) was treated at 50 MOI of φSaBov-Cas9-nuc for 8h. Viable cells were recovered on BHI plates. Data points indicate the average of triple measurements which were repeated in three independent experiments. Asterisk indicates statistical significance in student t-test, compared to the results from φSaBov-Cas9-null (p < 0.001). (D) A mixture of *S. aureus* strain CTH96pGFP and CTH96Δnuc (1:1, each at 5 × 10^4 CFU) was treated with φSaBov-Cas9-nuc at MOI of 50 for 8h. Viable cells were recovered from BHI plates. Pictures showing expression of green fluorescent protein were obtained under UV wave length using Gel Doc system (Bio-Rad). Results shown are a representative picture repeated in three independent experiments. (E) A sterile empty antibiotic disc was inoculated with a suspension of CTH96 or CTH96Δint and treated with the φSaBov-Cas9-nuc at various MOIs for 8h. Viable cells were recovered by blotting discs onto BHI plates. Results shown are a representative picture repeated in three independent experiments.
mice skin were shaved, depilated, decolonized with 70% alcohol, and colonized with CTH96pGFP (2 × 10^5 CFU) followed by treatment with the φSaBov-Cas9-nuc or φSaBov-Cas9-null at MOI of 500. After 24 h, infected skin was excised and homogenized. Viable cells were recovered by plating serially diluted homogenates onto BHI plate. The specificity of φSaBov-Cas9-nuc was evaluated by the proportion of viable cells expressing green fluorescent protein in total viable cells. Data points indicate the average of triple measurements in individual mice (n = 9). Asterisk indicates statistical significance in student t-test, compared to the results from φSaBov-Cas9-null (p < 0.001).

![Figure 4](https://www.nature.com/scientificreports/)  
**Figure 4. The efficacy of φSaBov-Cas9-nuc in vivo murine skin infection.** The backs of C57BL/6 mice were intradermally infected with a suspension of (A) CTH96pGFP (1 × 10^5 CFU) or (B) a mixture of CTH96pGFP and CTH96Δnuc (1:1, each at 5 × 10^4 CFU) for 6 h, followed by treatment with the φSaBov-Cas9-nuc or φSaBov-Cas9-null at MOI of 500. After 24 h, infected skin was excised and homogenized. Viable cells were recovered by plating serially diluted homogenates onto BHI plate. The specificity of φSaBov-Cas9-nuc was evaluated by the proportion of viable cells expressing green fluorescent protein in total viable cells. Data points indicate the average of triple measurements in individual mice (n = 9). Asterisk indicates statistical significance in student t-test, compared to the results from φSaBov-Cas9-null (p < 0.001).

Prevention of toxins contaminations in phage lysates. The strain RF122 harbors 10 superantigens (sec, seg, sei, selm, sein, selo, selu, sell, tst1, seix) and 11 cytolysins (hla, hlb, hlgA, hlgB, hlgC, lukD, lukE, lukG, lukH, lukK, lukF) genes (Supplemental Figure S5) which could be contaminated in the phage lysates generated from the strain RF122. Furthermore, our previous studies demonstrated that induction of φSaBov from strain RF122 generated transducting phage particles harboring genetic segments specifically associated with MGEs such as νSaBov, νSaγ, νSa3, and SaPII some of which contain superantigens (sec, seg, sei, selm, sein, selo, selu, sell, tst) and cytolysins (hla, lukD, lukE) and transferred these virulence genes to other S. aureus strains by generalized transduction23,24. To prevent contamination of these phage lysates and potential spread of toxin genes by the φSaBov-based delivery system, 10 superantigens and 11 cytolysins present in the chromosome of RF122Δnuc φSaBov-Cas9-nuc were removed by allelic exchange using a modified pMAD-secY system, resulting in RF122-19Δnuc φSaBov-Cas9-nuc. The deletions were confirmed by PCR analysis (Supplemental Figure S5). As expected, phage lysates generated from the strain RF122Δnuc showed superantigenicity as indicated by significantly higher count per minute (cpm) as a result of incorporation of [³H]-thymidine into dividing cellular DNA (Fig. 6A). Consistently, incubation with phage lysates generated from the strain RF122Δnuc induced cytotoxicity by cytolysins in phage lysates as demonstrated by intercalation of propidium iodide into cellular DNA (Fig. 6B). By contrast, phage lysates generated from the strain RF122-19Δnuc did not induce any superantigenicity or cytotoxicity (Fig. 6A and B).

Expansion of host specificity of φSaBov. The φSaBov has a narrow host range highly specific to bovine CC151 lineage of S. aureus as shown in phage spot test (Fig. 7A, φSaBov panel), thereby the efficacy of φSaBov-Cas9-nuc to human pandemic clonal lineage of S. aureus (ST1, ST5, ST8 and ST36) was minimal or had no effect (Fig. 7B, black bars). Host specificity of phage is primarily determined by phage-encoded tail fiber proteins interacting with receptors on host cells such as membrane proteins or cell wall carbohydrates25–27. We
investigated the possible expansion of host ranges of φSaBov by complementing the gene encoding the tail fiber protein (Tif) from φ11 (orf50) which has a broad spectrum of host range in several pandemic clones of human S. aureus isolates in the United States (Fig. 7A, φ11 panel). A pMK4 shuttle vector harboring the tail fiber protein gene of φ11 (pTF11) was constructed and transformed into the strain RF122-19ΔnucφSaBov and RF122-19ΔnucφSaBov-Cas9-nuc, resulting RF122-19ΔnucφSaBov-pTF11 and RF122-19ΔnucφSaBov-Cas9-nuc-pTF11, respectively. Phages induced from RF122ΔnucφSaBov-pTF11 (φSaBov-pTF11) showed improved clear zone of lysis in phage spot tests against pandemic human clonal lineages (ST1, ST5, ST8 and ST36) of S. aureus strains (Fig. 7A, φSaBov-pTF11 panel), compared to that of φSaBov. Phage absorption assay showed that the complementation of the Tif from φ11 significantly improved the phage absorption to the pandemic human clonal lineage, albeit not as high as to the CC151 (Fig. 7C). Consistently, phages induced from RF122ΔnucφSaBov-Cas9-nuc-pTF11 significantly improved the killing effect of CRISPR/Cas9 against ST1, ST5, ST8 and ST36 lineages of S. aureus ranging from 1.51 to 3.15 order of magnitude (Fig. 7B, gray bars).

Discussion

Conventional antibiotics, targeting proteins of critical bacterial cellular pathways, are often rendered ineffective due to bacteria either acquiring episomes harboring resistance genes or accruing spontaneous mutations in targets6. The CRISPR/Cas9 antimicrobials have become an attractive alternative due to the advantages of sequence-specific killing without disturbing the microbiome and multiplex features of spacer sequences to simultaneous target multiple genes, thereby preventing development of resistant mutants29. Despite promising results, a therapeutic use of CRISPR/Cas9 antimicrobials is still far from being practical due to the shortcomings in efficiency of delivery and safety aspects of phage-based delivery systems12,14. In this study, we demonstrated a novel genetic engineering strategy to enhance the efficacy and safety of phage-based delivery systems by integrating CRISPR/Cas9 system into the genome of a temperate phage to improve the delivery to target cells, complementing phage tail fiber protein to extend the host spectrum, and removing virulence genes from the host strain to prevent contamination by toxins and spread of virulence genes.

We chose the φSaBov lysogenized in S. aureus strain RF122 as a candidate for phage-based CRISPR/Cas9 delivery system because induction of φSaBov from the strain RF122 generated an exceptionally high number of transducing phage particles harboring the phage genome. Indeed, integration of CRISPR/Cas9 system into the genome of φSaBov significantly enhanced the efficacy of S. aureus specific-killing by φSaBov-Cas9-nuc to nearly complete decolonization in vitro under both nutritionally enriched and limited conditions and more than two orders of magnitude CFU reduction in an in vivo murine skin infection experiments. Interestingly, efficient packaging of the φSaBov genome is highly specific to the RF122 background, and not reproduced in other strains such as RN4220 (Supplemental Figure 6) or MW223. These results suggest the presence of genetic elements uniquely present in the chromosome of RF122 promoting phage DNA excision and replication. Phage DNA excision, replication, and packaging are controlled by complex mechanisms involving multiple factors encoded in the phage genome and host chromosome. Upon induction of phage by SOS signals, phage-encoded rint1 and rint2 activate transcription of phage-encoded integrase (Int), excisionase (Xis), and unknown host encoded factors such as IHF.

Figure 5. The efficacy of φSaBov-cas9-nuc in decolonization of S. aureus from surface of skin. The back skin of mice was shaved, depilated, and decolonized with 70% alcohol, followed by recolonization with CTH96pGFP (2 × 10⁴ CFU). After 6h, φSaBov-Cas9-nuc or φSaBov-Cas9-null at MOI of 500 was topically applied in solution (A) or in mixing with a hydrogel (B). Following treatment for 24h, infected skins were dissected and homogenized to determine the viable cell count. Data points indicate the average of triple measurements in individual mice (n = 8). Asterisk indicates statistical significance in student t-test, compared to the results from φSaBov-Cas9-null (p < 0.001).
...and Fis to initiate site-specific recombination at the attachment site (att site)28,30,31. Genome sequence comparison of RF122, MW2, and RN4220 revealed several unique integrases, transposases, and integrative and conjugative elements associated with MGEs and reminiscent of inactivated phage present in the chromosome of RF12232. We are currently investigating the role of these unique genetic elements present in RF122 on efficient phage DNA excision and packaging events by φSaBov.

Our results showed that the application conditions may greatly affect the efficacy of CRISPR/Cas9 mediated killing effect. When applied in the dry skin condition, the φSaBov-Cas9-nuc was unable to decolonize S. aureus from the skin surface, presumably due to the low water activity suppresses transcriptional and translational activities of S. aureus in which the machinery of CRISPR/Cas9 antimicrobial could not be expressed. By contrast, when mixed with a hydrogel to support water activity, the φSaBov-Cas9-nuc successfully decolonized S. aureus from the infected skin surface. It is also possible that intravenous administration of CRISPR/Cas antimicrobials delivered by phage lysates may evoke immune responses by transducing phage particles or bacterial products remaining in the phage lysates, resulting in antibody production, decreasing efficacy, and potential allergic reactions. Therefore, the most practical application of CRISPR/Cas antimicrobials delivered by phage lysates would be topical application to the condition supporting water activity such as infected tissues or the contaminated surfaces of medical and culinary devices and food products. Recently, the United States Food and Drug Administration approved phage cocktails against Listeria monocytogenes for use in ready to eat food as generally recognized as safe, further encouraging topical applications33.

Phage lysates generated by induction or propagation of temperate phage to the host strain harboring a plasmid or phagemid containing CRISPR/Cas system are mixtures of bacterial components including bacterial DNA, proteins, and cell wall components, as well as transducing phage particles. As demonstrated in Fig. 5, phage lysates generated from the strain RF122Δnuc contained superantigens and cytolysins expressed from the chromosome of RF122. These aspects will clearly raise a regulatory compliance concern on pharmaceutical use of phage lysates containing CRISPR/Cas antimicrobials in Western clinical settings. To alleviate this concern, we generated RF122 containing multiple toxin gene deletions resulting in loss of 10 superantigen and 11 cytolysin genes (RF122-19Δnuc) by using the modified pMAD-secY temperature sensitive shuttle vector system established in this study. Therefore, phage lysates generated from RF122-19Δnuc did not show any harmful effects associated with superantigens and cytolysins. Furthermore, RF122-19Δnuc can be used as a virulence factor-free host strain to propagate φSaBov-Cas9-nuc, without risk of spreading superantigen and cytolysin genes by temperate phage-mediated generalized transduction. One may still argue that the virulence factor-free host strain would not be an ultimate solution because phage lysates generated from RF122-19Δnuc may still contain other uncharacterized virulence factors in host chromosomal segments. Our previous study showed that excision and packaging of host chromosomal segments by φSaBov was highly specific to MGEs within the strain RF122 background, and not in other S. aureus strain backgrounds such as RN4220 and MW223. A recent study demonstrated that the integrase and terminase small subunit (TerS) encoded in Staphylococcus aureus pathogenicity islands (SaPI) induced
sequence-specific excision of host chromosome unlinked to phage DNA which packaged into transducing phage particles by the terminase large subunit (TerL) encoded by the helper phage $\phi$SaBov. Genome sequence analysis of strain RF122 showed a single copy of the terL gene associated with the genome of $\phi$SaBov, and two copies of the terS genes, one with $\phi$SaBov and the other with SaPIbov132. We are currently seeking to remove the redundant integrases and TerS gene to prevent excision of host chromosome mediating generalized transduction by $\phi$SaBov.

Figure 7. Expansion of host specificity of $\phi$SaBov by complementing the tail fiber protein. (A) For phage spot test, 10 μL of phage lysates (of $\phi$11, $\phi$SaBov, or $\phi$SaBov complemented with the tail fiber protein of $\phi$11 ($\phi$SaBov-pTF11) containing $1 \times 10^8$ pfu/ml was inoculated onto the lawn culture of several pandemic human clones of $S$. aureus. (B) To measure the efficacy of CRISPR/Cas9 system delivered by $\phi$SaBov complemented with the tail fiber protein of $\phi$11, a mid-exponential culture of pandemic human $S$. aureus clones (1 × 10^5 CFU) was treated with phage lysates of $\phi$SaBov-Cas9-null, $\phi$SaBov-Cas9-nuc, or $\phi$SaBov-Cas9-nuc pTF11 at MOI of 100 for 8 h. The bar graph indicates the average and SEM of Log CFU of viable cells, combined from triple measurements of three independent experiments (n = 9). (C) The phage absorption was determined by calculating the proportion of unbound phage from the total phage input (2.5 × 10^7 pfu) to several pandemic human clones of $S$. aureus (5 × 10^7 CFU) at the MOI of 0.005. Asterisk indicates statistical significance in student t-test, compared to the results from $\phi$SaBov (p < 0.001).
Phage absorption, an important process that determines the host specificity of phage, is mediated by the phage tail module. The phage tail module is typically composed of tape measure protein, minor tail protein, baseplate protein, cell wall hydrolase, and tail fiber protein. The baseplate protein is linked with the tail fiber protein which can be extended out to search the host receptors such as lipopolysaccharides or the outer membrane porin protein C. This contact triggers conformational changes in the baseplate protein of phages, causing irreversible binding of tail fibers to the outer core of lipopolysaccharides and penetration of inner tail tube to bacterial membrane allowing ejection of phage DNA. A recent study in Pseudomonas aeruginosa Pap1 phage demonstrated that a single nucleotide mutation in phage tail fiber protein resulted in altered host specificity. These results suggest that host specificity of phage could be modulated by altering the phage tail module. The tail fiber protein of φ11 (orf50) contains 370 amino acid residues in which the N-terminal part of amino acid residues from 1 to 120 are highly similar to the putative tail fiber protein of φSaBov (orf56) with 74% identity (85/120) and 85% positivity (102/120). Recent studies demonstrated that the baseplate protein of φ11 (Gp45) plays an essential role in phage absorption to S. aureus by mediating interaction with the N-acetyl-glucosamine (GlcNAc) in the peptidoglycan. Interestingly, the baseplate protein of φSaBov (orf51) showed high similarity with that of φ11 (Gp45) with 92% identity (587/636) and 96% positivity (615/635). These suggest a possibility that the baseplate protein is linked with the N-terminal part of tail fiber protein and the C-terminal part of tail fiber protein interact with the specific receptor in the host. This could explain the partial improvement of host specificity of φSaBov by complementing the plasmid harboring the gene encoding tail fiber protein of φ11 (pTFφ11) because the high amino acid sequence homology in the baseplate proteins and the N-terminal part of tail fibers proteins between φSaBov and φ11 allows the interchangeable linkage of tail fiber protein from both φSaBov and pTFφ11 to the baseplate protein of φSaBov, but causes competitions which results in partial improvement of host specificity. These results suggest that elucidating the interaction between the baseplate protein and tail fiber protein may lead to generate a chimeric tail fiber protein that bridges the C-terminal receptor recognition domain of tail fiber protein to the baseplate protein of phage for developing recombinant phage delivery system for CRISPR/Cas9.

Currently, we are pursuing either complete replacement of the tif gene within the φSaBov genome with that of a broad host range phages such as φ11, φ13, and φNM1 by allelic exchange or integration of CRISPR/Cas9 system into the genome of broad host range phage.

A phage therapy carrying CRISPR/Cas antimicrobials undoubtedly has great potential for alternative therapeutics, supplemental to conventional antibiotics, and prophylactic measurement against increasing antibiotic resistant pathogens. The genetic engineering strategy on both phage and host genome established in this study will be useful to create an efficacious and safe CRISPR/Cas9 antimicrobials platform broadly applicable to MRSA and other important pathogens.

**Methods**

**Bacterial strains and growth conditions.** All strains and plasmids used in this study are listed in Supplemental Table 1. Staphylococcus aureus strains were cultured in tryptic soy broth (TSB) or agar (TSA) plates (Difco) supplemented with chloramphenicol (10 μg/mL, Sigma-Aldrich) as necessary. Escherichia coli were grown in Luria-Bertani (LB) broth and agar plates supplemented with ampicillin (100 μg/mL, Sigma-Aldrich) as necessary.

**Plasmid construction.** All oligos used in this study are listed in Supplemental Table 2. Synthetic oligos (CRISPR_F/CRISPR_R) containing promoter, pre-crRNA, and direct repeats flanked with BbsI sites (CRISPR array) were annealed, ligated into pMK4 digested with BamHI and EcoRI, resulting pKS1. Synthetic oligos (spacernuc/spacenurc) containing a spacer sequence specific to the nuc gene followed by protospacer-adjacent motif (NGG) were annealed and ligated into pKS1 digested with BbsI, resulting pKS2. The tracrRNA and the cas9 genes were amplified from the genomic DNA of Streptococcus pyogenes SF370 (ATCC) using oligos (tracrRNA/cas9r), followed by digestion with AflII and EagI, and ligation into corresponding sites in pMK4 digested with BbsI. To program tracrRNA and the cas9 gene specific to S. aureus, CRISPR array specific to the nuc gene was amplified from pKS2 using oligos (leaderf/drr), digested with EagI and SbfI, and cloned into corresponding sites in pKS3, resulting pKS4.

**Allelic exchange construct.** Integration of CRISPR-Cas9 system into the genome of φSaBov and marker-less deletion mutants of the nuc gene and 21 virulence genes were generated by allelic exchange using modified pMAD-secY temperature sensitive shuttle vector system established in this study by introducing a new multi-cloning site, a GFPuv reporter gene, a chloramphenicol resistant gene (cat), and an anti-sense secY gene controlled by a tetracycline inducible promoter in the pMAD system to improve screening process of allelic exchange (Supplemental Figure S1). Briefly, upstream and downstream fragments of target gene were amplified and cloned in modified pMAD-secY system in E. coli, followed by electroporation into S. aureus strains. The first homologous recombination was induced by culturing at 43 °C (non-permissive temperature for the replication of pMAD-secY), followed by culturing at 37 °C to promote the second recombination, resulting in allelic exchange.

The mutant candidates were screened by growth in TSA plate supplemented with anhydrous tetracycline (0.5 μg/mL), loss of GFP expression, and no growth in TSA plate supplemented with chloramphenicol, indicating the second recombination.

**Phage lysates.** Phages were induced from the mid-exponential culture of strains by adding mitomycin C (1 μg/mL, Sigma-Aldrich) which induced clear lysis typically in 3 hours incubation at 30 °C with 80 rpm. The lysates were sterilized with syringe filers (0.22 μm, Nalgene). Phage lysates were generated by propagating phage to the mid-exponential culture of the same strains from which phages were initially induced, followed by filter sterilization of lysates. The number of transducing phage particles (TP) was determined by calculating the
plaque-forming unit using soft agar (0.5%) overlay method or quantitative real time PCR. Briefly, phage lysates were treated with excessive Dnase I (Sigma-Aldrich) to remove chromosomal DNA contamination, followed by DNA extraction from phage particles using DNeasy kit (Qiagen) as described previously23,24. Quantitative real time PCR reaction was performed using SYBR green I master mix (Applied Biosystems), primer sets specific to phages, and a serial dilution of phage DNA templates. The absolute copy number of phage DNA was calculated by interpolation of the threshold cycle from phage DNA template to the standard curves generated from cloned plasmid templates.

**In vitro efficacy tests.** The phage stock generated from the mitomycin C induction was propagated to the RF122-19ΔnucoSaBov-Cas9-nuc strain for 5 times to remove the mitomycin C contamination in the phage lysates. The mid exponential culture of recipient strains was harvested by centrifugation and adjusted to 1 × 10^8 CFU/mL in PBS. A test tube killing assay was performed in 1 mL of reaction mixtures consisting of 100 μL of recipient cell suspension, 20 μL of serially diluted phage lysates, and 880 μL TSB, and incubated at 37°C. The number of viable cells at each time point was determined by serial dilution and plating onto TSA plates. For *in vitro* killing under nutritionally limited conditions, an empty antibiotic disc was placed in sterile petri dish and inoculated with 100 μL of recipient cell suspension in PBS (1 × 10^8 CFU), followed by 20 μL of serially diluted phage lysates. After an 8 hr incubation at 37°C, the viable cells were recovered by blotting the disk onto TSA plates.

**In vivo efficacy tests.** All animal experiments were performed in compliance with a protocol reviewed and approved by the Institutional Animal Care and Use Committee at the Mississippi State University (14–040). The back of C57BL/6 mice (6 to 8 week old, female, Harlan laboratory) were shaved with electric razor, depilated with Nair cream, and decontaminated with 70% ethanol swab. For intradermal infection, 100 μL of phage lysates prepared from RF122Δnuc strain for 5 times to remove the mitomycin C contamination in the phage lysates. After an 8 hr incubation at 37°C, the viable cells were recovered by blotting the disk onto TSA plates.

**Phage spot and absorption assay.** For phage spot assay, recipient strains grown to the mid-log phase in BHI were harvested, adjusted to 1 × 10^8 CFU/mL, mixed with soft agar (0.5% w/v), and overlaid on the BHI plate. Ten microliters of phage lysates (1 × 10^8 pfu/ml) was dropped onto the lawn culture.

For phage absorption assay, 100 μL of the mid-exponential phase recipient strain (S. aureus 5 × 10^8 CFU/ml) was mixed with 100 μL of phage lysates (2.5 × 10^8 PFU/ml) in phage buffer (50 mM Tris-HCl pH 7.5, 100 mM NaCl, 10 mM CaCl2) and incubated for 15 min at 37°C. The phages bound to the bacterial cells was removed by centrifugation at 13,000 rpm for 15 min. The PFU of the supernatant containing the unbounded phage was determined by the soft agar (0.5%) overlay method. The phage absorption (% input) was calculated by [1 − (the number PFU of unbound phage / 1 × 10^5 input PFU)] × 100. Each absorption assay was repeated at least three times.

**Toxin detection in phage lysates.** Heparinized human venous blood was collected from healthy volunteers. All methods used in this study were carried out in accordance with the approved guidelines and all experimental protocols were approved by the Institutional Review Board for Human Subjects at the Mississippi State University (12–041). Informed written consent was obtained from all volunteers. Peripheral blood mononuclear cells (PBMCs) were isolated by density gradient centrifugation using Ficoll-Histopaque (Sigma-Aldrich). Purified PBMCs were adjusted to 2 × 10^8 cell per well in 96 well cell culture plate in RPMI1640 medium supplemented with 10% FBS. Phage lysates prepared from RF122Δnuc or RF122-19Δnuc was added to the wells. To detect superantigens in phage lysates, proliferation of T cell was measured using a [3H]-thymidine incorporation assay as described previously25. To detect cytotoxins in phage lysates, cytotoxicity of cells were measured using propidium iodide incorporation assay using LIVE/DEAD Cell-mediated Cytotoxicity kit (ThermoFisher).

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Author Contributions

K.S.S. and J.Y.P. conceived the project; K.S.S. and Y.H.P. designed experiments; J.Y.P. and B.Y.M. performed experiments; and K.S.S., J.A.T., J.W.P., and Y.H.P. performed the data analyses; K.S.S., B.Y.M., Y.H.P., and J.A.T. wrote the paper.

Additional Information

Supplementary information accompanies this paper at http://www.nature.com/srep
Competing Interests: Provisional patent application has been filed.

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