Bacteriophage Therapy for the Prevention and Treatment of Fracture-Related Infection Caused by *Staphylococcus aureus*: a Preclinical Study

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ABSTRACT Although several studies have shown promising clinical outcomes of phage therapy in patients with orthopedic device-related infections, questions remain regarding the optimal application protocol, systemic effects, and the impact of the immune response. This study provides a proof-of-concept of phage therapy in a clinically relevant rabbit model of fracture-related infection (FRI) caused by *Staphylococcus aureus*. In a prevention setting, phage in saline (without any biomaterial-based carrier) was highly effective in the prevention of FRI, compared to systemic antibiotic prophylaxis alone. In the subsequent study involving treatment of established infection, daily administration of phage in saline through a subcutaneous access tube was compared to a single intraoperative application of a phage-loaded hydrogel and a control group receiving antibiotics only. In this setting, although a possible trend of bacterial load reduction on the implant was observed with the phage-loaded hydrogel, no superior effect of phage therapy was found compared to antibiotic treatment alone. The application of phage in saline through a subcutaneous access tube was, however, complicated by superinfection and the development of neutralizing antibodies. The latter was not found in the animals that received the phage-loaded hydrogel, which may indicate that encapsulation of phages into a carrier such as a hydrogel limits their exposure to the adaptive immune system. These studies show phage therapy can be useful in targeting orthopedic device-related infection, however, further research and improvements of these application methods are required for this complex clinical setting.

IMPORTANCE Because of the growing spread of antimicrobial resistance, the use of alternative prevention and treatment strategies is gaining interest. Although the therapeutic potential of bacteriophages has been demonstrated in a number of case reports and series over the past decade, many unanswered questions remain regarding the optimal application protocol. Furthermore, a major concern during phage therapy is the induction of phage neutralizing antibodies. This study aimed at providing a proof-of-concept of phage therapy in a clinically relevant rabbit model of fracture-related infection caused by *Staphylococcus aureus*. Phage therapy was applied as prophylaxis in a first phase, and as treatment of an established infection in a second phase. The development of phage neutralizing antibodies was evaluated in the treatment study. This study demonstrates
that phage therapy can be useful in targeting orthopedic device-related infection, especially as prophylaxis; however, further research and improvements of these application methods are required.

**KEYWORDS** bacteriophages, *Staphylococcus aureus*, fracture-related infection, implant, rabbit

The use of fracture fixation devices has had an enormously positive impact on patient care, with shorter hospital stays and a more rapid return to function. However, a major drawback to this technological advance is the risk of developing device-related infections (1). The incidence of fracture-related infection (FRI) after internal fracture fixation ranges between 1 and 2% for closed fractures, and up to 30% for open fractures (2–5). While a variety of microorganisms can be involved, staphylococci such as *Staphylococcus aureus* are the most common causative pathogens (6, 7). Their ability to adhere to foreign objects such as fracture fixation devices, and to form antibiotic tolerant biofilms, contributes to their pathogenicity (7). As a consequence, treatment is challenging and consists of both surgical debridement and long-term antibiotic treatment. In most cases, the implant needs to be removed as the biofilm mode of growth renders conventional antibiotic treatments ineffective (6). Moreover, in chronic or late-onset cases there is a relatively high risk of treatment failure, leading to additional surgical revisions, longer length of hospital stay and increased morbidity (8). In case of antibiotic resistant pathogens, such as methicillin-resistant *S. aureus* (MRSA), treatment options may be even further restricted, and the risk of treatment failure may even be higher (9).

For the above-mentioned reasons, alternative antimicrobial strategies are being investigated, including bacteriophage (phage) therapy. The therapeutic potential of phages to support treatment of challenging cases of orthopedic device-related infections has been demonstrated in a number of case reports and series over the past decade (10–16). However, many unanswered questions remain concerning the optimal application protocol (17). Phage therapy is most often applied locally to patients with a musculoskeletal infection, whereby phage suspensions are prepared in a buffered solution and applied directly into the wound (10, 12–14, 18). This approach generally requires repeated daily dosing by an external draining system for a period of up to 10 days postoperatively. This places an additional burden on patients as they must remain in the hospital for a longer period of time with a percutaneous drain, which is prone to superinfection (12). Furthermore, preclinical models have shown that the induction of phage-neutralizing antibodies is common and can have detrimental effects on the local phage titer when treatment is prolonged (19, 20). However, it is not clear how these laboratory observations translate to phage treatments in patients. Furthermore, given the variety of phage application protocols and the diversity of phages, there is no universal predictive model of antibody induction by phages. It has been suggested that encapsulation of phages in a carrier may protect them from the adaptive immune system (19). Furthermore, analogous to antibiotic delivery vehicles such as antibiotic-loaded collagen sponges, the development of a carrier that allows a sustained release of phages would be an important clinical improvement reducing the burden on the patient, reducing risk of superinfection, and requiring fewer bedside interventions (21, 22). This may be a clinically attractive approach, although to date, only one case report has been published in the literature (15).

The aim of this study was to provide a proof-of-concept of phage therapy in a clinically relevant rabbit model of FRI caused by *S. aureus*, whereby the phage was applied as prophylaxis in a first phase, and as treatment of an established infection in a second phase. Phage therapy was applied in a prophylaxis mode in saline, and in a treatment of established infection mode as conventional delivery of repeated doses of phage suspension in saline through a subcutaneous access tube, or a single application in a phage-loaded hydrogel.
RESULTS

Animal welfare. In total, 42 animals were included in this study. Two rabbits died during induction of anesthesia for the index surgery due to unknown reasons. Two animals in the treatment study were culture-negative at revision surgery ($n=1$, Group 3 and $n=1$, Group 4), which made it impossible to assess the effect of the intervention on the infection and were, therefore, euthanized. The group allocation and study design for the prevention and treatment study are presented in Fig. 1. One animal (Group 4) died overnight and was found dead in its cage on the final study day and retained within the study. Three animals (2 animals from Group 3, 1 animal from Group 4) were euthanized 1 day prior to the scheduled euthanasia due to acute diarrhea and were retained within the study population. Parasitological analysis and bacteriology of fecal samples did not yield any identifiable cause of the acute diarrhea. All other animals tolerated surgery and survived the complete study duration. After replacement surgeries, a total of 38 rabbits were included in the final analysis, with seven to eight rabbits per group.

Prevention study. The quantitative bacteriological results from the prevention study are displayed in Fig. 2. Compared to the control group (Group 2), the group receiving phage in saline (Group 1) showed a significant reduction in bacterial load at all sampled locations (soft tissue, implant, bone and overall). The infection was eradicated in six out of eight animals receiving phage in saline. In the control group, infection was eradicated in only one out of seven. The isolated pathogens in the group that received phage in saline (Group 1) remained susceptible to ISP (intravenous...
staphylococcal phage, see Materials and Methods--Bacteriophage therapy), with a median EOP (efficiency of plating, see Materials and Methods--Phage susceptibility tests) of 1.60 (P25-P75:1.15 to 2.10), indicating no resistance development.

**Treatment study.** The results from the treatment study with the rifampicin-resistant strain are displayed in Fig. 3. The bacterial load in the soft tissue was mostly eradicated in all groups, i.e., in 6/7 control animals, 7/8 animals that received the phage-loaded hydrogel and in 6/8 animals that received phage in saline. In the bone, no infection was found in 1/8 animals that received phage in gel and in 2/8 animals that received phage in saline. The bacterial load on the implant was eradicated in 2/7 control animals, in 4/8 animals that received the phage in gel and in 3/8 animals that received phage in saline. The infection was fully eradicated in one animal in the group that received the phage-loaded hydrogel (Group 4) and in two animals in the group that received phage in buffer (Group 3). In the control group receiving systemic antibiotic therapy (Group 5), all animals were infected at euthanasia. No significant differences were found in the rate of culture positive to negative or in bacterial load between groups.

The application of phage in saline via a subcutaneous access tube was complicated due to the superinfection of the site where the tube was inserted. All animals in the group receiving phage in saline showed pus surrounding the access tube and along the drain. However, cultures after sonication of the subcutaneous access tube and drain were negative in six of these animals. In two animals receiving phages through the tube, cultures were positive for pathogens other than *S. aureus*.

**Phage susceptibility and phage-neutralizing antibodies.** Aliquots of spleen, kidney, liver, bone, soft tissue, and implant were screened for the presence of ISP at euthanasia. However, ISP could not be recovered from any testing material after propagation with the inoculating *S. aureus* JAR060131 RifR strain.

Bacterial isolates retrieved after euthanasia from the groups that received phage therapy in the treatment study were tested for their susceptibility to the applied phage. The median EOP was 1.95 (P25-P75:0.58–3.18) for the group that received phage in saline and 0.7 (P25-P75:0.55–0.87) in the group that received the phage-loaded hydrogel, indicating that the retrieved isolates remained susceptible to ISP after treatment with phage therapy (Fig. 4A). There was no significant difference between groups.

The presence of phage neutralizing antibodies in plasma at euthanasia was tested in the treatment study (Fig. 4B). The mean percentage of neutralized phages in the group receiving phage in saline (Group 3) was 45% ± 40%. Five out of eight animals receiving phage in saline developed neutralizing antibodies. No phage neutralization could be found at euthanasia in the group that received the phage-loaded hydrogel
(Group 4) or in the Control group receiving systemic antibiotics only (Group 5). This difference was statistically significant.

**Clinical observations.** In the prevention study, mean white blood cell (WBC) counts were within the normal range in both the phage treated and the control group throughout the study period (i.e., from index surgery until euthanasia) (Fig. 5A). In both groups, the mean WBC count at euthanasia did not significantly differ from the mean WBC count at the index surgery. The mean percentage weight change between euthanasia and index surgery is displayed for both groups in Fig. 5B. No significant differences were found comparing the WBC counts and weight changes between groups.

In the treatment study, the median WBC counts did not significantly differ between groups at index surgery and were within the normal range across all groups (Fig. 5C). In the control group (Group 5), there was no significant difference between the median WBC count at index surgery (day 0) and the median WBC count at euthanasia (day 28). In the group that received phage in saline (Group 3) and the group that received the

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**FIG 3** Quantitative bacteriological evaluation of soft tissue, implant and bone and the sum of all three is shown for the control animals receiving systemic antibiotics alone \( (n = 7) \), animals treated with antibiotics plus phage therapy through a subcutaneous access tube \( (n = 8) \) and animals treated with antibiotics plus phage-loaded hydrogel \( (n = 8) \). All animals received treatment 2 weeks after inoculation (i.e., established infection). The median and the interquartile range are shown for each group. No statistically significant \( (P < 0.05) \) results were found between groups. CFU, colony forming units.

**FIG 4** Phage susceptibility and neutralization tests. A. Efficiency of plating for the isolates retrieved after euthanasia of the animals in the treatment group. The median and interquartile range are plotted for each group. There was no statistically significant \( (P < 0.05) \) difference between the group receiving phage in saline or the group receiving phage in hydrogel. B. Percentage of neutralized phage particles. The differences between groups were statistically significant \( (P = 0.001) \), as indicated with an asterisk. For the group receiving phage in saline, a circle is drawn around the animals that were not infected at euthanasia.
phage-loaded hydrogel (Group 4), there was a significant difference (not shown) between the median WBC count at index surgery and the median WBC count at euthanasia. The median WBC counts at euthanasia did not differ significantly between groups. Considering the mean percentage weight change between the index surgery and euthanasia (Fig. 5D), no significant difference was observed between groups.

**DISCUSSION**

From a therapeutic viewpoint, phage therapy can be considered a natural, specific and self-limiting strategy and is currently being revisited as a sustainable approach to target (multidrug) resistant bacteria and difficult-to-treat cases. Although several reports have shown promising clinical outcomes in patients with orthopedic device-related infections, (11, 12, 14, 16), questions remain regarding the optimal application protocol, systemic effects of phage therapy and the impact of the immune response, specifically phage-neutralizing antibodies (17, 23). This proof-of-concept study evaluated the application of phage therapy for the prevention and treatment of FRI in a clinically relevant rabbit model.

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**FIG 5** White blood cell counts and weight change in the prevention and treatment study. A. Mean white blood cell counts for the animals during the prevention study. The mean and standard deviation are plotted for each group. B. Percentage weight change between euthanasia and index surgery for the animals in the prevention study. The mean and standard deviation are plotted for each group. WBC: white blood cell count. C. Median white blood cell counts for the animals in the treatment study. The median and interquartile range are plotted for each group. D. Percentage weight change between euthanasia and index surgery for the animals in the treatment study. The mean and standard deviation are plotted for each group. WBC, white blood cell count.
Prevention study. The intraoperative application of phage in saline was highly effective in preventing infection, as six out of eight animals (75%) were not infected at euthanasia. This compares favorably with the control group, in which only one out of seven animals (14%) was infection-free at euthanasia. To our knowledge, this is the only study that assessed the efficacy of phage therapy (in suspension [saline]) for the prevention of FRI caused by *S. aureus* in a clinically relevant fracture model. Two studies evaluated the use of phage therapy in a similar setting in murine models, but applied phages via local delivery systems, such as a polyethylene oxide hydrogel loaded with a phage cocktail against *P. aeruginosa* (24) and a hydroxypropyl methylcellulose coating containing a *S. aureus* phage and/or antibiotics (25). Both studies showed a statistically significant reduction in bacterial load compared to controls. Furthermore, in the study by Kaur et al., the use of a dual coated implant was more effective in reducing bacterial adherence compared to single coated or naked implants (25). The concomitant use of antibiotics with phages has been shown to work synergistically, resulting in better treatment outcomes (26). In fact, the local administration of antibiotics is gaining importance in the field of FRI for both prevention and treatment purposes. It may, therefore, be foreseen that the combination of phages and antibiotics should be investigated for direct application into surgical wounds. There may be synergistic benefits in the combination, as well as reduced risk of resistance developing. Bacterial resistance to phages generally involves cell surface receptor modifications, which hamper phage adsorption (27). Such modifications may thus lead to an increased susceptibility to antibiotics (28–30). On the other hand, some antibiotics that affect bacterial protein synthesis (such as aminoglycosides) have been shown to interfere with the phage life cycle, resulting in an attenuation of phage replication (31). Synergistic or antagonistic interactions between phages and antibiotics therefore depend on several factors, such as the class of antibiotic that is applied and its mechanism of bacterial inhibition, the receptor involved in phage adsorption and the order of treatment (i.e., simultaneous or sequential administration) (32–34). Further research is required to evaluate the extent and relevance of this phenomenon before phage therapy, alone or in combination with antibiotics, could become widely adopted in clinical practice (31).

The efficacy of phage in saline, as shown in our study, supports our initial hypothesis that prevention would require immediate release of phage and rapid distribution throughout the wound, whereas encapsulation of phage into a carrier – such as a hydrogel – functions to delay and control release. The phage-loaded hydrogel was therefore not tested in a prophylaxis setting. Indeed, in a prophylaxis setting, the bacterial inoculum is in planktonic mode, and rapid killing of bacteria before a biofilm may form, could be a relatively easier target. In the treatment setting, with biofilm, potentially fibrous tissue formation and less easy access to the surgical site, eradication of infection is clearly more challenging. Of course, in clinical reality, the target bacteria are not known in a prophylaxis setting. The clinical implementation of phage irrigation for prophylaxis would require phage cocktails to cover a broad range of potential contaminating microorganisms (based on local epidemiology). That is also the case, to a certain degree, for the local administration of antibiotics, for instance for the prevention of FRI after an open fracture where broad spectrum antibiotics may be used but may not cover all potential pathogens adequately. Nevertheless, our study supports the concept that phages could offer good protection in a prophylactic setting.

Treatment study. Infection was eradicated in one animal and in two animals of the groups that received the phage-loaded gel or phage in saline, respectively, while all animals in the control group were infected. However, these results were not statistically significant. No phages could be recovered from the dissected tissues in both treatment groups. This may potentially be attributed to the wash-out period of 1 week after cessation of treatment prior to euthanasia, which may have also allowed animals that were less infected at the end of treatment to relapse and regrow high numbers of bacteria.
**Phage in saline.** As stated above, no significant bacterial load reduction was found for animals that received phage in saline through a subcutaneous access tube. Furthermore, the application of phages was complicated due to the superinfection of the site where the access tube was inserted. That is, abscesses were found surrounding the tube, which were culture negative in six out of eight animals, while in two animals other bacterial species were isolated (other than *S. aureus*). This was not found in the group receiving the phage-loaded hydrogel, which again highlights the fact that the presence of an access tube or draining system is at risk of superinfection. This risk may be higher in a preclinical setting compared to a clinical setting where the conditions are more suitable to maintain sterility of the wound during patient care. To our knowledge, this is the first preclinical study that used a subcutaneous access tube to apply phage therapy. Previously published preclinical studies generally use repeated local injections of phage in suspension, either directly into the infected site (35) or intraperitoneally (36). Although these studies seem to show greater efficacy of phage therapy in treating implant-related bone infection, their route of administration (through repeated local administrations) does not seem clinically realistic. That is especially the case in patients with FRI or osteomyelitis, where the infection may be extensive, including an entire bone. A local delivery system (such as a hydrogel) may be more useful in such cases as it can be applied directly to the infected site and allows a sustained release. Furthermore, because such a delivery system only needs to be applied once intraoperatively, the risk of superinfection decreases.

**Neutralizing antibodies.** A major concern for the application of phage therapy is the induction of phage neutralizing antibodies (21). In our study, phage neutralizing antibodies were detected only in the group receiving phage in saline. In this group, a mean percentage of 45% ± 40% of phages were neutralized at plasma dilution 1:100. The intensity of anti-phage responses depends on several factors, such as the phage strain that is applied, with possible cross-reactivity between similar phages, the administration route and the overall immune status of the patient. Furthermore, higher phage titers and repeated administrations induce faster, stronger and more efficient anti-phage responses compared to a single administration (19).

The induction of phage neutralizing antibodies during initial phage therapy does not necessarily imply worse treatment outcomes (19, 20, 37). In fact, in our study, the animals that did not develop phage neutralizing antibodies were still infected. Plasma of two animals showed phage neutralization of more than 90%. One of these animals was not infected at euthanasia. We therefore cannot conclude that there is a correlation between our results and the development of phage neutralizing antibodies. An efficient and specific immune response also requires sufficient time (19). Several in vivo studies have shown that the neutralizing antibodies that were produced during the first phage administrations may cause a decrease in local phage titer during a second round of phage therapy and can thus potentially contribute to treatment failure (38). However, it is not known how this extrapolates to phage therapy in human subjects (19, 39). In a report of four patients with a musculoskeletal infection who were treated locally with phage therapy (including ISP) for seven to 10 days through an external draining system, no neutralizing phage antibodies were found during and up to 3 months after treatment (12), which may indicate a discrepancy between the human and rabbit immune response to phage treatment. Another study evaluated the anti-phage activity in sera from 122 patients who were treated at the phage therapy unit in Wroclaw, Poland (20). These patients received phage therapy for a variety of infectious diseases, including bone infections, and causative pathogens. In this study, higher phage inactivation rates did not correlate with a worse clinical outcome, although since the anti-phage activity of sera after phage therapy varied from patient to patient, it is difficult to generalize these results (20). Furthermore, the phages that were used were not purified (phage lysates were used), and phage therapy was continued for several weeks, which may have contributed to the results (20). This is supported by a recent case study by Dedrick et al. (40). They report on the intravenous administration
of a phage cocktail in an 81-year-old patient with bronchiectasis and refractory *Mycobacterium abscessus* lung disease. After 2 months, the phage neutralizing immune response increased, resulting in limited therapeutic efficacy as *M. abscessus* counts increased (40). Interestingly, the same administration protocol was used in a 15-year-old immunocompromised posttransplant patient with cystic fibrosis (41). In this patient no phage neutralizing antibodies were found, and the infection was eradicated after phage treatment (41). While there are other differences between both case reports such as age and cystic fibrosis status, the most striking difference was the immune status (40).

These studies underline the fact that knowledge on phage neutralizing antibodies is still very limited and the extreme diversity of phages and administration protocols does not allow us to draw general conclusions (19, 20, 37, 39). To compensate for the effects of neutralizing antibodies, it may be useful to adjust dosing to avoid unnecessary exposure of the immune system to phages and to monitor the immune reaction during therapy (19). Furthermore, phage encapsulation in for instance a phage-loaded hydrogel may protect phages from (early) immune neutralization (10, 19, 21, 39, 42). Indeed, in our study, no phage neutralization was detected in the group that received the phage-loaded hydrogel. Further investigation is therefore warranted to fully understand the mechanisms underlying this as it may prove to be a key benefit of biomaterials-based encapsulation strategies to extend the window for phage efficacy in therapeutic applications.

**Phage-loaded hydrogel.** A possible trend of minor bacterial load reduction (1 log reduction in median CFU) was observed with the phage-loaded hydrogel on the implant (where the infection was eradicated in 4/8 animals), which was achieved in only 2/8 rabbits receiving antibiotics alone, although this was not statistically significant. Similar results were found by Cobb et al. who applied an alginate hydrogel loaded with a CRISPR-Cas9-modified *S. aureus* bacteriophage (10⁷ PFU/ml) in a rat model of implant-associated osteomyelitis (43). The bacterial load in the soft tissue was significantly reduced in the groups receiving phage therapy or antibiotic therapy, but the effect of phage treatment was comparable to that of antibiotics. However, this was not observed in the bone, as only a high dose of phosphomycin (3g) could significantly reduce the bacterial load (43). The authors attributed this to the relatively low phage titer in the hydrogel (10⁷ PFU/ml) (43). In general, phage loading inside biomaterials consists of higher phage titers compared to the phage titers used in recent phage treatment protocols applied in saline suspensions (12, 21). It is hypothesized that with higher titers within the biomaterials, a sustained release of adequate phage titers may be maintained resulting in prolonged therapeutic efficacy. However, evidence regarding the optimal or minimal effective phage titer is scarce and is subject to phage-specific as well as patient-specific variability (21). It may also be that in our study, the dual application of antibiotic and phage masked the contribution of each. We did not include a phage treatment only group, as clinical treatment, even with phage therapy, still includes standard of care antibiotic therapy. It could be that the antibiotic therapy reduced the bacterial load rendering fewer bacterial hosts to support propagation of phage within the wound. *In vitro* data suggest that when combining antibiotic and phage, it may be beneficial to apply the phage initially, and follow that with antibiotic therapy to ensure maximal effect (26, 44). Further development of the phage-loaded hydrogel may benefit from dual application, with controlled and staggered release of antibiotic and phage.

A recent case report describes the use of a phage-loaded hydrogel as a salvage procedure to treat a human patient with an infected knee megaprosthesis (15). During a DAIR (debridement antibiotics and implant retention) procedure, a Defensive Antibacterial Coating (DAC) hydrogel loaded with phages against *S. aureus* (10⁹ PFU/ml) was applied (15). This study suggests the practical feasibility of the application of a phage-loaded hydrogel carrier, especially in cases where a DAIR procedure is the only option and the prosthetic surface is large. Further research is required to assess the
COMPATIBILITY OF DIFFERENT PHAGES WITH A HYDROGEL CARRIER AND ITS EFFICACY TO ERADICATE THE INFECTION (15). ALTHOUGH THE SAME PHAGE TITER OF \(10^9\) PFU/ml (AS IN THE AFOREMENTIONED CASE STUDY) WAS APPLIED IN OUR STUDY AND THE ISP WAS STABLE WITHIN THE HYDROGEL (DATA NOT SHOWN), WE DID NOT OBSERVE A STATISTICALLY SIGNIFICANT REDUCTION OF THE BACTERIAL LOAD IN VIVO, COMPARED TO THE CONTROL GROUP. FURTHERMORE, NO PHAGES COULD BE RECOVERED FROM THE TISSUES (BONE, SOFT TISSUE, IMPLANT OR ORGANS) SUGGESTING THAT PHAGE WAS NOT PRESENT IN THE TISSUE FOR AN EXTENDED PERIOD OF TIME. EMBEDDING, ENCAPSULATING, OR ADSORBING PHAGES INTO CARRIERS OFFERING CONTROLLED RELEASE OVER A LONGER PERIOD OF TIME MAY THEREFORE BE BENEFICIAL IN TREATING ESTABLISHED INFECTION. FURTHER RESEARCH, INVOLVING DIFFERENT PHAGE TYPES, CONCENTRATIONS, OR DELIVERY VEHICLES, IS THEREFORE ADVISABLE.

CONCLUSIONS

This study provides a proof-of-concept regarding the application of phage therapy in a clinically relevant model for FRI. The intraoperative application of phage in saline prevented the development of FRI compared to control animals that only received antibiotic prophylaxis. However, further research is required to evaluate whether phage therapy can be applied in clinical prevention settings. In a treatment setting, phage therapy via a subcutaneous access tube approximates the clinical situation, but this approach was complicated by superinfection in our model and the development of phage neutralizing antibodies. Phage neutralization was not found in the plasma of the animals that received the phage-loaded hydrogel, which may indicate that encapsulation of phages into a carrier such as a hydrogel limits their exposure to the adaptive immune system. Further research and improvements of these application methods are required.

MATERIALS AND METHODS

Animal care and use, surgical procedure, and study design. This study was approved by the Ethical Committee of the canton of Grisons in Switzerland (approval numbers 07_2019, 03E_2020, 14_2020). All procedures were performed in an AAALAC (Association for Assessment and Accreditation of Laboratory Animal Care International)-approved facility and according to Swiss animal protection laws and regulations. Skeletally mature specific pathogen-free (SPF) female New Zealand White rabbits (Charles River, Sulzfeld, Germany) between 20 and 32 weeks of age and with a mean body weight of 3.3 ± 0.8 kg (range 3.0 kg to 4.1 kg) were used in this study. All animals were screened prior to entry into the study and found to be healthy after a clinical examination and evaluation of the hematocrit and WBC count (Vet ABC, Sci animal care, Viernheim, Germany). All animals were allowed to acclimatize to their surroundings for 2 weeks prior to the start of the study. During this time, they were group housed with a 12h dark – 12h light cycle, fed with hay, lettuce and supplemental feed for rabbits (Granovit AG, Lucens, Switzerland).

The study design is displayed in Fig. 1. All animals underwent index surgery, including a humeral osteotomy and plate osteosynthesis (Fig. 6A and B). The animal experiments were performed in two phases. First, we tested phage therapy in a prevention setting (Fig. 1A), whereby bacterial inoculation and phage delivery were applied during the same procedure. All animals received antibiotic prophylaxis, to simulate current clinical practice. Test groups received either phage in saline (Group 1) or no treatment (Group 2), with a target of eight rabbits per group. In the second phase of this study, we assessed the ability of phage therapy to support conventional antibiotic therapy of an established infection (Fig. 1B). The same animal model was used as in the prevention study, but prophylactic antibiotics were omitted and the bacterial infection was allowed to develop for 2 weeks before treatment commenced. Treatment groups included phage in saline with concomitant antibiotic therapy (Group 3), a phage-loaded hydrogel with concomitant antibiotic therapy (Group 4), and antibiotics only (Group 5), with a target of eight rabbits per group. Group sizes were calculated in both studies based on an effective difference of 80% (i.e., an infection rate reduction of 80%). For all groups, outcome was measured by quantitative bacteriology, radiographic evaluation and blood work, including WBC count and the detection of phage neutralization caused by phage-neutralizing antibodies in blood plasma.

Surgery was performed as described previously by Arens et al. (45). In summary, a middiaphyseal osteotomy was created with a 0.44 mm Gigly saw (RISystem AG, Landquart, Switzerland) and fixed with a seven-hole locking plate and six 2 mm locking screws. Inoculation was performed by pipetting 34 μl of bacterial suspension of S. aureus JAR060131 onto the central screw hole overlying the osteotomy and to the adjacent proximal and distal screw holes. In the prevention study (Fig. 1A), all animals received a single shot of cefuroxime (Sandoz AG, Basel, Switzerland) intravenously, at a dosage of 18.75 mg/kg (weight-adjusted dosage based on human medicine for a nominal human with a bodyweight of 80 kg receiving a standard dosage of 1500 mg) (46). In the animals that received phage therapy (Group 1), the first layer was temporarily closed after
(MHB containing 20% glycerol) were prepared and stored at −20°C. The antibiotic-resistant isolate was recultured (strain here named JAR060131RifR) in fresh MHB without antibiotics. Frozen glycerol stocks were incubated statically at 37°C for 24 h. Next, the highest concentration with visible bacterial growth was used to prepare MHB containing rifampicin (Sigma-Aldrich, St. Louis, MO, USA) at concentrations ranging from 0.25 to 256 mg/liter. Bacterial suspensions (40 mg/kg four times per day) and rifampicin orally (40 mg/kg two times per day) for 7 days after the re-surgery. During this week, the animals of Group 3 received phage suspended in normal saline as 1.3 ml K3E 1.6 mg EDTA/ml, Sarstedt AG, Nümbrecht, Germany) was collected through the ear vein at 20°C for the phage neutralization assay. Weight was measured at surgery, after 3 days and weekly thereafter. Radiographs of the operated limb were taken in two planes after each surgery to check alignment and fixation, and at the end of the study to check for radiographic signs of infection. After the study period, all animals were humanely euthanized using intravenously administered pentobarbital (Esonarikon, Streuli pharma AG, Uznach, Switzerland). Animal welfare. After the initial surgery, the animals were single-housed for the entire duration of the study. All animals were checked daily by a veterinarian or experienced animal caretaker. Blood samples were taken preoperatively, 3 days postoperatively and weekly thereafter until the end of the study period. The WBC count was determined (Vet ABC, Scil animal care, Viernheim). Blood plasma (micro tube 1 ml) was collected with a syringe. In Group 4, 800 µl of phage-loaded hydrogel (ISP titer of 108 PFU/ml) was applied on the surface of the plate, after which the wounds were closed. The control animals (Group 5) only underwent debridement and irrigation of the wound, after which the incisions were closed. All animals in the treatment groups (Groups 3–5) received nafcillin subcutaneously (40 mg/kg four times per day) and rifampicin orally (40 mg/kg two times per day) for 7 days after the re-surgery. During this week, the animals of Group 3 received phage suspended in normal saline as 1 ml injections (108 PFU/ml) directly into the infected site via the subcutaneous access tube, two times per day. After a subsequent 1-week wash-out period with no antibiotic or phage treatment, all animals were euthanized and bacterial analysis was performed.

Bacteria and inoculum preparation. The clinical S. aureus strain JAR 060131 (CCOS number 890, Culture collection of Switzerland, Wädenswil, Switzerland), isolated from a patient with an orthopedic device-related infection, was used for preparation of the inoculum. This strain was used for the prevention study and was susceptible to cefuroxime, which was used as antibiotic prophylaxis. The treatment study was initially set up using the same bacterial strain. However, antibiotic treatment was effective in seven out of eight control animals (88%, results not shown). To have a better model to evaluate the effects of phage therapy in a treatment setting, this experiment was repeated with a rifampicin-resistant isolate of the initial strain. To generate the rifampicin-resistant mutant, JAR060131 strain was cultured in Müller-Hinton broth (MHB, Oxoid Ltd., Hampshire, United Kingdom) containing rifampicin (Sigma-Aldrich, St. Louis, MO, USA) at concentrations ranging from 0.25 to 256 mg/liter. Bacterial suspensions were incubated statically at 37°C for 24 h. Next, the highest concentration with visible bacterial growth was recultured (strain here named JAR060131RifR) in fresh MHB without antibiotics. Frozen glycerol stocks (MHB containing 20% glycerol) were prepared and stored at −20°C for further examinations.
Resistance to rifampicin was confirmed by the disk diffusion zone of inhibition (ZOI) method following the EUCAST instruction, using 5 μg rifampicin disks (Oxoid Ltd., Hampshire, United Kingdom). The ZOI was measured directly after generating rifampicin resistance, and after 4 and 24 weeks recovery from frozen storage. Stability of rifampicin resistance in JAR-060131RifR was further checked after serial passage (10 times) from one antibiotic-free MHA plate to a fresh plate. Each time, the zone of inhibition from a single colony was measured and was 0 mm compared to the parent control strain JAR060131, which had a 34/35 mm ZOI indicative for a stable rifampicin resistance in JAR060131RifR. Furthermore, this strain was susceptible to nafcillin, which was used in combination with rifampicin in the treatment study.

One day before surgery, an overnight culture (rifampicin sensitive or resistant, as appropriate) was prepared by suspending a single bacterial colony in 50 ml Tryptic Soy Broth (TSB; Oxoid Ltd., Hampshire, United Kingdom) at 37°C while shaking. From the overnight culture, a fresh logarithmic phase culture was prepared in TSB. Bacteria were washed twice in Phosphate buffered saline (PBS, Sigma-Aldrich, St. Louis, MO, USA) 1.5h prior to surgery and diluted in PBS to obtain a target inoculum of 2.0 × 10^8 CFU (CFU) of S. aureus. Immediately after inoculum preparation, quantitative cultures were performed by plating a dilution series to ensure the accuracy of the used inocula.

Bacteriophage therapy, ISP, a monophage targeting S. aureus was used with permission from the Eliava Institute, Georgia. This phage has been characterized in detail: its genetic sequence and therefore the confirmation of its strictly lytic profile and absence of toxins or antibiotic resistance genes is available (47, 48). Phage ISP has a contractile tail and belongs to the Kayvirus genus (Herelleitidae, Twortvirinaceae). ISP is commonly applied therapeutically in patients and has also been used in preclinical research (12, 49). The ability of ISP to infect S. aureus JAR060131 was tested using the spot test and double agar overlay method as previously described (50, 51).

Preparation of phage-loaded hydrogel. All reagents for the preparation of the emulsion-based hydrogel (EBH) were purchased from Sichma Aldrich, St. Louis, MO, USA. EBH was prepared 7 days prior to application. The EOP was determined, which is defined as the number of PFU a phage is able to produce when incubated with the initial (reference) bacterial strain.

Bacterial quantification. After euthanasia, the humerus was harvested. Soft tissue, abscesses and implants were collected and cultured. Soft tissue overlying the plate was homogenized (Omni TH, tissue homogenizer TH-02/TH21649) in 5 ml Tryptic Soy Broth (TSB; Oxoid Ltd., Hampshire, United Kingdom) at 37°C while shaking. From the overnight culture, a fresh logarithmic phase culture was prepared in TSB. Bacteria were washed twice in Phosphate buffered saline (PBS, Sigma-Aldrich, St. Louis, MO, USA) 1.5h prior to surgery and diluted in PBS to obtain a target inoculum of 2.0 × 10^8 CFU (CFU) of S. aureus. Immediately after inoculum preparation, quantitative cultures were performed by plating a dilution series to ensure the accuracy of the used inocula.

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Phage susceptibility tests. In the treatment group, each isolated pathogen at euthanasia was tested for susceptibility against ISP. The EOP was determined, which is defined as the number of PFU a phage is able to produce when incubated with the bacterial strain isolated at euthanasia relative to the number of PFU the phage is able to produce when incubated with the initial (reference) bacterial strain.

Phage titration and phage neutralization assay. In all animals that were treated with phage therapy, the presence of phages in the bone, soft tissue, surrounding the implant and in the organs (i.e., spleen, liver, kidneys) was measured at euthanasia according to the protocol described by Dufour et al. (52). The spleen, liver and kidneys were dissected as these are considered the major organs responsible for phage clearance, and homogenized (Omni TH, tissue homogenizer TH-02/TH21649) in 5–20 ml PBS (Sigma-Aldrich, St. Louis, MO, USA). An aliquot of each tissue sample was obtained and centrifuged at maximum speed for 5 min at 4°C to pellet cells and debris. The supernatant was then filtered (0.45 μm, Millex, Merck Millipore, Ireland). Serial dilutions were made in SM phage buffer followed by phage titrations using the double agar overlay method, as previously described (12). The JAR 060131 strain was used to perform titrations.

To determine if phage neutralization occurred by the development of neutralizing antibodies in the rabbits during treatment, phage neutralization by plasma was performed using the modified Adams protocol, as previously described (12). Samples were diluted 1:100 in normal saline at 37°C. 900 μl of diluted serum was incubated with 100 μl of phage (titer 10^10 PFU/ml) for 30 min at 37°C. After this time interval this mixture was diluted 100 times in cold normal saline and phage titration was performed using the double-agar overlay method.
**Data analysis.** Data were analyzed using SPSS statistics Version 27 (IBM, Chicago, IL, USA). Normality of continuous data was tested with the Shapiro-Wilk test and homogeneity of variances was tested using the Levene’s test. In case of parametric data, a one-way ANOVA or Student’s t-test (with either equal variance assumed or not) was used to compare differences between groups. In case of nonparametric data, the Kruskal-Wallis or Mann-Whitney U test was used, as appropriate. Paired data were evaluated using the paired samples t test or the Wilcoxon rank sum test, as appropriate. P values below 0.05 were considered significant. Data were visualized using GraphPad Prism 8 (GraphPad Software, San Diego, CA, USA).

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**REFERENCES**

1. Zimmerli W, Trampuz A. 2013. Biomaterial-associated infection: a perspective from the clinic. In Moriarty TF, Zaai SAJ, Busscher HJ (ed), Biomaterials associated infection: immunological aspects and antimicrobial strategies. Springer, New York, New York.

2. Gobbel C, Berger U, Schandelmair P, Greitbauer M, Schauwecker HH, Applegate B, et al. 2001. Rates and odds ratios for complications in closed and open tibial fractures treated with unreamed, small diameter tibial nails: a multicenter analysis of 467 cases. J Orthopaedic Trauma 15:415–423. https://doi.org/10.1097/00005131-200108000-00006.

3. Boxtma H, Broekhuizen T, Patka P, Oosting H. 1996. Randomised controlled trial of single-dose antibiotic prophylaxis in surgical treatment of closed fractures: the Dutch Trauma Trial. Lancet 347:1133–1137. https://doi.org/10.1016/S0140-6736(96)90606-6.

4. Patzakis MJ, Wilkins J. 1989. Factors influencing infection rate in open fracture wounds. Clinical Orthopaedics and Related Research 234: 36–40.

5. Metsemakers WJ, Kuehl R, Moriarty TF, Richards RG, Verhofstad MH, Borens O, Kates S, Morgenstern M. 2018. Infection after fracture fixation: current trends and microbiological concepts. Injury 49:511–522. https://doi.org/10.1016/j.injury.2016.09.019.

6. Depypere M, Morgenstern M, Kuehl R, Senneville E, Moriarty TF, Obremseyk WT, Zimmerli W, Trampuz A, Lagrou K, Metsemakers WJ. 2020. Pathogenesis and management of fracture-related infection. Clin Microbiol Infect 26: 572–578. https://doi.org/10.1016/j.cmi.2019.08.006.

7. von Eiff C, Jansen B, Kohnen W, Becker K. 2005. Infections associated with medical devices: pathogenesis, management and prophylaxis. Drugs 65: 179–218. https://doi.org/10.2165/00003495-200505020-00003.

8. Bezstarosti H, Metsemakers WJ, van Liest Hesham, Voskamp Lw, Kortnam K, McMally MA, Marais LC, Verhofstad MH. 2021. Management of critical-sized bone defects in the treatment of fracture-related infection: a systematic review and pooled analysis. Arch Orthop Trauma Surg 141: 1215–1230. https://doi.org/10.1007/s00402-020-03525-0.

9. Morgenstern M, Enrichs C, Militz M, Xie Z, Peng J, Stannard J, Metsemakers W-J, Schafer D, Alt V, Soballe K, Nerlich M, Buckley RE, Blauth M, Suk M, Leung F, Barla JD, Yokata K, Qing B, Kates SL. 2021. The antibiotic resistance profiles of nosocomial pathogens in Germany: an analysis of 1,000,000 isolates. Clin Microbiol Infect 27:295–304. https://doi.org/10.1016/j.cmi.2020.05.014.

10. Onsea J, Wagemans J, Pirnay JP, DI Luca M, Gonzalez-Moreno M, Lavigne R, Trampuz A, Moriarty TF, Metsemakers W-J. 2020. Bacteriophages in treatment of musculoskeletal infections: development of a standardized multidisciplinary treatment protocol. Viruses 11:891. https://doi.org/10.3390/v11100891.

11. Ferry T, Boucher F, Fevre C, Perpont T, Chateau J, Petitjean C, Josse J, Chadic C, L’hostis G, Lebouguer G, Laurent F, Lyon Bone and Joint Infect Study Group. 2018. Innovations for the treatment of a complex bone and joint infection due to XDR Pseudomonas aeruginosa including local application of a selected cocktail of bacteriophages. J Antimicrob Chemother 73:2901–2903. https://doi.org/10.1093/jac/dky263.

12. Ferry T, Lebouguer G, Fevre C, Henry Y, Conrad A, Josse J, Batailler C, Chadic C, Medina M, Lustig S, Laurent F, Lyon BJL Study Group. 2018. Salvage debridement, antibiotics and implant retention (“DAIR”) with local injection of a selected cocktail of bacteriophages: is it an option for an elderly patient with relapsing staphylococcus aureus prosthetic joint infection? Open Forum Infect Dis 5:soy269. https://doi.org/10.1093/ofid/foy269.

13. Ferry T, Batailler C, Petitjean C, Chateau J, Fevre C, Forestier E, Brosset S, Lebouguer G, Kolenda C, Laurent F, Lustig S. 2020. The potential innovative use of bacteriophages within the DAC(R) hydrogel to treat patients with knee megaprostheses infection requiring “debridement antibiotics and implant retention” and soft tissue coverage as salvage therapy. Front Med (Lausanne) 7:342. https://doi.org/10.3389/fmed.2020.00342.

14. Tkhlaishvili T, Winkler T, Muller M, Perka C, Trampuz A. 2019. Bacteriophages as adjunct to antibiotics for the treatment of periprosthetic joint infection caused by multidrug-resistant Pseudomonas aeruginosa. Antimicrob Agents Chemother 64. https://doi.org/10.1128/AAC.00924-19.

15. Onsea J, Uttybroek S, Chen B, Wagemans J, Loed C, Van Gerven L, Spriet I, Devolder D, Debayave Y, Depypere M, Dupont L, De Munter P, Peetersmans WE, Van Noort V, Merabishvili M, Pirnay JP, Lavigne R, Metsemakers W-J. 2021. Bacteriophage therapy for difficult-to-treat infections: the implementation of a multidisciplinary phage task force (the PHAGEFORCE study protocol). Viruses 13:543. https://doi.org/10.3390/v13080543.

16. Vogt D, Sperling S, Tkhlaishvili T, Trampuz A, Pirnay JP, Willy C. 2017. Beyond antibiotic therapy: future antiinfective strategies. Unfallchirurg 120:573–584. https://doi.org/10.1007/s00113-017-0374-6.

17. Gembara K, Dąbrowska K. 2021. Phage-specific antibodies. Curr Opin Biotechnol 68:186–192. https://doi.org/10.1016/j.copbio.2020.11.011.

18. Lusiak-Szelachowska M, Zaczek M, Weber-Dąbrowska B, Międzybrodzki R, Klak M, Fortuna W, Letkiewicz S, Rogóż P, Szufnarowski K, Joriczyk-Matysiak E, Owczarz B, Górski A. 2014. Phage neutralization by sera of patients receiving phage therapy. Viral Immunol 27:295–304. https://doi.org/10.1089/vim.2013.0128.

19. Rotman SG, Sumrall E, Ziaidou R, Grijsma DW, Richards RG, Elin D, Moriarty TF. 2020. Local bacteriophage delivery for treatment and prevention of bacterial infections. Front Microbiol 11:538060. https://doi.org/10.3389/fmicb.2020.538060.

20. Malik DJ, Sokolov IJ, Vinner GK, Mancuso F, Cinquerrui S, Vladisavljevic Ł, Łak M, Fortuna W, Letkiewicz S, Rogóź P, Szufnarowski K, Joriczyk-Matysiak E, Owczarz B, Górski A. 2014. Phage neutralization by sera of patients receiving phage therapy. Adv Colloid Interface Sci 249:100–133. https://doi.org/10.1016/j.cis.2017.05.014.
23. Uyttebroek S, Chen B, Onsea J, Ruythooren F, Debaveye Y, Devolder D. 2021. Lacent Infectious Diseases: Safety and efficacy of phage therapy in difficult-to-treat infections: a systematic review.

24. Wroe JA, Johnson CT, Garcia AJ. 2020. Bacteriophage delivering hydrogels reduce biofilm formation in vitro and infection in vivo. J Biomed Mater Res A 108:39–49. https://doi.org/10.1002/jbmr.a.36790.

25. Kaur S, Harjai K, Chhibber S. 2016. In vivo assessment of phage and linezolid based implant coatings for treatment of methicillin resistant S. aureus (MRSA) mediated orthopaedic device related infections. PLoS One 11: e0157626. https://doi.org/10.1371/journal.pone.0157626.

26. Wang L, Thilaishivil T, Trampuz A, Gonzalez Moreno M. 2020. Evaluation of staphylococcal bacteriophage Sb-1 as an adjunctive agent to antibiotic against rifampin-resistant staphylococcus aureus biofilms. Front Microbiol 11:62057. https://doi.org/10.3389/fmicb.2020.62057.

27. Labrie SJ, Samson JE, Moineau S. 2010. Bacteriophage resistance mechanisms. Nat Rev Microbiol 8:317–327. https://doi.org/10.1038/nrmicro2315.

28. Levin BR, Bull JJ. 2004. Population and evolutionary dynamics of phage therapy: the key is in the receptors. Curr Opin Biotechnol 68:123–129. https://doi.org/10.1016/j.copbio.2020.10.002.

29. Maciejewsk B, Olszak T, Drulis-Kawa Z. 2018. Applications of bacteriophages versus phage enzymes to combat and cure bacterial infections: an ambitious and also a realistic application? Appl Microbiol Biotechnol 102:2563–2581. https://doi.org/10.1007/s00253-018-8811-1.

30. Rohde C, Resch G, Pinay J-P, Blasdi B, Debarbieu L, Gelman D, Górska A, Hazan R, Huys I, Kakadzé E, Lobocka M, Maestri A, Almeida G, Makalata K, Malik D, Mašlanová I, Merabishvili M, Pantucek R, Rose T, Štveráková D, Van Raemdonck H, Verbeken G, Chanishivili N. 2018. Expert opinion on three phage therapy related topics: bacterial phage resistance, phage training and prophages in bacterial production strains. Viruses 10:178. https://doi.org/10.3390/v10040178.

31. Uyttebroek S, Chen B, Onsea J, Ruythooren F, Debaveye Y, Devolder D. 2020. Bacteriophage therapy in implant-related infections: an experimental model of Staphylococcus aureus induced osteomyelitis and soft tissue infection. PLoS One 14:e0220421. https://doi.org/10.1371/journal.pone.0220421.

32. Thilaishivil T, Lombardi L, Klett AB, Trampuz A, Di Luca M. 2018. Bacteriophage Sb-1 enhances antibiotic activity against biofilm, degrades exopolysaccharide matrix and targets persisters of Staphylococcus aureus. Int J Antimicrob Agents 52:842–853. https://doi.org/10.1016/j.ijantimicag.2018.09.006.

33. Arens D, Wilke M, Calabro L, Hackl S, Zeiter S, Zderic I, Richards RG, Moriarty TF. 2015. A rabbit humerus model of plating and nailing osteosynthesis with and without Staphylococcus aureus osteomyelitis. Eur Cell Mater 30: 148–161. https://doi.org/10.22203/ecm.v030a11.

34. Pueztler J, Metsemakers W-J, Arens D, Zeiter S, Kuehl R, Raschke MJ, Richards RG, Moriarty TF. 2018. Antibiotic prophylaxis with cefuroxime: influence of duration on infection rate with Staphylococcus aureus in a contaminated open fracture model. J Orthop Trauma 32:190–195. https://doi.org/10.1097/BOT.0000000000001053.

35. Merabishvili M, Monserez R, van Belleghem J, Rose T, Jennens S, De Vos D, Verbeken G, Vaneechoutte M, Pirnay J-P. 2017. Stability of bacteriophages in burn wound care products. PLoS One 12:e0182121. https://doi.org/10.1371/journal.pone.0182121.

36. Merabishvili M, Pirnay J-P, Verbeken G, Chanishivili N, Tediani M, Lashkhi N, Glonti T, Krylov V, Mast J, Van Parys L, Lavigne R, Volkaert G, Mattheus W, Verween G, De Corte P, Rose T, Jennens S, Zizi M, De Vos D, Vaneechoutte M. 2009. Quality-controlled small-scale production of a well-defined bacteriophage cocktail for use in human clinical trials. PLoS One 4:e004944. https://doi.org/10.1371/journal.pone.0004944.

37. Ngassam-Tchamba C, Duprez JN, Fergestad M, De Visscher A, L’Abée-Lund T, De Vliegher S, Wasteson Y, Touzain F, Blanchard Y, Lavigne R, Chanishivili N, Cassard M, Mainil J, Thié D. 2020. In vitro and in vivo assessment of phage therapy against Staphylococcus aureus causing bovine mastitis. J Glob Antibiotic Resist 10:1–12. https://doi.org/10.1016/j.jgar.2020.06.020.

38. Kabakadze E, Makalata K, Grdzeliashvili N, Bakuradze N, Goderdzishvili M, Kusradze I, Phoba M-F, Lunguya O, Loed C, Lavigne R, Jacobs J, Deborggraeve S, De Block T, Van Puyvelde S, Lee D, Coffey A, Sedrakyan A, Soentjens P, De Vos D, Pirnay J-P, Chanishivili N. 2018. Selection of potential therapeutic bacteriophages that lyse a CTX-M-15 extended spectrum beta-lactamase producing salmonella enterica serovar typhi strain from the Democratic Republic of the Congo. Viruses 10:172. https://doi.org/10.3390/v10040172.

39. Djebara S, Maussen C, De Vos D, Merabishvili M, Damanet B, Pang K, De Vos D, Verbeken G, Vaneechoutte M, Pirnay J-P. 2017. Processing phase therapy requests in a Brussels military hospital: lessons identified. Viruses 11:265. https://doi.org/10.3390/v11030265.

40. Dedrick RM, Freeman KG, Nguyen JA, Bahadirli-Talbott A, Smith BE, Wu AE, Ong AS, Lin CT, Ruppel LC, Parrish NM, Hatfull GF, Cohen KA. 2021. Potent antibiotic-mediated neutralization limits bacteriophage treatment of a pulmonary Mycobacterium abscessus infection. Nat Med 27:1357–1361. https://doi.org/10.1038/s41591-021-01403-9.

41. Dedrick RM, Guerero-Bustamante CA, Gariena RA, Russell DA, Ford K, Harris K, Gilmour KC, Soothill J, Jacobs-Sera D, Schooley RT, Hatfull GF, Spencer H. 2019. Engineered bacteriophages for treatment of a patient with a disseminated drug-resistant Mycobacterium abscessus. Nat Med 25:730–733. https://doi.org/10.1038/s41591-019-0437-z.

42. Dąbrowska K, Abedon ST. 2019. Pharmacologically aware phage therapy: pharmacodynamic and pharmacokinetic obstacles to phage antibacterial action in animal and human bodies. Microbiol Mol Biol Rev 83:e00012-19. https://doi.org/10.1128/MMBR.00012-19.

43. Cobb LH, Park J, Swanson EA, Beard MC, McCabe EM, Rourke AS, Seo KS, Olivier AK, Priddy LB. 2019. CRISPR-Cas9 modified bacteriophage for treatment of Staphylococcus aureus induced osteomyelitis and soft tissue infection. PLoS One 14:e0220421. https://doi.org/10.1371/journal.pone.0220421.

44. Labrie SJ, Samson JE, Moineau S. 2010. Bacteriophage resistance mechanisms. Nat Rev Microbiol 8:317–327. https://doi.org/10.1038/nrmicro2315.

45. Levin BR, Bull JJ. 2004. Population and evolutionary dynamics of phage therapy: the key is in the receptors. Curr Opin Biotechnol 68:123–129. https://doi.org/10.1016/j.copbio.2020.10.002.

46. Rohde C, Resch G, Pinay J-P, Blasdi B, Debarbieu L, Gelman D, Górska A, Hazan R, Huys I, Kakadzé E, Lobocka M, Maestri A, Almeida G, Makalata K, Malik D, Mašlanová I, Merabishvili M, Pantucek R, Rose T, Štveráková D, Van Raemdonck H, Verbeken G, Chanishivili N. 2018. Expert opinion on three phage therapy related topics: bacterial phage resistance, phage training and prophages in bacterial production strains. Viruses 10:178. https://doi.org/10.3390/v10040178.