Phage Therapy in the Era of Multidrug Resistance in Bacteria: A Systematic Review

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Abstract: Bacteriophages offer an alternative for the treatment of multidrug-resistant bacterial diseases as their mechanism of action differs from that of antibiotics. However, their application in the clinical field is limited to specific cases of patients with few or no other alternative therapies. This systematic review assesses the effectiveness and safety of phage therapy against multidrug-resistant bacteria through the evaluation of studies published over the past decade. To that end, a bibliographic search was carried out in the PubMed, Science Direct, and Google Scholar databases. Of the 1500 studies found, 27 met the inclusion criteria, with a total of 165 treated patients. Treatment effectiveness, defined as the reduction in or elimination of the bacterial load, was 85%. Except for two patients who died from causes unrelated to phage therapy, no serious adverse events were reported. This shows that phage therapy could be an alternative treatment for patients with infections associated with multidrug-resistant bacteria. However, owing to the phage specificity required for the treatment of various bacterial strains, this therapy must be personalized in terms of bacteriophage type, route of administration, and dosage.

Keywords: phage therapy; bacteriophages; multidrug-resistant bacteria; infections

1. Introduction

Bacteriophages were discovered by Frederick Twort and Félix d’Hérelle in 1915 and 1917, respectively. Since then, it has been suggested that the administration of these viruses could be used to treat bacterial infections. Initial results with phage therapy (PT) were promising; however, its application was limited owing to the discovery of penicillin in 1928 as well as the large-scale production of antibiotics in 1940 [1–4]. In recent years, the use of PT has recommenced owing to the increased bacterial resistance to antibiotics, which can arise because of mutations or be acquired by the acquisition of resistance-conferring genes via horizontal gene transfer (HGT). HGT may occur via conjugation, transformation, and transduction processes as well as via mobile genetic elements, such as transposons, and insertion sequences [5,6].

Infections caused by multidrug-resistant bacteria have become one of the leading causes of morbidity and mortality worldwide, with approximately 700,000 deaths each year [7,8]. The World Health Organization (WHO) has listed critical priority pathogens that require new antibiotics or therapeutic alternatives; these pathogens include multidrug-resistant bacteria that are especially dangerous in hospitals and nursing homes and for patients who need to be treated with ventilators and intravenous catheters [9]. According to the Organization for Economic Cooperation and Development (OECD), in Europe, North America, and Australia alone, 2.4 million people could die between now and 2050 if current bacterial resistance rates continue [10], which evidences that bacterial resistance poses...
a global public health problem [2–6,11]. Likewise, the discovery and development of new antibiotics have decreased because of the large amount of time and money required, resulting in increasingly difficult clinical management of infections and, in some cases, infections that are impossible to treat [9,11,12]. For these reasons, there is a need to look for alternatives for patients for whom conventional antibiotic therapy is ineffective. Thus, PT has become an alternative as its mechanism of action differs from that of antibiotics [13].

Lytic bacteriophages have had an impact on the clinical treatment of multidrug-resistant bacteria because of their capacity to naturally control bacterial populations [14,15]. These viruses provide novel advantages, such as the safe treatment of infections, as they are harmless to eukaryotic cells, do not cause harmful side effects, and demonstrate high host specificity. This is because they only replicate in the presence of bacteria causing the infection, thus reducing damage to the natural microflora. In addition, genetic exchange between phages rarely happens [16–18]. Therefore, as the frequency of therapy increases, the treatment doses and periods needed to achieve an optimal effect are expected to decrease [16,17]. Phage therapy has been used for the treatment of infections related to burn injuries or soft tissue and skin trauma, osteomyelitis, sepsis, bacteremia, and otitis media as well as urinary tract, pulmonary, and prosthetic device-associated infections. It is often used after the generally prolonged stay of the patients, who have been treated with different antibiotics without being able to eradicate the infection successfully [19,20]. However, effectiveness- and safety-related data for this type of therapy are scattered across individual clinical case reports owing to the few clinical studies conducted. In this review, we summarize the effectiveness- and safety-related data from studies conducted on human patients infected with various types of bacteria who underwent PT as an adjuvant or alternative therapy to antibiotics.

2. Materials and Methods
2.1. Search Strategy

To carry out the bibliographic search, three databases were selected: ScienceDirect, PubMed, and Google Scholar. The keywords used were phage therapy and multidrug resistant. The search was defined with the use of Boolean operators as follows: “phage therapy” AND “multidrug resistant”. The search was adjusted in each database according to the descriptors and using “phage therapy in the era of multidrug resistance in bacteria” as a reference. The search protocol was conducted by two independent reviewers.

2.2. Inclusion and Exclusion Criteria

The review included full-text articles and clinical case reports published in Spanish and English over the last ten years, that is, from 1 January 2011 to 25 April 2021. The main objectives of these studies were to describe the use of PT in patients infected with various bacteria, provided that, at the end of the study, it would be possible to tell whether there was a positive or negative response to the treatment.

The review also excluded bibliographic material that had not been reviewed by academic peers; articles written in a language other than Spanish or English; articles published outside the selected duration; bibliographic material, such as reviews, systematic reviews, posters, conferences, book sections, and perspectives; multidrug-resistance-based studies that did not use phage therapy; studies on metagenomics in bacteriophages without subsequent application in human patients; studies on PT in animals; and studies on the use of phages in foods.

The selection of articles was carried out considering the PRISMA statement (Preferred Reporting Items for Systematic Reviews and Meta-analyses) (Figure 1).
2.3. Data Extraction

The citations selected, along with their title, were imported into EndNote, a Bibliographic Reference Management Software, and Microsoft Office Excel, which was used to check duplicated results between the three databases. Duplicated records were eliminated.

2.4. Data Analysis

Evaluation of the records was carried out by reading the titles and the abstracts. All those that did not meet the inclusion criteria were discarded. Then, the full texts were analyzed to rule out those articles that did not meet the inclusion criteria, thus leaving us with the final records that were included in this review. Subsequently, the information of each article was collected in two Excel tables, whereas the most relevant information was captured in graphs for later analysis.

3. Results

The bibliographic search yielded a total of 1500 articles distributed as follows: 203 in PubMed, 306 in ScienceDirect, and 991 in Google Scholar published between 2011 and 2021. After excluding duplicated records (336 articles) and 1121 articles that did not meet the inclusion criteria, 27 articles were included in this review (Figure 1). Subsequently, information concerning the effectiveness and safety of PT was collected along with other findings that the authors considered relevant [21].

Of a total of 165 patients who underwent phage therapy, 141 (85%) showed a reduction in or complete elimination of the bacterial load accompanied by an improvement in the signs and symptoms, whereas PT was ineffective in the remaining 24 (15%). Only 21% (n = 35) of patients received PT combined with antibiotics, with a 100% success rate. The remaining 79% (n = 130) were treated exclusively with PT, with a success rate of 81%. Bacte-
rial resistance to phages was reported in 6 of the 27 articles reviewed [22–27]; in four studies, the initial therapy was modified by changing or including new bacteriophages [22–27], whereas in the remaining studies, resistance was measured at the end of the study.

The age of the patients receiving PT ranged from 2 to 88 years, with an average of 54 years. Three of these studies were conducted on minors (12%) [19,28,29], whereas 88% included patients over the age of 18. Most cases yielded positive results in terms of effectiveness (improving the medical condition), and adverse side effects were minimal. In addition, phage administration was not associated with the death of any of the patients (Table 1).

A total of 61 multidrug-resistant bacterial species were reported, including *Staphylococcus aureus* (24.5%, *n* = 15/61), *Pseudomonas aeruginosa* (22.9%, *n* = 14/61), *Klebsiella pneumoniae* (13.1%, *n* = 8/61), and *Acinetobacter baumannii* (8.1%, *n* = 5/61) (Figure 2).

![Figure 2. Number of infections caused by each of the multidrug-resistant bacteria mentioned in the articles.](image)

In most clinical cases, phage cocktails were administered (84%, *n* = 21), i.e., a combination of 2 to 12 bacteriophages, whereas in seven cases, a single phage (16%) was administered [23,26,30–32]. The routes of administration of the phage cocktails in these 27 cases were as follows: 4 topical (16%) [23,33–35], 7 intravenous (28%) [19,25,30,33–36], 4 in organs or cavities (16%) [22,24,32,37,38], 5 through inhalation (16%) [26,27,29,39], and 6 using more than one route of administration (24%) [28,31,40–43] (Table 2).
Table 1. Data collection, after reading the full texts of the selected articles.

| Author(s)/Reference | Country | Year of Publication | Multidrug-Resistant Bacteria | Site or Type of Infection | Patients (Number/Gender/Age) | Comorbidities/ Medical History | Adverse Effects | Effect of the Treatment |
|---------------------|---------|---------------------|------------------------------|---------------------------|-----------------------------|-------------------------------|------------------|------------------------|
| Ooi, M.L. et al. [37] | Australia | 2019 | Staphylococcus aureus | Recalcitrant chronic rhinosinusitis caused by a biofilm | 9/4 men, 5 women/18-70 years of age | Endoscopic sinus surgery | Diarrhea, epistaxis, oropharyngeal pain, cough, rhinalgia, and low blood bicarbonate level | Positive treatment effect: 2 patients were cured and the others showed reduced bacterial growth |
| Schooley, R.T. et al. [25] | U.S. | 2017 | Acinetobacter baumannii | Pancreatic pseudocyst infection | 1/male/68 years of age | Diabetic, necrotizing pancreatitis | Not reported | Positive treatment effect: patient showed continuous clinical improvement, woke up from coma, showed renal improvement, and was extubated |
| Jault, P. et al. [23] | France-Belgium | 2018 | Pseudomonas aeruginosa-Escherichia coli | Infection of wounds from burns on the skin | 12/men and women/over 18 years of age | Hospitalized patients, treated for burns in a burn unit | No adverse effects related to PT were observed | No/insufficient treatment effect owing to the administration of a very low treatment dose |
| Law, N. et al. [34] | U.S. | 2019 | Pseudomonas aeruginosa | Respiratory tract infection | 1/woman/26 years of age | Patient with cystic fibrosis, persistent respiratory failure, and renal failure | No adverse effects related to PT were observed | Positive treatment effect: pneumonia clinically resolved, baseline renal function was restored, and fever disappeared |
| Aslam, S. et al. [19] | U.S. | 2019 | Pseudomonas aeruginosa-Burkholderia dolosa-Mycobacterium abscessus | Lower respiratory tract infections | 3/1 man, 2 women/67, 28, and 57 years of age | Lung transplantation and progressive kidney failure | Intravenous and nebulized formulations of bacteriophages were well tolerated without adverse effects attributed to bacteriophage administration | A positive effect of the treatment was observed in two of the three patients (one male and female, each aged 67 years) as they did not develop further infection with Pseudomonas aeruginosa, and their respiratory status markedly improved. The 28-year-old patient died. However, this was not attributed to PT, but rather to a sequence of events that led to multi-organ failure, most plausibly attributable to postsplenectomy complications and liver disease. Hence, a verdict on the treatment was not possible. |
| Nir-Paz, R. et al. [35] | U.S. | 2019 | Acinetobacter baumannii-Klebsiella pneumoniae | Left tibial infection following trauma | 1/male/42 years of age | Grade IIIA bilateral open fractures of the lower extremities: left bicondylar tibial plateau fracture with compartment syndrome and right distal femoral fracture | No adverse effects associated with PT were observed | Positive treatment effect, with healing of the graft and elimination of subclinical bone pain in the left leg. During a follow-up period of 8 months after treatment, no Ab- or Kp-positive cultures were obtained from any site. |
| Author(s)/Reference | Country       | Year of Publication | Multidrug-Resistant Bacteria | Site or Type of Infection                                                                 | Patients (Number/Gender/Age) | Comorbidities/ Medical History                              | Adverse Effects                                                                                              | Effect of the Treatment                                                                                     |
|--------------------|---------------|---------------------|------------------------------|--------------------------------------------------------------------------------------------|------------------------------|-------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------|
| Tkhilaisshvili, T.  | Germany       | 2019               | *Pseudomonas aeruginosa*-    | Chronic recurrent periprosthetic infection of the knee and chronic osteomyelitis of the femur | 1/woman/80 years of age     | Metabolic syndrome (type II diabetes mellitus and obesity) and chronic renal failure | No adverse effects related to PT were observed                                                              | Positive treatment effect. The patient did not report pain in the right knee; the soft tissue at the surgical site did not show complications; and mobility was satisfactory. All periprosthetic tissue samples collected intraoperatively resulted in negative cultures. |
| et al. [33]         |               |                     | *Staphylococcus epidermidis* |                                                                                             |                              |                                                             |                                                                                                            |                                                                                                              |
| Corbellino, M.      | Italy         | 2019               | *Klebsiella pneumoniae*     | Infections of the gastrointestinal and urinary tracts                                       | 1/male/57 years of age       | Crohn’s disease, obstructive nephrolithiasis, right nephrectomy and radical cystectomy, and stage III chronic renal failure | No adverse effects related to PT were observed                                                              | Positive treatment effect. Long-lasting multi-site colonization by an MDR Kp strain in a patient with a single kidney, a cutaneous ureterostomy, and an indwelling ureteral stent resolved after a 3-week course of PT. |
| et al. [28]         |               |                     |                              |                                                                                             |                              |                                                             |                                                                                                            |                                                                                                              |
| Duplessis, C.       | U.S.          | 2017               | *Pseudomonas aeruginosa*    | Recalcitrant infection after ASD/VSD closures                                              | 1/male (child)/2 years of age| DiGeorge syndrome and complex congenital heart disease, including a type B interrupted aortic arch, posterior misalignment of a ventricular septal defect (VSD), sub-aortic stenosis, bicuspid aortic valve, and secundum atrial septal defect | Decompensation owing to anaphylaxis, which was subsequently attributed to progressive heart failure, although the release of endotoxins as a contributing factor could not be ruled out | Clinical improvement was seen after treatment for several days with PT, however, the patient decompensated and developed severe arrhythmias and cardiac and septic shocks. This turbulent worsening was attributed to the progressive accumulation of undrained fluids, a history of influenza infection, and end-stage heart failure. Finally, the child passed away. |
| et al. [36]         |               |                     |                              |                                                                                             |                              |                                                             |                                                                                                            |                                                                                                              |
| Qin, J. et al. [24] | China         | 2021               | *Klebsiella pneumoniae*     | Multifocal urinary tract infections                                                        | 1/male/66 years of age       | The patient’s cancerous bladder was partially resected      | No adverse effects related to PT were observed                                                              | Negative treatment effect. The phage cocktail did not work. It is believed that the heterogeneous bacteria that colonized the renal pelvis could not be effectively eliminated because the plagues could not reach them. |
| Tan, X. et al. [26] | China         | 2021               | *Acinetobacter baumannii*   | Lung infection                                                                             | 1/male/86 years of age       | Exacerbation of chronic obstructive pulmonary disease and type II diabetes | No adverse effects related to PT were observed                                                              | Positive treatment effect, with elimination of the pathogen and clinical improvement of the patient’s lung function |
| et al. [29]         | Georgia       | 2018               | *Achromobacter xylosoxidans*| Chronic lung infection                                                                     | 1/female/17 years of age     | Cystic fibrosis                                            | No adverse effects related to PT were observed                                                              | Positive treatment effect; dyspnea resolved and cough was reduced. Lung function: FEV1 increased. |

| Author(s)/Reference | Country     | Year of Publication | Multidrug-Resistant Bacteria | Site or Type of Infection | Patients (Number/Gender/Age) | Comorbidities/ Medical History | Adverse Effects | Effect of the Treatment |
|---------------------|-------------|---------------------|------------------------------|---------------------------|-----------------------------|-------------------------------|-----------------|------------------------|
| Leitner, L. et al. [38] | Georgia     | 2018                | Enterococcus spp. Escherichia coli, Streptococcus spp. | Urinary tract infection | 28/men/ over 18 years of age | Transurethral resection of the prostate (TURP), with complicated or recurrent uncomplicated UTI without signs of systemic infection | No adverse effects related to PT were observed | Positive effect of treatment in 60% of patients, with normalization of urine cultures |
| Cano, E.J. et al. [30] | U.S.        | 2020                | Klebsiella pneumoniae        | Knee joint prosthesis infection | 1/male/62 years of age | Diabetes mellitus, obesity, total knee arthroplasty, multiple bacterial infections | No adverse effects related to PT were observed | Positive treatment effect. The patient showed improvement and remained asymptomatic 34 weeks after completing the treatment, in addition to showing improvement of erythema, swelling, pain, range of motion, and function of his right lower extremity. |
| Gupta, P. et al. [44] | Varanasi, India | 2019            | Escherichia coli, Staphylococcus aureus, and Pseudomonas aeruginosa | Infection of chronic wounds (ulcers) that do not heal | 20/12-60 years of age | Cystic fibrosis and double lung transplantation | No adverse effects related to PT were observed | Positive treatment effect in 7 patients who completely healed; the others showed significant improvement in terms of wound healing |
| Lebeaux, D. et al. [45] | Paris, France | 2021                | Achromobacter xylosoxidans | Persistent lung infection | 1/male/12 years of age | Cystic fibrosis and double lung transplantation | No adverse effects related to PT were observed | Positive effect. Respiratory condition gradually improved, and the bacterial load decreased. |
| Nadareishvili, L. et al. [40] | Georgia     | 2020                | Staphylococcus aureus | Osteomyelitis of the sternum and parasternal abscess | 1/male/74 years of age | Coronary artery graft, type II diabetes mellitus, high blood pressure | Not reported | Positive treatment effect. The diameter of the wound decreased and the purulent fistula closed. |
|                      |             |                     | Staphylococcus aureus       | Chronic osteomyelitis of the right tibia | 1/female/ 60 years of age | Type II diabetes mellitus | Not reported | Positive treatment effect with complete wound closure |
|                      |             |                     | Burkhodera cepacia, S. aureus, and Enterococcus fucalis | Osteomyelitis with foot ulcer | 1/male/69 years of age | Type II diabetes mellitus | Not reported | Positive treatment effect. The patient improved and the ulcer healed. |
|                      |             |                     | Pseudomonas aeruginosa, then by Staphylococcus aureus and Serratia Marcescens | Laryngeal (post-surgical) infection | 1/male/68 years of age | Removal of the larynx due to carcinoma | Not reported | Positive treatment effect. The patient improved and infection was completely cured. |
| Patel, D.R. et al. [46] | India       | 2019                | Escherichia coli, Pseudomonas aeruginosa, Staphylococcus aureus, Klebsiella pneumoniae, Proteus species, Citrobacter freundii, Morganella morganii, Acinetobacter baumannii | Chronic wound infection of the skin | 48/34 men, 14 women/12-70 years of age | 27 diabetics, 8 with high blood pressure, 2 with tuberculosis, and 10 with amputations | Not reported | Positive treatment effect in 39 patients who were completely cured; 2 patients died |
| Author(s)/Reference | Country       | Year of Publication | Multidrug-Resistant Bacteria | Site or Type of Infection                  | Patients (Number/Gender/Age) | Comorbidities/Medical History                                                                 | Adverse Effects                                                                 | Effect of the Treatment                                                                                                                                 |
|---------------------|---------------|---------------------|------------------------------|-------------------------------------------|------------------------------|-----------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------|
| Fadlallah, A. et al. [31] | Georgia      | 2015                | Staphylococcus aureus        | Eye infection/bacterial keratitis         | 1/female / 65 years of age | Postoperative corneal abscess of the left eye and interstitial keratitis after craniotomy for acoustic neurinoma | No adverse effects related to PT were observed | Positive treatment effects, with stabilization of ocular signs and negative eye and nasal cultures                                                                                 |
| Jennes, S. et al. [41] | Belgium      | 2017                | Pseudomonas aeruginosa       | Acute kidney infection after injury       | 1/male / 61 years of age | Peritonitis owing to Enterobacter cloacae and severe abdominal sepsis with disseminated intravascular coagulation secondary to strangulated diaphragmatic hernia. The patient had a prolonged hospital stay complicated by peripheral gangrene of his extremities, which resulted in the amputation of the lower limbs and the development of large necrotic pressure ulcers. | No adverse effects related to PT were observed | Positive treatment effects; blood cultures were negative, CRP levels decreased, and fever disappeared. Kidney function recovered within a few days. However, the patient died after 4 months from a sudden refractory cardiac arrest in the hospital due to Klebsiella pneumoniae. |
| Kuipers, S. et al. [42] | Holland      | 2020                | Klebsiella pneumoniae        | Recurrent urinary tract infection         | 1/male / 58 years of age | Kidney transplant | No adverse effects related to PT were observed | Positive treatment effects. The symptoms of urethritis quickly disappeared. |
| Onsea, S. et al. [43] | Belgium      | 2019                | P. aeruginosa and S. epidermidis | Chronic osteomyelitis of the pelvis       | Patient 1 | Isolated fibrous tumor in the left pelvic region | Not reported | Positive treatment effects. C-reactive protein and white blood cell levels returned to normal. |
| Onsea, S. et al. [43] | Belgium      | 2019                | P. aeruginosa and S. epidermidis | Non-union of distal femur | Patient 2 | Polyttrauma after aggression, with open segmental fractures of the right femur | Not reported | Positive treatment effects. The levels of C-reactive protein and white blood cells returned to normal. Infection-free. |
| Onsea, S. et al. [43] | Belgium      | 2019                | S. agalactiae and S. aureus | Postoperative problems of the femur wound | Patient 3 | Polyttrauma after the collapse of a building, with crush injuries of the upper right leg and complex fractures of the femur | Not reported | Positive treatment effects. C-reactive protein and white blood cell levels returned to normal. |
| | | | E. faecalis | Infection of the surgical site with abscess formation and evolution to osteomyelitis of the femur | Patient 4 | Polyttrauma after traffic accident, with fracture of the femur | Local redness and pain | Positive treatment effects. C-reactive protein and white blood cell levels returned to normal. |
| Author(s)/Reference | Country | Year of Publication | Multidrug-Resistant Bacteria | Site or Type of Infection | Patients (Number/Gender/Age) | Comorbidities/ Medical History | Adverse Effects | Effect of the Treatment |
|---------------------|---------|---------------------|-----------------------------|---------------------------|-------------------------------|---------------------------------|----------------|------------------------|
| Wu, N. et al. [27]  | Shanghai, China | 2021 | Acinetobacter baumannii | Respiratory tract infection acquired in the hospital | 1/male/62 years of age | Critical condition due to COVID-19 (RNA-negative) and co-infection with Candida albicans and Ralstonia mannitolipita | Cytokine storm and fever 4 h after Φ phage administration | Positive treatment effects. Clinical improvement, discharged from the hospital on day 30. |
|                     |         |                    |                             | Respiratory tract and intubation wound infections acquired in the hospital | 1/male/64 years of age | Critical condition due to COVID-19 (RNA-negative) | No adverse effects related to PT were observed | Positive treatment effects. Clinical improvement and discharge from the hospital on day 9. |
|                     |         |                    |                             | Respiratory tract infection acquired in the hospital | 1/male/81 years of age | Critical condition due to COVID-19 (RNA-negative) and co-infection with Candida albicans and Candida glabrata | No adverse effects related to PT were observed | Positive treatment effects. The infection was eliminated; however, patient died on day 10 owing to respiratory failure caused by infection with Klebsiella pneumoniae. |
|                     |         |                    |                             | Respiratory tract and bladder infections acquired in the hospital | 1/male/78 years of age | Critical status due to COVID-19 (RNA-negative) and co-infection with Candida albicans, Klebsiella pneumoniae resistant to carbapenems, and Sphingomonas paucimobilis | No adverse effects related to PT were observed | Positive treatment effect with improvement; patient was discharged from the ICU on day 7; however, he died on day 40 owing to respiratory failure. |
| Bao, J. et al. [22] | Shanghai, China | 2020 | Klebsiella pneumoniae | Recurrent urinary tract infection | 1/female/ 63 years of age | Type II diabetes and hypertension | No adverse effects related to PT were observed | Positive treatment effect. The infection was completely eradicated. |
| Rose, T. et al. [39] | Brussels, Belgium | 2014 | Pseudomonas aeruginosa and Staphylococcus aureus | Burn infection | 9/4 men, 5 women / 27–88 years of age | Not reported | No adverse effects related to PT were observed | Negative treatment effect. Low bacterial load, almost unchanged. |
| Rubalskii, E. et al. [32] | Germany | 2020 | Staphylococcus aureus, Enterococcus faecium, Pseudomonas aeruginosa, Klebsiella pneumoniae, and Escherichia coli | Patients had infections associated with immunosuppression after organ transplantation or had infections of vascular grafts, implanted medical devices, and surgical wounds | 8/7 men, 1 woman | Not reported | No adverse effects related to PT were observed | Positive effect of treatment in 7 patients who are completely cured; the other showed a significant improvement in terms of decreased bacterial load. |
Table 2. Characteristics of the bacteriophages used for the treatment of infections.

| Author(s)/Reference | Phage/Cocktail Identification | Number of Phages in the Cocktail | Phage Order or Family | Route of Administration | Concentration Per Unit Dose/Duration of Treatment | Antibiotics |
|----------------------|-------------------------------|----------------------------------|-----------------------|-------------------------|------------------------------------------------|------------|
| Ooi, M.L. et al. [37] | Cocktail AB-SA01              | 3                                | Myoviridae            | Nasal irrigation         | $3 \times 10^5$–$10^9$ PFU, for 14 days       | NA         |
| Schooley, R.T. et al. [25] | AB-Navy1-AB-Navy4-AB-Navy71-AB-Navy97-AbTP3Φ1-AC4-C1P12-C2P21-C2P24, cocktails ΦPC-ΦIV-ΦIVB | 8                                | Myoviridae-Podoviridae (AbTP3Φ1) | Intravenous-Intracavitary | $5 \times 10^9$ PFU, for 2 to 16 weeks | Meropenem, fluconazole, and minocycline |
| Jault, P. et al. [23] | Cocktail PP1131               | 12                               | Not reported          | Topical                 | $1 \times 10^6$ PFU, for 7 days                  | NA         |
| Law, N. et al. [34]   | Cocktail ABPA01               | 4                                | Not reported          | Intravenous             | $4 \times 10^9$ PFU, for 8 weeks                | Ciprofloxacin, piperacillin–tazobactam, doripenem |
| Aslam, S. et al. [19] | Cocktails AB-PA01, AB-PA01-m1, Navy 1, Navy 2 | 4, 5, 2, 3                     | Not reported          | Intravenous             | Between $5.3 \times 10^6$ PFU and $5 \times 10^9$ PFU for 3 to 5 weeks | Piperacillin-tazobactam and colistin |
|                       | Cocktail AB-PA01              | 4                                | Not reported          | Intravenous             |                                                   | Piperacillin-tazobactam, tobramycin, and inhaled colistin |
|                       | BdPF16phi4281                | 1                                |                       |                         |                                                   | Meropenem, ceftazidime-avibactam, minocycline, and tobramycin |
| Nir-Paz, R. et al. [35] | ΦAbKT21phi3 and ΦKpKT21phi1   | 2                                | Not reported          | Intravenous             | $5 \times 10^7$ PFU for 5 days                  | Colistin and meropenem |
| Tkhiailashvili, T. et al. [33] | Local phage adjuvants       | Not reported                     | Not reported          | Intravenous (local administration system)       | $10^8$ PFU for 5 days                           | Colistin, meropenem, and ceftazidime |
| Corbellino, M. et al. [28] | vB_KpnM_GF                   | Not reported                     | Not reported          | Oral and intrarectal    | Did not report concentration, for 3 weeks       | NA         |
| Author(s)/Reference | Phage/Cocktail Identification | Number of Phages in the Cocktail | Phage Order or Family | Route of Administration | Concentration Per Unit Dose/Duration of Treatment | Antibiotics |
|---------------------|--------------------------------|----------------------------------|-----------------------|-------------------------|-----------------------------------------------|--------------|
| Duplessis, C. et al. [36] | Phage cocktail | 2 | Not reported | Intravenous | $3.5 \times 10^5$ PFU, did not specify duration of treatment | NA |
| Qin, J. et al. [24] | Phage cocktails $\Phi902, \Phi JD905, \Phi JD907, \Phi JD908, JD910$ | 5 | $Podoviridae, Myoviridae$ ($\Phi JD905$) | Irrigations | $5 \times 10^8$ PFU for 4 weeks | Piperacillin-tazobactam, imipenem, amikacin |
| Tan, X. et al. [26] | Unique Ab_SZ3 phage | 1 | $Siphoviridae$ | Airway (vibrating mesh nebulizer) | Increasing concentrations: $5 \times 10^6$ on day 1 to $2.5 \times 10^7$ PFU, day 2 with $10^8$ PFU, day 4 with $10^9$ PFU, and day 13 with $5 \times 10^{10}$ PFU for 16 days | Tigecycline and polymyxin E |
| Hoyle, N. et al. [29] | Phage cocktail | 2 | $Siphoviridae$ | Inhalation (compressor nebulizer) | $3 \times 10^8$ PFU for 20 days | Piperacillin-tazobactam |
| Leiter, L. et al. [38] | PYO cocktail | Not reported | Not reported | Intravesical instillation | $10^4$ to $10^5$ PFU for 7 days | NA |
| Cano, E.J. et al. [30] | Phage KpJH46$\Phi2$ | 1 | Not reported | Peripheral intravenous | 40 doses of $6.3 \times 10^{10}$ PFU/mL for 30 min plus minocycline | Minocycline |
| Gupta, P. et al. [44] | Phage cocktail | 3 | Not reported | Topical | 0.1 mL/cm² (a concentration of $10^9$ PFU/mL from 9 to 13 days) | NA |
| Author(s)/Reference | Phage/Cocktail Identification | Number of Phages in the Cocktail | Phage Order or Family | Route of Administration | Concentration Per Unit Dose/Duration of Treatment | Antibiotics |
|---------------------|-------------------------------|----------------------------------|-----------------------|-------------------------|-----------------------------------------------|-------------|
| Lebeaux, D. et al. [45] | JW Delta, JWT, and 2-1 cocktails (APC 1.1) | 3 | Not reported | Inhalation (vibrating mesh nebulizer) | $4 \times 10^{10}$ PFU (3 nebulizations/day of 5 mL of solution). Did not specify the duration of treatment. | Imipenem |
| | Cocktail 2, plus phage JW alpha (APC 2.1) | 4 | Not reported | Administered to the lungs using a fibroscope and through nebulization | $5 \times 10^9$ PFU/mL. Did not specify the duration of treatment. | NA |
| Nadareishvili, L. et al. [40] | Staphylococcal bacteriophage, PYO bacteriophage, and SES bacteriophage | 3 | Not reported | Topical and oral | Did not specify the concentration; 10 mL once a day for 20 days and then for 2 weeks | NA |
| | Staphylococcus bacteriophage and Intesti bacteriophage | 2 | Not reported | Topical and oral | Did not specify the concentration; 10 mL once a day for 20 days initially and then for 15 days | NA |
| | Staphylococcus bacteriophage and Intesti bacteriophage | 2 | Not reported | Topical and oral | Did not specify the concentration; 10 mL once a day for 20 days initially and then for 20 days | NA |
| | Pyo bacteriophage and Intesti bacteriophage | 2 | Not reported | Topical and oral | Did not specify the concentration; 10 mL once a day for 20 days initially and then for 20 days. Solcoseryl (epithelial regenerator) was also used. | NA |
### Table 2. Cont.

| Author(s)/Reference | Phage/Cocktail Identification | Number of Phages in the Cocktail | Phage Order or Family | Route of Administration | Concentration Per Unit Dose/Duration of Treatment | Antibiotics |
|---------------------|-------------------------------|----------------------------------|-----------------------|-------------------------|-----------------------------------------------|-------------|
| Patel, D.R. et al. [46] | Phage cocktail | Not reported | Not reported | Topical | Did not specify the concentration or the duration of treatment; 500 µL/cm², 5–7 applications | NA |
| Fadlallah, A. et al. [31] | Phage SATA-8505 (ATCC PTA-9476) | 1 | Not reported | Topical (eye drops and nasal spray) and general (intravenous) | Did not specify the concentration, for 4 weeks | NA |
| Jennes, S. et al. [41] | BFC1 Cocktail | 2 | Not reported | 50 microliters administered through intravenous infusion for 6 h over 10 days. Wounds were irrigated with 50 mL of BFC1 every 8 h for 10 days. | Did not specify the concentration, for 10 days | NA |
| Kuipers, S. et al. [42] | Anti-Klebsiella pneumoniae phages | Not reported | Not reported | Oral and bladder irrigation/intravesical | Did not report the concentration, for 8 weeks | NA |
| Onsea, S. et al. [43] | COCKTAIL BCF1 | 3 | Not reported | Topical and in marrow | 10⁷ PFU/mL for 7 days | Vancomycin, rifampin, moxifloxacin |
| | COCKTAIL BCF1 | 3 | Not reported | Topical | 10⁷ PFU/mL for 10 days | Vancomycin, colistin, fosfomycin |
| | COCKTAIL BCF1 | 3 | Not reported | Topical | 10⁷ PFU/mL for 9 days | Vancomycin, clindamycin, moxifloxacin |
| | PYO phage | Not reported | Not reported | Topical | 10⁷ PFU/mL for 7 days | Amoxicillin |
| Author(s)/Reference | Phage/Cocktail Identification | Number of Phages in the Cocktail | Phage Order or Family | Route of Administration | Concentration Per Unit Dose/Duration of Treatment | Antibiotics |
|---------------------|--------------------------------|----------------------------------|-----------------------|------------------------|-----------------------------------------------|-------------|
| Wu, N. et al. [27]  | Phage 1Φ (ΦAb124) 1           | 1 Podoviridae                    | Respiratory by inhalation | 10^8 PFU/mL for 20 min, two doses with 1 h interval | NA                                             |
|                     | Cocktail 2Φ (ΦAb124 and ΦAb121) 2 | Podoviridae and Myoviridae      | Respiratory by inhalation | 1 × 10^10 PFU, did not specify concentration | NA                                             |
|                     | Cocktail 2Φ (ΦAb124 and ΦAb121) 2 | Podoviridae and Myoviridae      | Respiratory by inhalation and topical with a wet compress | 1 × 10^10 PFU, did not specify concentration | NA                                             |
|                     | Cocktail 2Φ (ΦAb124 and ΦAb121) 2 | Podoviridae and Myoviridae      | Respiratory by inhalation | 1 × 10^10 PFU, did not specify concentration | NA                                             |
| Bao, J. et al. [22] | Cocktail I (SZ-1, SZ-2, SZ-3, SZ-6) against the CX7224 strain 5 | Not reported | Bladder irrigation | 5 × 10^8 PFU per mL of each phage for 5 days | NA                                             |
|                     | Cocktail II (Kp165, Kp166, Kp167, Kp158, and Kp169) for the CX8070 strain 5 | Not reported | Bladder irrigation | Did not specify the concentration, for 5 days | NA                                             |
|                     | Cocktail III (Kp152, Kp154, Kp155, Kp164, Kp6377, and HD001) 6 | Not reported | Bladder irrigation | SMZ-TMP administered orally (800–160 mg) twice daily, plus the cocktail for 5 days | NA                                             |
| Rose, T. et al. [39] | BFC-1 cocktail (P. aeruginosa phage 14/1, S. aureus phages ISP and PNM) 3 | Podoviridae and Myoviridae | Topical | 10^9 PFU/mL, did not specify the duration of treatment | Ceftazidime or meropenem in combination with amikacin, Vancomycin or linezolid. |
| Author(s)/Reference | Phage/Cocktail Identification | Number of Phages in the Cocktail | Phage Order or Family | Route of Administration | Concentration Per Unit Dose/Duration of Treatment | Antibiotics |
|---------------------|-------------------------------|---------------------------------|----------------------|-------------------------|-----------------------------------------------|-------------|
| Rubalskii, E. et al. [32] | *Staphylococcus phage CH1, Enterococcus phage Enf1, Pseudomonas phage PA5, Pseudomonas phage PA10* | 4 | *Myoviridae, Siphoviridae, Myoviridae* | Local | $1 \times 10^8$ PFU/ml | Cefepime, daptomycin, linezolid, tobramycin |
| | *Klebsiella phage KPV811, Klebsiella phage KPV15* | 2 | *Podoviridae, Myoviridae* | Respiratory by inhalation, via nasogastric tube | $1 \times 10^8$ PFU/ml | Ceftazidime, linezolid, avibactam, colistin, meropenem, cotrimoxazole, tobramycin |
| | *Staphylococcus phage CH1* | 1 | *Myoviridae* | Local application via drainage | $1 \times 10^9$ PFU/ml | Rifampicin, flucloxacillin |
| | *Staphylococcus phage CH1* | 1 | *Myoviridae* | Local application via drainage | $1 \times 10^9$ PFU/ml | Daptomycin |
| | *Staphylococcus phage Sa30, Staphylococcus phage CH1, Staphylococcus phage SCH1, Staphylococcus phage SCH11* | 4 | *Myoviridae, Myoviridae, Podoviridae, Myoviridae* | Local application via drainage, intranasal | $1\times10^9$ PFU/ml | Daptomycin |
| | *Staphylococcus phage Sa30* | 1 | *Myoviridae* | Locally, intraoperatively mixed with fibrin glue | $4 \times 10^{10}$ PFU/ml | Sultamicillin |
| | *Escherichia phage ECD7, Escherichia phage V18* | 2 | *Myoviridae* | Locally, intraoperatively mixed with fibrin glue | $4 \times 10^{10}$ PFU/ml | Clindamycin |
| | *Pseudomonas phage PA5, Pseudomonas phage PA10* | 2 | *Myoviridae* | Locally, intraoperatively mixed with fibrin glue | $4 \times 10^{10}$ PFU/ml | Colistin, ceftazidime, avibactam |
4. Discussion

This systematic review displays effectiveness- and safety-related data as well as some relevant findings that have been reported in PT studies in humans from studies published in English over the last ten years. Based on the findings from a total of 165 patients treated in 27 selected studies, it can be concluded that PT produced encouraging results for the treatment of infections caused by various bacterial species, especially those that are difficult to manage, such as infections caused by bacteria resistant to multiple antibiotics [22,25,28,34,41]. In 85% of the cases, the therapy was successful through the administration of either a single phage or a phage cocktail. One of the primary factors responsible for this success is the use of specific bacteriophages for each bacterial strain. Although 100% of the studies reported having conducted in vitro studies of bacteriophage activity at the beginning of the therapy, in 15% of the cases, the infection had not been resolved at the end of the therapy. Some authors attributed these failures to defective modes of administration [38], a low phage concentration [23], or co-infection with other species of bacterial strains [46], rather than to a low effectiveness of the therapy.

One of the main concerns with PT is the development of phage-resistant strains [19,25]. An in vitro study by Oechslin et al. found that the frequency of spontaneous phage resistance mutations in a susceptible strain of P. aeruginosa was around $10^{-7}$. Additionally, the same authors concluded that, although some mutations confer resistance to phages, these mutations can decrease bacterial fitness when compared to normal growth under less strict in vitro conditions in animals [47]. This could explain the success of PT in some patients despite the presence of phage-resistant bacteria [23,26], which would prove that a patient’s immune system plays an important role in the success of PT in the short, medium, and long terms.

Although PT is presented as an emerging strategy to fight multidrug-resistant bacterial infections, evidence suggests that coadministration of phages and antibiotics could be a more effective strategy for the elimination of infection [19]. In cases where combined therapy was administered, patients recovered from the infection with 100% success, whereas 88% of those patients who received PT alone recovered. This result can be explained as follows: combination therapy may decrease the selection of phage-resistant bacteria [47], in addition to increasing the sensitivity to antibiotics [25,47,48]. Additionally, antibiotics could act as adjuvants that minimize the appearance of other pathogens in immunocompromised patients.

Owing to serious public health threats posed by health care-associated infections (HAIs), such as those caused by E. coli, S. aureus, K. pneumoniae, coagulase-negative Staphylococcus, and Enterococcus faecalis [49], and given the call made by the WHO for a search for new alternatives in the treatment of multidrug-resistant bacteria, especially those belonging to the ESRAPE group [7], it is not surprising that infections caused by these pathogenic bacteria were more frequently treated with PT (50.8%). A single phage or a cocktail of phages proved effective in the treatment of these infections regardless of their sensitivity or resistance to antibiotics. However, phage cocktails were more widely used (84%) to increase bacterial sensitivity to therapy.

In all studies, the administration of PT was considered safe for patients. The age of the treated patients ranged between 2 and 88 years, with an average of 55 years. Duplessis et al. (2017) reported the case of a 2-year-old patient with DiGeorge syndrome and a series of comorbidities that were difficult to manage, in addition to recalcitrant bacteremia/sepsis owing to multidrug-resistant P. aeruginosa. Phage therapy was initiated in this patient to treat the infection. Although sterile blood cultures were obtained for P. aeruginosa in the short term, the treatment had to be interrupted owing to decompensation of the infant caused by anaphylaxis. A refractory infection led to the reinitiation of PT, which once again proved effective, resulting in sterile blood cultures 24 h after therapy onset. Unfortunately, the patient’s decompensation and worsening prognosis owing to severe arrhythmias as well as cardiogenic and septic shocks (caused by the same bacteria), possibly owing to a history of influenza infection and end-stage heart failure, among others (although none of
these effects were related to the administration of PT), led relatives to withdraw care, and the child died soon afterwards [36].

Jault et al. (2019) reported the death of an adult man over 70 years of age who was recruited to participate in the controlled clinical trial PhagoBurn, once the inclusion and exclusion criteria were met. The patient received PT topically for the treatment of *P. aeruginosa*, and his death occurred sometime after completing the treatment regimen, although no association with PT was found [23].

The potential development of adverse events associated with the concentration of endotoxin residues, i.e., the amount of bacterial endotoxin lipopolysaccharide released during phage-mediated lysis [19], has apparently been discarded based on reports that the maximum dose is below 5 units of endotoxin per kg of body weight per hour (required by international regulators), and this is reflected by the absence of serious adverse events during the administration of PT in all the studies. This confirms the safety of PT for the treatment of bacterial infections.

Additionally, bacteriophages offer new advantages, namely, their high specificity toward the host cell, which would reduce the damage to the normal microbiome of the patient and decrease colonization by other pathogens in the absence of in vivo drug interactions, bactericidal activity, minimal variability in pharmacokinetics and pharmacodynamics, agnostic bacterial targeting irrespective of the antibacterial susceptibility profile, minimal environmental footprints, and potential inducement of susceptible bacterial profiles [36]. Moreover, their ability to alter the formation of biofilms enhances the action of antibiotics in combination therapies [21]. In addition, free genetic exchange of phages rarely occurs [3,17,18]. Thus, as the frequency of therapy increases, the treatment doses and periods needed to achieve an optimal effect would be expected to decrease [17,18]. However, one of the main challenges that must be faced for the definitive implementation of phage therapy is the unification of criteria and standardized procedures that allow dealing with infections caused by multidrug-resistant bacteria. This is because the few studies that have been reported present a great variability of results because they not only depend on the multidrug-resistant bacteria, but also on the type and place of the infection that they are causing. This in turn affects the dosage and application of combination therapies, which makes comparison between studies difficult. All this highlights the need to generate guidelines with unified criteria to determine the validity of phage therapy.

5. Conclusions

Phage therapy has proven to be effective and safe for the treatment of infectious diseases caused by various bacterial species, including multidrug-resistant strains. This effectiveness has been demonstrated both through local and systemic administration, with no associated serious adverse events. However, much of the success of PT depends on the selection of specific bacteriophages for each bacterial species and strain, owing to their specificity toward the host cell, which is why extensive libraries of phages are required to personalize treatments. On the other hand, coadministration of antibiotics as adjuvants is recommended as it prevents infections by non-specific bacteria and helps eliminate bacteria already sensitized by the phages.

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