Viability, stability and biocontrol activity in planta of specific Ralstonia solanacearum bacteriophages after their conservation prior to commercialization and use

Belén Álvarez1, 2, Laura Gadea-Pallás1, Alejandro Rodríguez1, Begonya Vicedo3, Àngela Figàs-Segura1 and Elena G. Biosca1*

1 Departamento de Microbiología y Ecología. Universitat de València (UV). Valencia, Spain.
2 Departamento de Investigación Aplicada y Extensión Agraria. Instituto Madrileño de Investigación y Desarrollo Rural, Agrario y Alimentario (IMIDRA). Madrid, Spain.
3 Departamento de Ciencias Agrarias y del Medio Natural. Universitat Jaume I (UJI). Castellón, Spain.

* Correspondence: elena.biosca@uv.es

Abstract: Ralstonia solanacearum is a pathogen that causes bacterial wilt producing severe damage in staple solanaceous crops. Traditional control has low efficacy and/or environmental impact. Recently, the bases of a new biotechnological method by lytic bacteriophages vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2 with specific activity against R. solanacearum were established. However, some aspects remain unknown, such as the survival and maintenance of the lytic activity after submission to a preservation method as the lyophilization. To this end, viability and stability of lyophilized vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2 and their capacity for bacterial wilt biocontrol have been determined against one pathogenic Spanish reference strain of R. solanacearum in susceptible tomato plants in different conditions and making use of various cryoprotectants. The assays carried out have shown satisfactory results with respect to the viability and stability of the bacteriophages after the lyophilization process, maintaining high titres throughout the experimental period, also with respect to the capacity of the bacteriophages for the biological control of bacterial wilt, controlling this disease in more than 50% of the plants. The results offer good prospects for the use of lyophilization as a conservation method for the lytic bacteriophages of R. solanacearum in view of their commercialization as biocontrol agents.

Keywords: plant pathogenic bacterium; phage; bacterial wilt; biological control; lyophilization

1. Introduction

Ralstonia solanacearum [1, 2] is a phytopathogenic bacterium that mainly affects crops such as potato, tomato, eggplant and pepper, and some ornamental species such as geranium, producing vascular bacterial wilt disease and potato brown rot in tubers [3, 4, 5]. This pathogen has been reported as one of the most destructive in the world [4], due to its global extension and the importance of the damage crops, which are basic for human nutrition. It is also considered a quarantine bacterium in many countries, and a Select Agent in the USA [6, 7, 8].

This bacterial species was a member of the former “Ralstonia solanacearum species complex” presently divided into the closely related R. pseudosolanacearum, R. solanacearum and R. syzygii subsp. indonesiensis [2].

The infection usually occurs through the roots, since the pathogen is soil and water borne. Once inside the plant, R. solanacearum multiplies in the cortex and xylem, ascending to the aerial parts [9, 10, 11]. External symptoms produced by R. solanacearum in the host are mainly wilting, growth retardation, adventitious roots in the stem, and epinasty in the leaves [12]. Normally the disease progresses until plant death, due to xylem obstruction.
The great ability of *R. solanacearum* to survive in different environments that can constitute reservoirs for long periods, such as plant debris, water or soil, without altering the pathogenic character of the bacterium, makes it difficult the control of the disease in the field [13, 14, 15]. Physical and chemical methods developed to eradicate *R. solanacearum* from affected crops, as the use of fumigants and/or cupric compounds, showed variable results, besides having an impact in soils and contributing to the appearance of bacterial resistance [10, 14]. Biological control as the use of bacteriophage viruses can be an alternative to these methods, particularly by lytic bacteriophages (phages), since they can destroy the bacterial host. Their antibacterial potential against phytopathogenic bacteria allows their use as agricultural biocontrol agents in an environmentally respectful way [15-18].

Due to this potential, an efficient method of preservation and distribution is required, especially if the phages are to be marketed after large-scale production for field application. Lyophilization is one of the most commonly used conservation methods for microorganisms [19]. It allows them to be preserved for long periods while maintaining their viability. It is carried out in three phases, namely, the freezing of the sample to be treated, followed by drying that leads to the sublimation of the ice crystals [20, 21]. Applied to phages, the cost of transport and maintenance is reduced [21, 22] but, freeze drying is a great stress for viruses, since it can alter their survival and/or their lytic capacity, because it affects their structure, destabilizing it, especially protein compounds such as the capsid [23]. For this reason, lyophilization should be carried out with cryoprotective agents, which have the function of physically and/or chemically stabilize the compounds to be lyophilized in the freezing and drying processes [24]. There are many types of cryoprotectants. For bacteriophages or different types of viruses, the most common are disaccharides such as saccharose or trehalose [20, 24-27].

With respect to *R. solanacearum*, the foundations of a new biotechnological control method were recently established based on three lytic phages called vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2 with specific activity against the pathogen and biocontrol efficiency *in planta* [28-31]. However, with a view to their commercialization, there were some unknown aspects, such as the survival capacity and the maintenance of the lytic activity of these phages after subjecting them to a conservation method such as lyophilization. In this work, the viability and stability of the phages preserved by this method with various cryoprotectants against various strains of *R. solanacearum* were determined for different periods. In addition, the efficiency of the lyophilized phages in the biological control of the strain CFBP 4944 of this pathogen in susceptible tomato plants was evaluated under different conditions.

2. Materials and Methods

2.1. Bacterial strains and culture conditions

The *R. solanacearum* strains used in this work are in Table 1. Among them, the strain CFBP 4944 was the only one used in all the tests. The strains were kept in cryo-preserved stocks at -80°C with 25% (v/v) glycerol and were grown in Casaminoacids Peptone Glucose (CPG) medium (casaminoacids 0.1%, peptone 1%, glucose 0.5%) [32], with bacteriological agar (1.5%). The general medium Luria-Bertani (LB) (1% NaCl, 0.5% yeast extract, 1% tryptone) [33] was used as the liquid culture medium. The Semiselective Medium from South Africa (SMSA) (0.1% casaminoacids, 1% peptone, 0.5% glycerol, 1.5% agar, crystal violet 0.0005%, bacitracin 1,250 U/l, polymyxin sulfate B 600,000 U/l, penicillin 825 U/l, chloramphenicol 0.0005%, triphenyltetrazolium chloride 0.005%) [34] was used in some assays carried out with the strain CFBP 4944. SMSA is a semi-selective medium for *R. solanacearum* isolation.

The culture conditions of the bacterial strains were the following: in liquid medium, overnight cultures (16-18 h) in 5 ml of LB and incubated at 28°C with shaking (120 rpm), and in solid medium, incubation at 28°C for 48-72 h.
Table 1. Strains of *Ralstonia solanacearum* used in the present study.

| Strains                      | Host            | Country   |
|------------------------------|-----------------|-----------|
| IVIA 1 146-19, 1602.1, 2, 1670, 1674 | *Solanum tuberosum* | Spain     |
| IVIA 2297 4T2a               | Soil            | Spain     |
| EURS 48                      | *S. lycopersicum* | United Kingdom |
| EURS 67                      | *S. tuberosum*   | Belgium   |
| EURS 71                      | *S. tuberosum*   | France    |
| SMT 46                       | *S. lycopersicum* | Peru      |
| 203 Mexico                   | *S. lycopersicum* | Mexico    |

1IVIA: Valencian Institute for Agricultural Research (Spain), 2EURS: European Project CE-FAIR-PL97-3632, 3SMT: European Project CE-SMT4-CT97-2179.

2.2. Phages and amplification conditions

In the different assays, three specific *R. solanacearum* phages, vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2, isolated and purified from environmental water from different regions of Spain [28] were used. The three phages were separately kept with their bacterial host *R. solanacearum* strain CFBP 4944 in cryo-preserved stocks at -25°C in 25% (v/v) glycerol.

For the amplification of the phages, overnight cultures of *R. solanacearum* strain CFBP 4944 were performed and the optical density (OD) at 600 nm was adjusted to a value of 0.5. Then aliquots from the bacterial cultures were separately inoculated with each of the three phages and were incubated overnight at 28°C with shaking (120 rpm). After the incubation period, phage lysates were obtained for each of the phages, which were kept at 4°C until subsequent assays.

Before using the phages, each phage lysate was centrifuged at 10,000 rpm for 10 min, the supernatant was filtered through a sterile 0.22 μm pore diameter filter, and the titration of the supernatant was performed by the double layer agar method. For this, serial ten-fold dilutions were made in SM buffer (50 mM TrisHCl pH 7.5, 100 mM NaCl, 10 mM MgSO4 and 0.01% gelatin) [35]. Volumes of 100 μl of the dilutions were mixed with volumes of 200 μl of a suspension of the strain CFBP 4944 (OD600nm 0.5), and these mixtures were inoculated into CPG top agar (or soft CPG) tubes that were poured onto solid CPG medium. The plates were incubated at 28°C for 24-48 h, and then the plaques formed were counted to calculate the titre of the phage as plaque-forming units (PFU)/ml.

2.3. Stability assays of one single phage lyophilized with various cryoprotectants

Suspensions of phage vRsoP-WF2 intended for lyophilization were initially prepared with three cryoprotective agents: 50% glycerol [36], and 0.1 M and 0.5 M saccharose and trehalose [22, 37].

The suspensions of vRsoP-WF2 in the different cryoprotectants were distributed in sterile glass vials for lyophilization (200 μl per vial) and frozen for 1 h at -80°C while the lyophilizer was tempered at -51.5°C. The vials with the frozen viral suspensions were introduced in a lyophilizer (Virtis) operating at a vacuum pressure of 15.9 mTorr for 18 h [21]. Subsequently, the lyophiles were then sealed with a vacuum pressure of 1.7 mTorr, and stored at 4°C until use.

The viability of phage vRsoP-WF2 after lyophilization with different cryoprotectants was calculated according to the following formula:

\[
\text{Viability} = \frac{\text{Initial V}}{\text{Final V}} \times 100
\]

with “Initial V” being the number of PFU/ml before lyophilization, and “Final V” the number of PFU/ml after lyophilization and resuspension of the phages. Titration was performed in triplicate after rehydration of the viral particles in the same initial volume of 200 μl of sterile distilled water by the double layer agar method as abovementioned.
2.4. Stability assays of one single phage lyophilized or conserved at 4°C over time

To compare the stability of the phage vRsoP-WF2 over time after lyophilization with respect to conservation at 4°C, suspensions of the phage were simultaneously prepared either with saccharose and trehalose 0.5 M for lyophilization, or in buffer SM and the liquid culture medium LB for conservation at 4°C. Stability was monitored during the storage period of vRsoP-WF2 lyophiles and non-lyophilized suspensions after 0, 15, 30 and 60 days. Countings were performed as indicated above.

2.5. Stability assays of the three phages lyophilized with one cryoprotectant

To proceed with the lyophilization of phages vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2, 0.5 M trehalose was selected as the cryoprotective agent. Thus, 1/100 dilutions of the phage stocks were performed in 0.5 M trehalose to obtain a concentration of 10⁸ PFU/ml, and 200 µl of each of the viral suspensions were transferred to sterile glass vials. Lyophilization procedure was as abovementioned. Then, viability and stability of the three phages at different times were determined. Phage lyophiles were resuspended in 200 µl of sterile distilled water and titrations were performed by the double layer agar method. Phage viability was determined by calculating the difference between the titres (PFU/ml) of each of the lyophilized phages at each time “Final V”, and those of the non-lyophilized phages used as controls “Initial V”.

At the same time that the viral suspensions were disposed for lyophilization, additional aliquots were prepared for the three phages in LB liquid medium and in SM buffer, which were kept at 4°C to compare viability and stability under different conservation methods.

Titrations were performed for the three phages, either lyophilized or kept at 4°C in LB medium and in SM buffer at 0, 1, 2, 5, 8 and 12 weeks of storage in these conditions. All the assays were performed in triplicate.

2.6. Effect of lyophilization on the lytic activity of the three phages

To determine the influence of lyophilization on the activity of the R. solanacearum phages, qualitative assays of bacterial lysis were performed with a collection of phage-sensitive R. solanacearum strains listed in Table 1. Bacterial suspensions were prepared in Phosphate Buffered Saline (PBS) 10 mM pH 7.2 for each of the different strains, which were spread onto the surface of CPG plates. Then, drops of 5 µl of the viral suspensions of non-lyophilized phages (controls) and phages lyophilized after 1 and 4 weeks were inoculated onto the surface of the same plates. The assays were performed in triplicate for the three phages.

2.7. Biocontrol assays in planta

2.7.1. Susceptible host plants and growing conditions

For the biocontrol assays, tomato plants (Solanum lycopersicum L.) variety Roma, sensitive to R. solanacearum [13, 38, 39] were used.

Seeds were sown using vermiculite as a substrate, and cultivated for 30 days before the biocontrol assays. Since vermiculite requires the administration of nutrients for the correct plant development, they were watered with Hoagland’s nutrient solution [40] throughout the experimental period. The plants were kept during the assays in a plant growth chamber (Panasonic) at 26°C, with 12 h light and 8 h darkness for 15 days, under conditions of biological containment.

2.7.2. Biocontrol assays with the three phages

Inoculations were performed with R. solanacearum and/or the mixture of the three phages either non-lyophilized or lyophilized.

The strain CFBP 4944 was grown in LB liquid medium for 16-18 h and the suspensions adjusted to OD₆₀₀ₙₐₜ = 0.1 (approximately 10⁸ CFU/ml), then serial ten-fold dilutions were made to a final concentration of 10⁵ CFU/ml for plant inoculation. Regarding phage suspensions, the concentration of each one was adjusted to 10⁸ PFU/ml, and for mixtures,
phage suspensions were combined at a final concentration of 10⁸ PFU/ml. For the in planta inoculations, suspensions were prepared with mixtures of the bacteria and the three phages at a final bacteria:phage concentration of 10⁵ CFU/ml:10⁸ PFU/ml. Positive and negative controls were included, consisting of R. solanacearum suspensions at the same final concentration, and PBS or mixtures of the three phages either lyophilized or non-lyophilized, respectively. The assays were carried out in sets of 10 tomato plants per treatment, in triplicate.

For inoculations, a bevel wound was made in the stem of each plant with a sterile scalpel, between the cotyledons and the first petiole, and volumes of 5 µl of each mixture, suspension or PBS were applied. Inoculated plants were incubated in a growth chamber for 15 days, and the appearance of bacterial wilt symptoms was daily monitored.

From symptomatic plants, R. solanacearum reisolation was performed to confirm the presence of the pathogen. Thus, stem samples above the inoculation point were taken, superficially disinfected, and cut up in PBS buffer to obtain the plant extracts. Direct platings were made on SMSA and incubated at 28°C for 48-72 h. Colonies with R. solanacearum morphology were purified and PCR-identified according to [41].

Plants were also analysed for phage detection. The obtained plant extracts were filtered through 0.22 µm pore membranes and mixed with R. solanacearum suspensions adjusted to OD₆₀₀nm = 0.5, then the mixtures were added to melted CPG top agar, which was poured onto CPG plates and incubated at 28°C for 24-48 h.

2.8. Statistic analysis

Data (PFU/ml) from the viability and stability assays of the phages either lyophilized or preserved in LB medium or SM buffer at 4°C, which were performed in triplicate, were processed by an analysis of variance ANOVA with the GraphPad Prism 6 program after their logarithmic transformation. P values ≤ 0.05 were considered statistically significant.

3. Results

3.1. Viability and stability of phage vRsoP-WF2

3.1.1. Phage viability after lyophilization

To investigate the effect of the lyophilization process on the survival and viability of the phage vRsoP-WF2, three cryoprotectants were initially tested: 50% glycerol, 0.1 M and 0.5 M saccharose, and 0.1 M and 0.5 M trehalose, the titre of the phage being determined before and after lyophilization. The results obtained in this initial test are shown in Figure 1. Phage vRsoP-WF2 did not survive the lyophilization process when 50% glycerol was used as a cryoprotective agent or, at least, survival fell below the detection limit (not shown). On the contrary, saccharose and trehalose made it possible to recover viable viral particles in high numbers at the two concentrations tested, these being higher in the case of 0.5 M for both disaccharides. Thus, vRsoP-WF2 countings after lyophilization were from 3.50 x 10⁷ PFU/ml in the control to 3.01 x 10⁶ and 4.65 x 10⁶ PFU/ml in 0.1 M and 0.5 M trehalose, and to 3.26 x 10⁶ and 7.24 x 10⁶ PFU/ml in 0.1 M and 0.5 M saccharose, respectively.
Based on these data, saccharose and trehalose at 0.5 M were selected to repeat the test. The results confirmed the favourable vRsoP-WF2 survival after lyophilization with these two cryoprotectants. Phage countings were from $2.35 \times 10^7$ PFU/ml in the control to $5.90 \times 10^6$ PFU/ml and $1.74 \times 10^6$ PFU/ml in the lyophiles prepared in 0.5 M trehalose and 0.5 M saccharose, respectively. Therefore, apparently 0.5 M trehalose worked better than 0.5 M saccharose as a cryoprotectant, since the reduction in viability of phage vRsoP-WF2 was less than a logarithmic order.

The appearance of the lyophiles prepared in 0.5 M saccharose was less compact and more crystalline than in 0.5 M trehalose, which were more compact and with a gritty aspect.

3.1.2. Stability of phage vRsoP-WF2

Lyophiles of vRsoP-WF2 were kept at 4°C and their viability assayed over time to evaluate their conservation in the medium-long term. Additionally, control suspensions of the phage prepared from the same stock used for the lyophiles in liquid LB and in SM buffer and kept at 4°C without lyophilization were included.

The results showed maintenance in the countings of the lyophilized phage vRsoP-WF2 throughout the experimental period of 60 days (Figure 2), after an initial decrease of around one log unit, and a subsequent slight decrease at around 30 days. Stability was greater in 0.5 M trehalose than in 0.5 M saccharose, with a concentration of viable viral particles of about $10^6$ PFU/ml and $10^5$ PFU/ml respectively at the end of the experimental period. Therefore, trehalose as a cryoprotectant in lyophilization allowed not only a greater survival of the phage vRsoP-WF2 lyophiles but, also a greater stability of the phage particles, at least in the first 2 months post-lyophilization.
Figure 2. Stability of phage vRsoP-WF2 lyophilized with trehalose or saccharose at 0.5 M and stored at 4°C over time. Titration was carried out by the double layer agar method on CPG plates. Data are the means for assays performed at least in triplicate, and error bars indicate variation as the standard deviation. Asterisks above bars denote statistically significant differences between the two signaled columns: * p < 0.05, ** p < 0.01, *** p < 0.001.

When the stability of the lyophilized vRsoP-WF2 phage particles kept at 4°C were compared with that of the phage suspensions of the controls kept in LB medium and in SM buffer at 4°C, the results showed greater viability and stability of the viral particles throughout the same experimental period (Figure 3).

Figure 3. Stability of phage vRsoP-WF2 preserved in LB medium and in SM buffer at 4°C over time.Titration was carried out by the double layer agar method on CPG plates. Data are the means for assays performed at least in triplicate, and error bars indicate variation as the standard deviation.

The viability of the vRsoP-WF2 phage suspensions did not undergo initial decreases as observed when the viral particles were lyophilized. In fact, the titre of these suspensions did not drop below the initial logarithmic order (10^7 PFU/ml) throughout the 60 days. However, there were small decreases in the concentration of viable viral particles over time. Comparing both media, despite the high stability of phage vRsoP-WF2 in LB medium and SM buffer, this was slightly higher in SM. The countings of vRsoP-WF2 stabilized in LB medium after 15 days while in SM buffer a smooth but more progressive decrease was observed.
3.2. Viability and stability of phages vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2

3.2.1. Phage viability after lyophilization

To verify the viability and survival of the three phages over time after lyophilization, the viral suspensions were titrated before and after this process with 0.5 M trehalose as a cryoprotective agent.

The viral suspensions were titrated before their lyophilization, as controls, and then the lyophiles obtained at time zero and periodically for 12 weeks to determine the viability of the phages. As it can be seen in Figure 4, the three phages maintained their viability after lyophilization with trehalose but, this was lower than that of the controls. In the case of phage vRsoP-WF2, the titre initially decreased one logarithmic order, from $10^8$ to $10^7$ PFU/ml, then maintained over time. In phage vRsoP-WM2, the decrease was more pronounced, going from $10^8$ to $10^6$ and $10^5$ PFU/ml, and staying around the latter value. In the case of vRsoP-WR2, there was a reduction of two logarithmic units, from $10^8$ to $10^6$ PFU/ml, remaining at this level throughout the experimental period. Thus, the survival of phage vRsoP-WM2 was lower than that of the others, under the conditions tested.

![Figure 4](image)

**Figure 4.** Stability of phages vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2 lyophilized with trehalose at 0.5 M and stored at 4°C over time. Titration was carried out by the double layer agar method on CPG plates. Data are the means for assays performed at least in triplicate, and error bars indicate variation as the standard deviation. Asterisks above bars denote statistically significant differences between the two signaled columns: * p < 0.05, ** p < 0.01, *** p < 0.001.

3.2.2. Stability of phages vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2

To compare the stability of the lyophiles of the different phages with that of other conservation methods, at the same time that the lyophiles were prepared, phage suspensions from the same stocks were prepared in LB medium and SM buffer, and kept at 4°C. Thus, each time the lyophilized phages were titrated, these phage suspensions at 4°C were titrated in parallel. The results showed a greater stability and viability of the viral particles of these suspensions at 4°C throughout the same experimental period (Figure 5). The viability of phages vRsoP-WF2 and vRsoP-WR2 was not reduced, contrarily to their viability as lyophiles for the same period (Figure 4), in fact, their titres maintained values around $10^8$ PFU/ml throughout the 12 weeks. However, the viability of the phage vRsoP-WM2 decreased one logarithmic order, from $10^8$ to $10^7$ PFU/ml, although this reduction was lower than that observed when lyophilized (Figure 4).
3.3. Effect of lyophilization on the lytic activity of the phages vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2 against different bacterial strains

To determine the maintenance of the lytic activity of the phages after their lyophilization at different times, *R. solanacearum* strains of various sources and geographical origins were used (Table 1) in assays with the non-lyophilized and the lyophilized phages.

The strain CFBP 4944 was used as a control, since it has been shown that it is lysed by the three phages, both the non-lyophilized and the lyophilized (Figures 4 and 5). It was observed that all the tested strains were lysed by the three phages before and after having been subjected to the lyophilization process. Of these, the phage vRsoP-WM2 was comparatively the most affected by this process, since it showed the lowest lytic activity against most of the tested strains, while vRsoP-WF2 and vRsoP-WR2, both non-lyophilized and lyophilized, maintained their activity against the different strains of *R. solanacearum*.

3.4. Bacterial wilt biocontrol assays in susceptible plants inoculated with *R. solanacearum* and mixtures of vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2

To determine the potential of the lyophilized phages to control the bacterial wilt disease, biocontrol assays were carried out with tomato plants inoculated on the stem simultaneously with the pathogen and the three phages lyophilized for 4 weeks comparatively to a mixture of the non-lyophilized. Strain CFBP 4944 was used in the assays, and the respective controls were included. Results were obtained as the percentage of plants with wilting symptoms with respect to the total number of inoculated plants.

Plants treated with bacteria and phages showed a reduction in wilting symptoms with respect to plants inoculated only with bacteria (Figure 6). The treated plants did not reach 100% wilting in any of the assays, contrarily to the positive controls, which completely wilted within the second week post-inoculation. Wilting values for treated plants were around 33.3%, either those inoculated with the pathogen and a mixture of phages lyophilized for 4 weeks or a mixture of the non-lyophilized. Regarding the rate of appearance of bacterial wilt symptoms, an initial variation could be observed between the phages lyophilized for 4 weeks and the non-lyophilized (Figure 6), although at the end of the assay results were comparable, and it could be argued that the phages can prevent and control bacterial wilt almost equally. Negative control plants, inoculated with PBS or the
non-lyophilized phages or the phages lyophilized for 4 weeks, remained asymptomatic with 0% wilting throughout the experimental period.

**Figure 6.** Bacterial wilt biocontrol assays in planta inoculated with *Ralstonia solanacearum* strain CFBP 4944 and a mixture of the phages vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2 either non-lyophilized or lyophilized after 4 weeks. Mixtures were co-inoculated at final concentrations bacteria:phages of $10^5$ CFU/ml:$10^8$ PFU/ml. Positive controls (Rsol) were at $10^5$ CFU/ml. Negative controls (mixtures of phages) were at $10^8$ PFU/ml. Additional negative controls were performed with PBS. Plants inoculated with negative controls, either mixtures of phages or PBS did not develop wilting symptoms (not shown). Data are the means for assays performed in triplicate, and error bars indicate variation as the standard deviation. Asterisks above bars denote statistically significant differences between the two signaled columns: * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$.

Tomato plants with wilting symptoms were processed, and the pathogen was reisolated on SMSA. The selected colonies with typical reddish coloration and morphology were PCR identified as *R. solanacearum*.

The detection of phages after 15 days of inoculation was possible in the control plants inoculated with non-lyophilized phages, where plaques could be observed on the plates. No plaque was detected from plants inoculated with the mixture of lyophilized phages, either with the mixture of bacteria and non-lyophilized phages, or the mixture of bacteria and phages lyophilized after 4 weeks, in the selection of plants analysed.

4. Discussion

The growing interest in phages as biocontrol agents for bacterial diseases of plants makes it necessary to develop methods of phage preservation that are reliable in the medium and long term to allow their commercialization. One of the methods considered most effective in terms of preserving the characteristics of the microorganisms and their ease of storage and transport is lyophilization, since it could allow both the maintenance of the viral particles and their readily distribution [22, 26].

Since a biocontrol method of bacterial wilt based on the use of the phages vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2 due to their lytic activity against *R. solanacearum* was developed and patented [28-31], the viability and stability of the three phages after subjecting them to an aggressive freeze-drying preservation process as the lyophilization became a necessary objective in view of their commercialization.

Lyophilization implies a strong stress for the viral particles, which can lead to a great loss of viability unless suitable cryoprotective or lyoprotective agents are used. Based on the literature [22, 25, 36, 42], an initial assay was performed with the phage vRsoP-WF2 testing three cryoprotectants. One of them was 50% glycerol, which did not lyophilize coinciding with previous studies and the viability of vRsoP-WF2 dropped below the detection levels. This fact demonstrated that without the presence of an adequate cryoprotective agent, lyophilization is a very aggressive process able of destroying microorganisms. This has been related to the formation of ice crystals during freezing, in addition to
the increase in osmolarity resulting from an increase in internal solute concentration, and to the denaturation of molecules due to dehydration [22, 43]. On the contrary, when saccharose and trehalose, disaccharides described as good cryoprotectants in other phages [22-25, 37, 42, 44, 45] were used, the results revealed that both allowed to maintain high concentrations of viable viral particles at the two concentrations tested (0.1 M and 0.5 M), being higher at the highest concentration and in the presence of trehalose. In this case, the cryoprotection mechanisms provided by these two disaccharides have been described as involving vitrification, i.e. the conversion of a material to solid devoid of crystalline structures and, on the other hand, the formation of a matrix that prevents the aggregation of microorganisms and mechanical stress by ice crystals [22]. In addition, it has a low hygroscopicity due to the lack of internal hydrogen bonds, which allows for greater plasticity in the formation of these bonds on proteins [22, 46] and, in this way, it can prevent the inactivation of phages during the lyophilization process [47].

Based on these results, the assay was repeated only with 0.5 M saccharose and 0.5 M trehalose, confirming the results of the first assay and thus trehalose as the best cryoprotective agent for vRsoP-WF2, coinciding with previous studies on other phages [22].

On the other hand, the stability of vRsoP-WF2 was monitored over time in order to evaluate medium to long term preservation. The results showed a slightly higher stability in the presence of trehalose than in saccharose 60 days after lyophilization. These differences were not very pronounced but, they could be marked at longer times, so it would be necessary to continue the study in the longer term.

When these results were compared with those obtained with controls maintained in LB medium and in SM buffer at 4°C, it was observed that the stability was greater for these controls within the experimental period, and pronounced losses of viability as those obtained with the lyophilization process did not occur. However, lyophilization has several advantages, such as the smaller volume of sample to be preserved as opposed to the larger volumes of microbial suspensions in liquid media; greater ease of transport and shipping; lower maintenance costs because it does not require subzero temperatures; and a lower risk of contamination, since lyophilized viral particles are dehydrated and in vacuum conditions, unlike stocks of viral suspensions in liquid media.

Based on the results abovementioned, the effect of lyophilization on the viability and stability of vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2 was addressed using trehalose 0.5 M. The viability of the three phages after lyophilization tended to slight decreases with respect to the non-lyophilized controls. Thus, phages vRsoP-WF2 and vRsoP-WR2 presented reduced viability values, around 1 and 1.5 logarithmic units, and the phage vRsoP-WM2 reductions of around 2 and 3 logarithmic units, with respect to the controls and under the conditions tested. Among the three phages, vRsoP-WM2 seemed to be more affected by the lyophilization than vRsoP-WF2 and vRsoP-WR2, possibly because it is more sensitive to the desiccation and/or freezing treatments suffered during the process, or maybe trehalose at the concentration tested is not the best cryoprotectant for this phage, hindering adequate preservation and survival.

In this sense, also after lyophilization, the titres of the three phages decreased in comparison with their respective controls, even if a cryoprotectant was added [36, 48]. In contrast, the titres of the phages that were directly conserved in LB medium and in SM buffer at 4°C did not decrease, except for vRsoP-WM2. Therefore, this phage exhibits greater sensitivity to conservation at low temperature than the other two phages, similarly to what happened in relation to conservation by lyophilization. Overall, even with the exception of vRsoP-WM2, the phages preserved at 4°C have remained more stable than those lyophilized, at least for a period of three months. In fact, phages can be kept stable for long periods under low temperatures [21, 44, 46, 49]. However, lyophilization has other valuable advantages already mentioned.

The biocontrol potential of the phages after lyophilization in susceptible host plants was evaluated. To verify the efficacy of a cocktail of the three R. solanacearum phages after lyophilization, biocontrol assays were carried out in tomato plants co-inoculated in the stem with the pathogen and a cocktail of phages either lyophilized or not. In the assays,
biocontrol percentages were higher than 65% in the plants treated with the non-lyophilized phage cocktail, whereas the biocontrol percentages with the cocktail of lyophilized phages at 4 weeks were higher than 50%. In contrast, the plants inoculated only with R. solanacearum presented wilting symptoms earlier than those treated with phages, with 100% wilting in less than 2 weeks. Plants treated only with phages did not develop any disease symptoms, indicating their safety and suitability as biocontrol agents. Up until now, there are no other references on the effect of a conservation method as the lyophilization on the biocontrol efficacy in planta of R. solanacearum phages.

In summary, the work carried out has made possible to increase the knowledge on the viability of lyophilized and non-lyophilized phages vRsoP-WF2, vRsoP-WM2 and vRsoP-WR2, and on their interaction with R. solanacearum. It has been found that they remain fairly stable when lyophilized with trehalose 0.5 M, at least during a period of 90 days. Further, the three phages retain their lytic activity against R. solanacearum and biocontrol efficacy in susceptible tomato plants, providing determining of their preventive potential or therapeutic action on the bacterial wilt disease.

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