Fast and reliable procedure developed to generate soft rot 
Pectobacteriaceae (Pectobacterium spp. and Dickeya spp.)
Tn5 mutants resistant to bacteriophage infection

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Abstract A simple and fast procedure has been developed to generate soft rot Pectobacteriaceae (SRP: Pectobacterium spp. and Dickeya spp.) Tn5 mutants in genes encoding receptors used by bacteriophages to interact with their hosts, for the follow-up studies. The procedure is inexpensive and does not require any specialized tools and/or dedicated technical support. The neomycin-resistant SRP Tn5 mutants are generated via conjugation with a transposon donor Escherichia coli ST18 strain (requiring 5-aminolevulinic acid (5-ALA) to survive) carrying pFAJ1819-mini-Tn5-neoR. The conjugation is done on solid medium supplemented with 5-ALA. After conjugation bacterial cells are collected, suspended in liquid bacterial medium, added to the suspension containing lytic bacteriophages and incubated for the additional 30 min with shaking (120 rpm). During this stage, the transposon recipients (Pectobacterium spp. and/or Dickeya spp. Tn5 mutants), susceptible to bacteriophage infection are lysed. Likewise, due to the lack of 5-ALA in the growth medium, E. coli ST18 (transposon donor) cells die at this stage. Finally, after incubation, the bacterial mutants with the Tn5 insertions, resistant to phage infection are selected on solid growth medium supplemented with neomycin. The Tn5 insertion sites are sequenced to acquire knowledge about the Tn5-disturbed genes and their putative function in phage-host interactions. The proposed assay allows generation of a number of immediately-available Tn5 mutants expressing phage-resistant phenotypes in a short time (ca. 48 h) that can be later characterized for various other phenotypic features. In this study, as a proof-of-concept, this method has been used to generate Dickeya solani IPO2222 Tn5 mutants resistant to infection caused by the lytic bacteriophage φD5.

Keywords Bacterial viruses · Survival · Phage · Resistance · Transposon mutagenesis · Phage receptors

Bacteriophages (phages) are viruses able to infect and kill bacteria (Abedon 2009). Due to their obligatory parasitic nature, phages are completely dependent on their bacterial hosts to survive and to propagate in the environment (Labrie et al. 2010). Survival and ecological success of bacteriophages is therefore solely linked to their ability to infect the host cell and, once infection has happened, to complete their life cycle within the host. All successful viral infections begin with host recognition and adsorption: an irreversible binding (attachment) of phage tail proteins to the specific components on the surface of bacterial cells named receptors (Bertozzi Silva et al. 2016).

The presence of a specific receptor on the host as well as its three-dimensional structure determine whether a...
particular bacterium is susceptible to phage infection or remains resistant (Lindberg 1973). Although a number of studies has targeted analyses of receptors involved in interaction of model bacterial viruses (e.g. phages λ and T4) with the model bacteria (viz. *Escherichia coli*), still relatively little is known about receptors used by environmental bacteriophages that infect hosts other than *E. coli* (Rakhuba et al. 2010). Specifically, the knowledge about receptors used by environmental bacteriophages to infect plant pathogenic bacteria is scarce (Bertozzi Silva et al. 2016). A fundamental understanding of the molecular mechanisms governing viral adsorption to the host cell is crucial to assess the role of bacteriophages in environment in adaptation and evolution of their hosts (Samson et al. 2013). Likewise, the knowledge about receptors utilized by bacteriophages is of use when designing viral-based control strategies for bacterial pathogens both in food production and in agriculture (Roach and Debarbieux 2017).

The aim of this study was to develop a simple and fast procedure to generate bacteriophage-resistant soft rot *Pectobacteriaceae* (SRP) Tn5 mutants. These bacteria are important necrotrophic pathogens infecting a number of commercially-relevant crops worldwide (Charkowski 2018; Pérombelon 2002). The presence of SRP bacteria in crops is linked to increasing losses in agriculture under different climatic conditions (Charkowski et al. 2020) and consequently SRP have been for a long time recognized among the top-ten most important plant pathogenic bacteria (Mansfield et al. 2012). For many years SRP have been a useful model for studying molecular aspects of the plant-bacterium interactions for both monocotyledonous and dicotyledonous plants (Hugouvieux-Cotte-Pattat et al. 1996; Sepulchre et al. 2007), they were however not extensively used as a model to study bacteriophage-bacterial host interactions (Czajkowski 2016, 2019). Analyses of receptors present on the surface of SRP bacteria that are used by bacteriophages for interaction/infection may broaden our understanding of *Pectobacterium* spp. and *Dickeya* spp. adaptation in environment as well as their evolution in the presence of bacterial viruses.

To develop and validate a proposed procedure, for the proof-of-concept experiments, the SRP pathogen emerging in Europe, *D. solani* type strain IPO2222 (van der Wolf et al. 2014), together with its corresponding lytic bacteriophage ϕD5, characterized in detail in the previous studies (Czajkowski et al. 2013, 2014) were chosen. Unless specifically noted, well-established molecular biology and microbiological methods were used for all described experiments (Sambrook et al. 1989).

For this study, the bacteriophage ϕD5 was propagated on their wild type bacterial host (*D. solani* IPO2222) and titer as described earlier (Czajkowski et al. 2013). The adjusted stock concentration of ϕD5 phage particles was $10^9$ plaque forming units (pfu) mL$^{-1}$ in Luria-Bertani broth (LB broth-Miller, Sigma-Aldrich). The experimental methodology to generate bacterial mutants was based on a random transposon mutagenesis of *D. solani* genomes with a mini-Tn5 transposon. To mutate bacterial genomes, conjugation of *D. solani* IPO2222 (transposon recipient) with *Escherichia coli* S18 (transposon donor, requiring 5-aminolevulinic acid (5-ALA) to survive) (Thoma and Schobert 2009) carrying pFAJ1819 (obtained from The Belgian Coordinated Collections of Microorganisms - BCCM, Brussels, Belgium) (Xi et al. 1999) suicide vector with mini-Tn5-neo$^R$ was performed as previously described (Czajkowski et al. 2017). Briefly, *E. coli* strain ST18 carrying pFAJ1819-Tn5-neo$^R$ was

![Fig. 1 The scheme of the procedure used to generate phage-resistant mutants of *Dickeya solani* strain IPO2222](image-url)
grown in Luria-Bertani broth (LB broth-Miller, Sigma-Aldrich) supplemented with 50 µg mL⁻¹ of 5-aminolevulinic acid (5-ALA, Sigma-Aldrich) for 16 h at 37 °C with shaking (120 rpm) and *D. solani* strain IPO2222 was grown under the same growth conditions and for the same time but at 28 °C. For the conjugation assay, the equal volumes of the donor (2 ml) and recipient (2 ml) strains, collected after incubation, were individually centrifuged for 3 min. at 8000 x g, washed twice with sterile Ringer’s buffer (Merck) and after a second wash, resuspended each in the 500 µl of sterile Ringer’s buffer. The donor (0.5 ml) and recipient strain (0.5 ml) cultures were mixed together with the equal proportions (1:1), vortexed vigorously for 1 min. and centrifuged for 3 min. at 8000 x g to sediment the cells. The joint bacterial pellet was resuspended in 15 µl of sterile Ringer’s buffer and the resulting bacterial suspension (ca. 25 µl) was dropped onto the Luria-Bertani agar (LB agar-Miller, Sigma-Aldrich) plate supplemented with 50 µg mL⁻¹ of 5-ALA. Such an inoculated plate was incubated for 6 h at 37 °C for conjugation between donor and recipient to occur. After conjugation, bacterial cells (ca. 10⁸ – 10⁹ colony forming units (cfu) mL⁻¹) were collected from the agar plate using a sterile inoculation loop and resuspended in 500 µl of LB broth following addition of a 500 µl volume of ϕD5 suspension in LB broth containing 10⁹ plaque forming units (pfu) mL⁻¹. Such a bacterial suspension (1 ml) was incubated for 30 min. at 28 °C with shaking (120 rpm) to allow phages to infect and to lyse susceptible *D. solani* Tn5 mutant cells. After this time, 100 µl volumes were separately plated on LB agar plates without 5-ALA and supplemented with 50 µg mL⁻¹ of neomycin (Sigma-Aldrich) following incubation for 24–48 h at 28 °C to select *D. solani* mutants containing Tn5 insertions, resistant to ϕD5 infection. The scheme of the procedure is shown on Fig. 1.

The obtained phage resistant Tn5 mutants selected in this procedure were screened using a plaque formation assay and host challenge assay (Clokie and Kropinski 2009) to confirm the phage resistant phenotype of the each
individual Tn5 mutant analyzed. Additionally, the Tn5 mutants were selected for sequencing the transposon insertion sites as previously described (Lisicka et al. 2018).

In the preliminary assay using the proposed procedure, the 15 D. solani Tn5 mutants, resistant to bacteriophage ϕD5 were generated and analyzed in time shorter than 48 h. All the D. solani mutants were confirmed to be resistant to infections caused by ϕD5 bacteriophage. Example results are shown in Fig. 2; Table 1. The Tn5 insertions were found in genes coding for proteins associated with bacterial cell membrane/cell wall, for example: in gene coding for a PTS mannose transporter subunit IID (mutant 9ST), methionine ABC transporter permease (mutant 11ST), sulfate ABC transporter ATP-binding protein (mutant 12ST), spermidine/putrescine ABC transporter ATP-binding protein (24ST) as well as in the couple of cases the transposons were inserted into the genes coding for hypothetical proteins with unknown functions (e.g. in mutant 15ST).

In conclusion, the proposed procedure allows an immediate (time less than 48 h) generation of SRP Tn5 mutants resistant to bacteriophage infections for the follow-up studies. These studies may include various phenotypic analyses of phage resistant bacterial strains, investigations of phage-bacteria ecological interactions as well as comparative studies targeting adaptation and fitness of bacterial strains resistant/susceptible to phage infections in environment. The major advantage of the proposed procedure is that the assay can be completed in a short time and that it does not require the screening of a large collection of Tn5 mutants to find bacteriophage resistant ones. Likewise, the proposed procedure does not require any expensive laboratory tools and consumables and therefore it may be performed in virtually every microbiological laboratory possessing standard microbiological equipment.

It is postulated that due to the universal nature of selection of bacteriophage-resistant Tn5 mutants, this procedure can be easily adapted to generate mutants of other bacterial species and resistant to other lytic bacteriophages of interest.

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Compliance with ethical standards The results presented in this manuscript did not involve any protected and/or endangered species, field studies, human participants, specimens or tissue samples, or vertebrate animals, embryos or tissues.

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