An improved conjugation method for Pseudomonas syringae

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An improved conjugation method for Pseudomonas syringae

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Abstract

- 20 In order to achieve saturating transposon mutagenesis of the genome of plant
- 21 pathogenic strains of *Pseudomonas syringae* we needed to improve plasmid
- 22 conjugation frequency. Manipulation of the growth stage of donor and recipient cells
- 23 allowed the required increase in frequency and facilitated conjugation of otherwise
- 24 recalcitrant strains.

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Strains of Pseudomonas syringae (P.s.) cause a wide range of economically 28 important plant diseases worldwide. A number of P. s. pathovars from distinct 29 phylogroups cause bacterial canker of cherry (Prunus avium) including P. syringae 30 pv. morsprunorum (Psm) races 1 and 2 and P. syringae pv. syringae (Pss) (Hulin et 31 32 al., 2020). Bioinformatic analyses have identified shared sets of genes putatively involved in the pathogenicity of Psm and Pss on cherry (Hulin et al., 2018). As a 33 34 functional screen for genes in Psm and Pss regulating colonisation of, and persistence in, woody tissue we initiated an unbiased approach using saturating 35 transposon mutagenesis. We proposed to use the method of TnSeq (transposon 36 mutagenesis with next-generation sequencing) (Wetmore et al., 2015), which 37 requires a high density of transposon insertion into the genome. 38 To achieve saturating mutagenesis a high conjugation frequency needs to be 39 achieved for the transfer of the plasmid containing the transposon into the recipient 40 Pseudomonas strain. Using routine protocols for the cherry pathogens we grew 41 overnight cultures of Pss 9644, Psm R2 MH001 (formally R2 leaf) and R1 5244 42 (Hulin et al., 2018) and *E. coli* APA752 containing the *mariner* plasmid pKMW3 43 (Wetmore et al., 2015), cultured in LB broth for 18 h at 25°C or 37°C for E. coli 44 respectively. One ml of each overnight culture was diluted 1 in 10 and grown for 6 h 45 to reach log phase (0.8 OD₆₀₀) (Sup Fig. 1). Equal cell numbers (500 µl each of 0.8 46 OD₆₀₀) of *Pseudomonas* strains and *E. coli* were combined and allowed to conjugate 47 on 0.45µm nitrocellulose filters (Millipore) placed on LB agar plates supplemented 48 with 0.0625 mg/mL diaminopimelate (in sterile distilled water) at 30°C. After 6 h cells 49 were scraped off the filters, serially diluted and plated onto KB agar with 25µg/mL 50

kanamycin (Kan) and either 100 μg/ml nitrofurantoin (Nf, in DMSO) for *Pss* 9644 and *Psm* R2 MH001 or 100 μg/ml rifampicin (Rif, in methanol) for *Psm* R1 5244. Using this standard protocol we achieved a maximum of 2.6x10² transconjugants (CFU) per ml of conjugation mixture (Fig.1) which was insufficient to allow adequate saturation of the genome required for TnSeq screens (Wetmore *et al.* 2015). We therefore repeated the procedure but increased incubation time on the conjugation plates to 24h which increased the number of transconjugants 100-fold to a maximum of 1.5x10⁴ CFU/ml for *Psm* R2 MH001. However, the conjugation frequency was much lower in *Psm* R1 5244 and *Pss* 9644 rendering it difficult to proceed with mapping the transposon mutant library and *in planta* experiments.

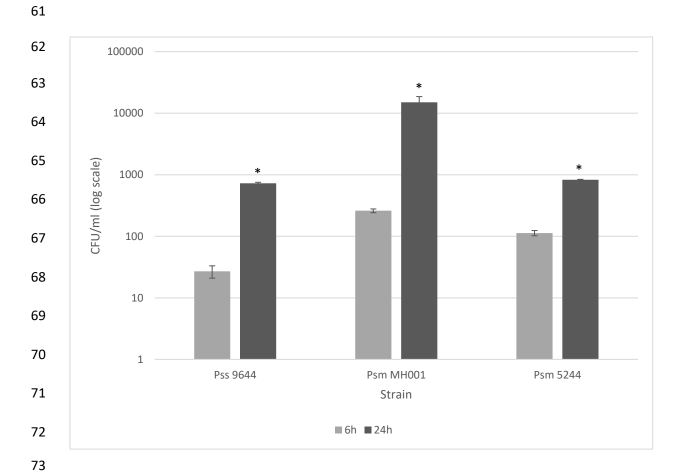


Figure 1. Effect of varying the time for conjugation on the frequency of transfer of the *mariner* plasmid pKMW3 into *Pseudomonas* strains using the Wetmore

et al. (2015) method. Extension of the routine 6h incubation on conjugation plates to 24h led to a statistically significant increase in conjugation frequency in all three strains, indicated by * above the bars, as determined by a Student's t-test (p<0.05). Data show means of three replicates ± standard error of mean (SEM) and are displayed as log₁₀ cfu per ml of conjugation mixture, using strains of *Pseudomonas* syringae pv. morsprunorum R2 (*Psm*); *P. syringae* pv. syringae (*Pss*).

We next examined the effect of changing the growth stage of the donor and recipient. We used a combination of donor and recipient in stationary phase with an OD600 1.5 (18 h) and log phase with an OD600 0.8 (6 h) (Fig. 2), with a 24h conjugation incubation time. The conjugation frequency of pKMW3 into *Pseudomonas* strains using the donor cells in log phase and the recipient cells in stationary phase resulted in a ~1000 fold increase in conjugation frequency to a maximum of 2.7x10⁷ CFU/ml (Fig.2). This new conjugation frequency was considered adequate to make the TnSeq libraries. To confirm that plasmid pKMW3 had transferred to the transconjugant cells, we amplified a section of the transposon using a standard PCR protocol with primers pKMW3F-5'GATGTCCACGAGGTCTCT3', pKMW3R-5'GTCGACCTGCAGCGTAC3' (Wetmore et al., 2015). A region of 100 bp was obtained in ten randomly selected transconjugants (data not shown).



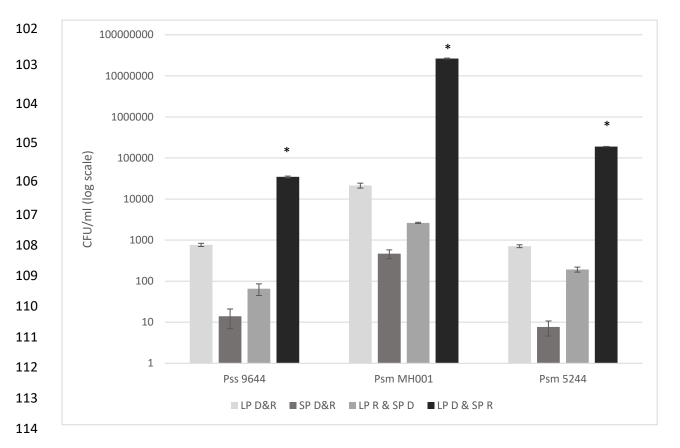


Figure 2. Effect of varying the growth stage of donor (D) and recipient (R) cells on the frequency of conjugation of the *mariner* plasmid pKMW3 into *Pseudomonas* strains. Incubation for log phase and stationary phase multiplication was for 6h (OD600 0.8) and 18h (OD600 1.5) respectively using strains of *Pseudomonas syringae* pv. *morsprunorum* R2 (Psm) and *P. syringae* pv. *syringae* (Pss). Using a combination of log phase (LP) donor cells and stationary phase (SP) recipient cells resulted in a statistically significant increase in conjugation frequency for all three strains as indicated by the *, determined by a within-strain comparison of means by a Student's t-test (p<0.05). Data show means of three replicates ± standard error of mean (SEM) and are displayed as log₁₀ CFU/ml of conjugation mixture.

To expand this study, we tested our revised protocol on other *P. syringae* strains that have previously exhibited low conjugation rates. We used *P. syringae* pv. *phaseolicola* (*Pph*) 1448A (Joardar et al., 2006), a good conjugator as a control and *Pph* 1302A (Taylor et al., 1996), which has been very recalcitrant to plasmid conjugation in the past. We also tested additional recalcitrant strains - *P. syringae* RMA1, a pathogen of *Aquilegia vulgaris* (Hulin et al., 2018) and the cherry pathogen *Pss* 9097 (Hulin et al., 2018) (Fig. 3). The new method allowed transconjugants to be obtained at a reasonable frequency (1302A 4.7x10³ CFU/ml; RMA1 2.1x10³ CFU/ml; 9097 1.7x10² CFU/ml) with strains that had very low or no transconjugants using the original Wetmore et al. (2015) method.

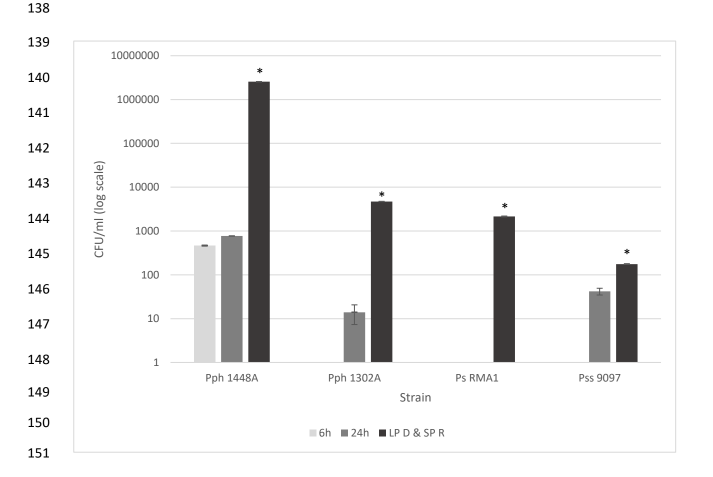


Fig. 3. Effects of varying incubation period on conjugation plates and the growth phase of donor and recipient cells on the frequency of conjugation of the *mariner* plasmid pKMW3 into diverse *Pseudomonas* strains. The combination of log phase donor cells and stationary phase recipient cells resulted in a statistically significant increase in conjugation frequency for all four *Ps* strains; isolates of *P. syringae* pv. *phaseolicola* (*Pph*); *P. syringae* (*Ps*) and *Pseudomonas syringae* pv. *syringae* (*Pss*). *Pph* 1302A, *Pss* 9097 produced no transconjugants at 6h and *Ps* RMA1 none at 6 or 24h. Statistical significance as indicated by the * was determined by a within-strain comparison of means using a Student's t-test (p<0.05), means are given of three replicates ±SEM and are displayed as log₁₀ CFU/ml of conjugation mixture.

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Data underlying this article can be accessed at http://researchdata.uwe.ac.uk/583/

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