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Fluidic resistance control enables high-throughput establishment of mixed-species biofilms

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ABSTRACT
Bacteria often live in communities of mixed species embedded in a self-produced extracellular matrix of polysaccharides, proteins and DNA, termed biofilms. The BioFlux microfluidic flow system is useful for studying biofilm formation in different media under flow. However, analyzing the architecture and maturation of biofilms under flow requires a proper seeding, which can prove difficult when working with bacteria of different sizes, motile bacteria or aiming for a high number of replicates. Here we developed an efficient protocol that exploits viscosity tuning and seeding indicator dyes to improve seeding and allow for high-throughput examination and visualization of consistent mono- and mixed-species biofilm developments under flow.

METHOD SUMMARY
An improved protocol for seeding bacteria in the BioFlux microfluidic flow system was developed using glycerol and Nigrosin to increase the viscosity and stain the media, respectively. This new protocol does not alter biofilm development and allows consistent mixed-species biofilm formation in a copious number of replicates.

KEYWORDS:
biofilm analysis • biofilm device • biofilm formation • biofilm seeding • BioFlux • CLSM • microfluidic device • microfluidic flow systems • mixed species biofilm • shear stress

The BioFlux microfluidic flow system (Fluxion Biosciences) has been developed in order to study biofilm formation in continuous flow over a broad range of controlled fluid shear, approximate natural biofilms known from, for example, medical devices or some human infections, while allowing simultaneous studies of up to 96 replicates. Biofilms established in the BioFlux system can be examined continuously with optical microscopy or confocal laser scanning microscopy (CLSM), enabling monitoring of attachment, maturation and potential dispersal under flow, while requiring minimal volumes of media compared with other continuous flow systems [7]. The small volume applied compared with other systems such as drip flow reactors makes it suitable for screening of biofilm inhibitory agents under flow [9]. Precise seeding is, however, required in order to avoid time waste and running cost of unsuccessful replicates. Seeding is performed by reversing the direction of the fluidic flow and allowing the inoculum to enter the viewing chamber from the outlet well, without contaminating the upstream channel and inlet well (Figure 1), which can prove challenging without hands-on experience. Moreover, recent research focuses on interspecies interactions in multispecies biofilms and the competition, cooperation, synergy and new microniches they promote [10], imposing further challenges on the model systems applied. Difficulties when seeding mixed bacterial communities efficiently and precisely are currently a significant constraint of the BioFlux system, which limits the reproducibility of high-throughput experiments. The swimming speed of bacterial cells affects settling dynamics [11], and motility also enables the cells to settle at unintended positions and inconsistently between channels. In addition, inadvertent residual flow during the attachment period, and different settling rates among bacterial cells of different sizes in the inoculum (both in the well and within the flow channel), cause difficulties when using this system to obtain consistent growth, especially for mixed-species biofilms. We here present an improved protocol for seeding the BioFlux system, designed to circumvent the challenges experienced in analysis of mixed-species cultures. By using glycerol to increase the viscosity of the seeding media and Nigrosin stain to visualize the inoculum, we enhanced throughput and producibility of bacterial seeding in the BioFlux microfluidic flow system.
According to Stokes’ law, settling velocity varies greatly with particle size and is related to fluid viscosity (Eq. 1).

\[ v = \frac{2 (\rho_p - \rho) g R^2}{9 \mu} \]

Where \( v \) is the settling velocity (m/s), \( R \) is the radius of the particle (m), \( \mu \) is the dynamic viscosity (kg/m * s), \( g \) is the gravitational acceleration (m/s²), \( \rho_p \) is the mass density of the particles and \( \rho \) is the mass density of the fluid (kg/m³).

Increasing the medium viscosity with glycerol reduces gravity-induced flow and equalizes the settling time and simultaneously decreases unwanted flow during the attachment period. As bacteria are small and seeding is often performed with a few cells, they can be difficult to observe via microscopy at low magnification. Staining the inoculum provides a simple way for the operator to verify that the inoculum has been introduced in the viewing area quickly and easily, without reverting from low to high magnification microscopy (Figure 1C). This ensures correct bacterial positioning and improves the consistency of seeding, thus also the consistency of biofilm formation, and enables controlled high-throughput biofilm growth in this flow system.

A doubling in viscosity can be achieved at 20% glycerol concentration [12,13], which will lead to a halving of both settling velocity and residual flows due to gravity and surface tension effects. The same pressure driving flow will lead to lower flow speeds in the channel and result in better control of the position of the inoculum front. Longer settling times will lead to superior species mixing during inoculum introduction.

In this study we used fluoresecently tagged strains of Pseudomonas putida, Paenibacillus amylolyticus and a wrinkled variant of Xanthomonas retroflexus in order to demonstrate the improvements provided by this new protocol. These bacteria represent a great model, due to their ability to grow under the same conditions, and exemplify the challenges met during seeding of motile or different-sized bacteria.

P. amylolyticus and X. retroflexus are soil isolates known to engage in a dynamic multispecies biofilm with strong synergistic effects when grown in a static four-species community alongside Microbacterium oxydans and Stenotrophomonas rhizophila [14]. The recently isolated and characterized wrinkled variant of X. retroflexus, with enhanced biofilm formation, has shown augmented mutualistic behaviour when forming dual-species biofilms with P. amylolyticus and also a changed spatial organization compared to that of co-cultures containing the ancestral variant [15]. However, as P. amylolyticus is considerably larger than wrinkled X. retroflexus, correct seeding of this combination of strains and other similar ones can prove challenging (Figure 2A–D). By increasing the media viscosity and adding stain (Nigrosin) to the inoculum, we were able to successfully introduce this co-culture and establish a dual-species biofilm in a consistent high-throughput manner in the BioFlux system (Figure 3). This procedure is directly applicable to future studies of different combinations of bacterial species, enabling studies of mixed-species biofilms under continuous flow.

P. putida and X. retroflexus were used to analyze the improved BioFlux seeding procedure, based on addition of stain and enhanced viscosity, under high-throughput conditions (Figure 4). Despite the fact that the bacterial inoculum was introduced into the center of the viewing chamber, control channels at normal viscosity had in some cases upstream bacterial growth after incubation (Figure 4D–F). This was completely avoided by increasing the viscosity during seeding (Figure 4A–C). Moreover, measurements of the distance from the beginning of the biofilm formation and to the downstream edge of the viewing chamber showed improved restriction of the growth to a well-defined area of the channels and reduction of the standard deviation (n = 6) from 1368 μm to 101 μm (Figure 5).

The bacterial viability was unchanged following incubation in glycerol (Supplementary Figure 1) and the Nigrosin stain disappeared quickly upon initiating the fluidic flow, indicating that this and glycerol are removed alongside the spent media. During the experiments we acquired time-lapse videos of the biofilm formation and did not observe any obvious side effects on biofilm development by using glycerol and Nigrosin for seeding (data not shown).
mixed-species biofilms also indicated that this protocol affects neither single- nor dual-species *P. putida* and *X. retroflexus* biofilms (Figure 6).

Even though studies of bacterial growth under continuous flow can be challenging, fluid shear has great impact on the phenotype and is relevant for understanding biofilm development in natural habitats [3,5]. Mixed-species communities have traditionally not received much attention, but a recent acknowledgement of microbial communities being heterogeneous, coordinated and affected by social interactions have emphasized the importance of community-intrinsic properties [16]. With this new protocol (found in the supplementary material) we aim to enable improved understanding of unique properties emerging in communities exposed to flow and reduce the gap between studies of mono- and mixed-species communities.

### Bacterial Strains & Conditions

*P. bagley* _gfpmut3-Gen*<sup>+</sup>_ was introduced into *P. putida* KT2442 by a mini-Tn7 system previously described by Choi and Schweizer [17]. *P. amylolyticus* and *X. retroflexus* were originally isolated simultaneously from the same specific soil environment [18] and characterization of wrinkled variant as well as introduction of fluorescent markers were performed in a previous study [15]. All Gram-negative strains were chromosomally tagged using a Tn7-based system, while the Gram-positive *P. amylolyticus* harbours plasmid pCM20 encoding _gfpmut3_. All strains were grown in premixed Tryptic Soy Broth (TSB) (Sigma-Aldrich). In order to prevent plasmid loss, single-species cultures of *P. amylolyticus* and dual-species cultures of *P. amylolyticus* and *X. retroflexus* were supplemented with 5 μg/ml erythromycin. All overnight cultures were grown at 250 R.P.M. at 24°C.

### Viability in High Viscosity

Overnight cultures of *P. amylolyticus*, *P. putida* and *X. retroflexus* were diluted equally in either glycerol (final concentration 50%) or PBS. In order to mimic the protocol setup, cells were incubated at room temperature for 45–60 min and were subsequently quantified by CFU counting on solidified TSB (1.5% agar).

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**Figure 2.** Consistent mixed-species biofilms when seeding at high viscosity. Representative confocal laser scanning microscopy tile regions of biofilm formation by *P. amylolyticus* (GFP, green) and a wrinkled variant of *X. retroflexus* (mCherry, red) 18 h post seeding with scale bars representing 100 μm. Seeding performed at normal viscosity (A–D) showed inferior establishment of mixed-species biofilms compared with seeding at increased viscosity (E–H). (I) Quantitative analysis indicated that the amount *P. amylolyticus* was more consistent when seeding at high viscosity and that the general biomass was increased overall (Mann-Whitney U test; n = 6, error bars represent standard error).

**Figure 3.** High-throughput dual-species biofilm formation with bacteria of different sizes at high viscosity. Representative confocal laser scanning microscopy images of *P. amylolyticus* (GFP, green) and a wrinkled variant of *X. retroflexus* (mCherry, red) 20 h post seeding. Five out of six channels in a column is represented as tile regions, with scale bars representing 200 μm (A–C) or as single snaps, with scale bar representing 100 μm (D–E). White lines represent the side walls of the viewing chamber.
Figure 4. Consistent high-throughput multispecies biofilm formation after seeding at high viscosity. Representative confocal laser scanning microscopy images of biofilm formation by *P. putida* (GFP, green) and a wrinkled variant of *X. retroflexus* (mCherry, red) 18-h post seeding. Biofilms formed in channels from a column (6 channels) with increased viscosity while seeding (A–C) were more consistent and well defined than those in channels from a column seeded at normal viscosity (D–F). Bacterial growth upstream of the seeding area was observed in some channels seeded at normal viscosity (illustrated in E); this was not observed in any channels seeded at high viscosity. Scale bars represent 500 μm.

**BIOFILM CULTIVATION IN THE BIOFLUX MICROFLUIDIC FLOW SYSTEM**

*X. retroflexus* and *P. putida* were tested as single- and dual-species biofilms by adjusting overnight cultures to OD$_{600}$ = 0.3 followed by addition of glycerol (final concentration 27.5%) or fresh medium (control) and Nigrosin (Sigma-Aldrich) to create working cultures with OD$_{600}$ = 0.15. A reverse flow at 4 dyn cm$^{-2}$ was provided until Nigrosin was visible in the viewing chamber of a 48-well plate in the BioFlux 1000 device. Subsequently, the inoculum was incubated for 15 min without flow in order to attach, before the inlet well was filled and the flow initiated at 0.15 dyn cm$^{-2}$ for an 18-h period. A dual-species combination of *X. retroflexus* and *P. amylolyticus* was performed similarly, but with an increased start concentration of *P. amylolyticus* at OD$_{600}$ = 0.45, with an attachment period lasting 30 min and a 20-h incubation period. The BioFlux microfluidic flow system was located in a room cooled to 20°C without the heater turned on.

**SEEDING VERIFICATION, CONFOCAL IMAGING & IMAGE ANALYSIS**

Priming of the channels with media, correct seeding and image acquisition of bacterial inoculum stained with Nigrosin was monitored and performed using a BioFlux 1000Z system, including an Axio Inverted Observer Microscope Z1 (Carl Zeiss Inc.) equipped with a 10X objective.

Time-lapse videos and final image acquisition of biofilm formation were monitored and analyzed using a confocal laser scanning microscopy instrument (Zeiss LSM 800, Carl Zeiss Inc.) equipped with a 20X objective. Images were acquired with a 20-min interval with excitation wavelengths at 561 and 488 nm applied for mCherry and GFP, respectively, while maximum emission wavelengths for mCherry and GFP were 617 and 506 nm, respectively.

Image preparation, loading and quantification of biomass volume (μm$^3$) and area (μm$^2$) were analyzed as described previously [19], using the free statistic software R [20].

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**AUTHOR CONTRIBUTIONS**

MFH, AMT, JSM, HLR and MB were all involved in the development of study design and concept, data analysis and interpretation. *gfpmut3* was introduced into *P. putida* by JSM. Acquisition of data was performed by MFH, AMT and HLR. Initial draft of the manuscript was written by MFH, and all authors contributed to revisions.

**FINANCIAL & COMPETING INTERESTS DISCLOSURE**

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SUPPLEMENTARY DATA
To view the supplementary data that accompany this paper please visit the journal website at: www.future-science.com/doi/suppl/10.2144/btn-2018-0150

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