Characterization of a novel air–liquid interface biofilm of *Pseudomonas fluorescens* SBW25

Anna Koza, Paul D. Hallett, Christina D. Moon and Andrew J. Spiers

1SIMBIOS Centre, Level 5 Kydd Building, University of Abertay Dundee, Bell Street, Dundee DD1 1HG, UK
2Scottish Crop Research Institute, Invergowrie, Dundee DD2 5DA, UK
3AgResearch Limited, Grasslands Research Centre, Private Bag 11008, Palmerston North, New Zealand

Pseudomonads are able to form a variety of biofilms that colonize the air–liquid (A–L) interface of static liquid microcosms, and differ in matrix composition, strength, resilience and degrees of attachment to the microcosm walls. From *Pseudomonas fluorescens* SBW25, mutants have evolved during prolonged adaptation–evolution experiments which produce robust biofilms of the physically cohesive class at the A–L interface, and which have been well characterized. In this study we describe a novel A–L interface biofilm produced by SBW25 that is categorized as a viscous mass (VM)-class biofilm. Several metals were found to induce this biofilm in static King’s B microcosms, including copper, iron, lead and manganese, and we have used iron to allow further examination of this structure. Iron was demonstrated to induce SBW25 to express cellulose, which provided the matrix of the biofilm, a weak structure that was readily destroyed by physical disturbance. This was confirmed *in situ* by a low (0.023–0.047 g) maximum deformation mass and relatively poor attachment as measured by crystal violet staining. Biofilm strength increased with increasing iron concentration, in contrast to attachment levels, which decreased with increasing iron. Furthermore, iron added to mature biofilms significantly increased strength, suggesting that iron also promotes interactions between cellulose fibres that increase matrix interconnectivity. Whilst weak attachment is important in maintaining the biofilm at the A–L interface, surface-interaction effects involving cellulose, which reduced surface tension by \( \sim 3.8 \text{ mN m}^{-1} \), may also contribute towards this localization. The fragility and viscoelastic nature of the biofilm were confirmed by controlled-stress amplitude sweep tests to characterize critical rheological parameters, which included a shear modulus of 0.75 Pa, a zero shear viscosity of 0.24 Pa s\(^{-1}\) and a flow point of 0.028 Pa. Growth and morphological data thus far support a non-specific metal-associated physiological, rather than mutational, origin for production of the SBW25 VM biofilm, which is an example of the versatility of bacteria to inhabit optimal niches within their environment.

INTRODUCTION

The formation of biofilms by bacteria is a key strategy in the colonization of natural environments, though biofilms are only one of a range of bacterial assemblages which occur. These range from isolated surface-attached bacteria, monolayers of associated bacteria forming microcolonies, and larger and more complex structures, including differentiated biofilms, to poorly attached or free-floating flocs and slime. Biofilm research has largely focused on submerged flow cell biofilms in which a surface-attached exopolysaccharide polymer matrix-based structure develops away from the surface into the flow of a nutrient and oxygen-rich growth medium, and where fluid flow and mass transfer affect biofilm development, structure and rheology (for recent reviews see Hall-Stoodley et al., 2004; Ramey et al., 2004; Branda et al., 2005; Battin et al., 2007; Danhorn & Fuqua, 2007; Flemming et al., 2007, and references therein). In contrast, the partially saturated fluid-filled pores and networks found in soils, and the vascular systems and cavities found in plants, are significantly more complex environments for biofilm-colonizing bacteria and might be better approximated in...
the laboratory by low-flow-rate flow cells and static liquid microcosms with variable levels of oxygen.

We have been interested in the ability of bacteria to colonize the surface layer of static bodies of liquid, otherwise known as the air–liquid (A–L) interface (Gehrig, 2005; Spiers et al., 2002, 2006; Ude et al., 2006). In this favourable niche, nutrient and oxygen gradients are opposing, with access to abundant oxygen from the air above and nutrients in solution from below. The colonization of the A–L interface requires a floating biofilm in which attachment at the meniscus to solid surfaces may be limited [A–L biofilms have been referred to as ‘pellicles’ (Branda et al., 2005)]. Both medical and environmental isolates of *Escherichia* and *Salmonella* displaying the rough, dry and rugose (rdar) phenotype are capable of producing cellulose matrix-based, well-attached and robust biofilms which colonize the A–L interface of static liquid microcosms (Römling, 2005). In a survey of plant-associated, soil- and river-isolated pseudomonads, 76% of isolates were able to form A–L biofilms after selection in static liquid microcosms, indicating that this is an evolutionarily deep-rooted ability with significant ecological advantages (Ude et al., 2006). These biofilms can be categorized on the basis of phenotype and physical robustness into physically cohesive (PC), floccular mass (FM), waxy aggregate (WA) and viscous mass (VM) classes (Spiers et al., 2006) (these are described in Table 1). Of these, the VM-class biofilms are the most fragile and appear not to rely on significant levels of meniscus growth

Mutants of *Pseudomonas fluorescens* SBW25 have been isolated that produce biofilms similar to those of the rdar mutants of *Escherichia* and *Salmonella*. In one particular SBW25 mutant known as the Wrinkly Spreader (WS), the single-base-pair mutation responsible for the overexpression of partially acetylated cellulose and the expression of an as-yet-unidentified fimbriae-like attachment factor necessary for biofilm formation have been identified (Spiers et al., 2002, 2003; Spiers & Rainey, 2005; Bantinaki et al., 2007). In the WS, bacterial attachment at the meniscus region (the air–liquid–solid surface interface) leads to horizontal growth along the A–L interface (A–L biofilm meniscus growth may be similar in many respects to that seen in flow-cell systems where the biofilm is also directly attached to a solid surface). The physical strength of the biofilm resulting from interactions between cellulose fibre, attachment factor and lipopolysaccharide, plus the relative hydrophobicity of the structure, maintain it at the A–L interface (Spiers & Rainey, 2005). Furthermore, when grown on agar plates, the WS forms a large, flat and wrinkled colony that is dramatically different from the smaller, smooth and rounded colony of wild-type SBW25. The Complementary Biofilm Forming Strain (CBFS) biofilm mutant was isolated from a cellulose-deficient SBW25 strain (Gehrig, 2005). This produces a thinner and more finely structured biofilm than the WS, and a distinctive colony morphology different from those of both the WS and wild-type SBW25. Although the underlying genetics of the CBFS are yet to be characterized.

### Table 1. Classes of A–L interface biofilms produced by pseudomonads

Biofilm attributes were compiled from Spiers et al. (2006), Ude et al. (2006) and A. J. Spiers & A. Koza, unpublished observations. Strength, ability to withstand weight applied to the top of the biofilm; Resilience, response to applied physical disturbance such as gentle or vigorous mixing; Attachment, connection to the microcosm vial walls in the meniscus region.

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Biofilm class</th>
<th>WA</th>
<th>FM</th>
<th>PC</th>
<th>VM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Occurrence Structure</td>
<td>Rare</td>
<td>Common</td>
<td>Common</td>
<td>Common</td>
<td>Common</td>
</tr>
<tr>
<td>Strength</td>
<td>Strong</td>
<td>Strong</td>
<td>Medium</td>
<td>Strong</td>
<td>One or more large, viscous masses</td>
</tr>
<tr>
<td>Resilience</td>
<td>Good</td>
<td>Dissociates into flocs, but flocs are hard to destroy</td>
<td>Very good</td>
<td>Weak</td>
<td>Very poor</td>
</tr>
<tr>
<td>Attachment and meniscus growth</td>
<td>Good</td>
<td>Some attachment and growth</td>
<td>Very good attachment and growth</td>
<td>Very poor attachment and growth</td>
<td>Evidence for EPS but not for cellulose</td>
</tr>
<tr>
<td>EPS</td>
<td>No evidence</td>
<td>Cellulose and evidence for other EPS</td>
<td>No evidence of EPS, possible cell–cell interactions</td>
<td>No evidence of EPS, possible cell–cell interactions</td>
<td>No evidence of EPS, possible cell–cell interactions</td>
</tr>
</tbody>
</table>
fully, the biofilm structure may be due to cell–cell interactions mediated by the expression of haemin storage system (Hms)-associated or regulated structures on the cell surface (Gehrig, 2005). Both the WS and CBFS form PC-class biofilms, as they produce physically cohesive, well-attached and strong biofilms at the A–L interface [in addition to these two biofilms, SBW25 also produces a flow-cell cellulose-matrix-based biofilm (Villavicencio, 2000)].

We have discovered that SBW25 also produces a VM-class biofilm in static liquid microcosms in the presence of iron and various other metal ions. Here, we characterize the SBW25 VM biofilm in terms of induction, matrix component, strength and rheology, and attachment and localization to the A–L interface. Unlike the WS and CBFS biofilms formed by mutants derived from SBW25, this SBW25 VM biofilm appears to be a physiological response of the wild-type strain rather than the result of mutation.

**METHODS**

**Bacteria, culture and microcosms.** Bacterial strains are listed in Table 2 and were cultured in King’s B (KB) medium (King et al., 1954) at 18–20 °C for all assays. KB plates contained 1.5 % (w/v) agar. Microcosms were 6 ml KB in 30 ml universal glass vials inoculated with 60 µl of an overnight shaken KB culture and were incubated statically with the lids loosely fixed to allow good aeration. Ferrous and ferric iron were provided by FeSO₄ and FeCl₃, respectively. Other metals were added in the form of CuSO₄, MnCl₃, Pb(NO₃)₂ and ZnCl₂. KB–Fe contained 1 µM FeCl₃, and iron-chelated KB (KB-DP/T) contained 20 µM 2,2-dipyridyl (Sigma) and 0.1 µM Tiron (1,2-dihydroxybenzene-3,5-disulfonic acid; Fisher Chemicals).

**Growth, colony morphology, twitching, bacterial density and hydrophobicity analyses.** Bacterial growth and cell numbers were determined by OD₆₀₀ measurements using a Spectronic Helios Epsilon spectrophotometer (Thermo Fisher Scientific) with 10 mm optical-path cuvettes. Colony morphology was examined on KB-DP/T and KB-Fe plates. Twitching was assessed by measuring the expansion of colonies across the agar–dish interface of 20 ml KB-Fe and KB-DP/T plates after 24 h incubation. The density of the compressed VM biofilm (i.e. cells and matrix but excluding liquid) was calculated from the weight of the pellet (P) and volume of the supernatant (S) after centrifugation of 6 ml KB-Fe microcosms at 16000 g for 1 min [P/(6–S); g ml⁻¹]. The relative hydrophobicity (H₅) of SBW25 cells was determined in KB medium using a modification of the microbial adherence to hydrocarbons (MATH) assay (Spiers & Rainey, 2005).

**Tests for biofilm-forming mutants.** Ten colonies were chosen randomly from KB plates spread with mature VM biofilm material from each of five independent KB-Fe microcosms incubated for 24 h. Each of the 50 colonies was used to inoculate KB-Fe and KB-DP/T microcosms, which were inspected for growth and VM biofilm formation after 24 h. A 1 ml sample of biofilm material was also recovered by centrifugation at 16000 g for 1 min from five independent KB-Fe microcosms incubated for 24 h. The supernatants were discarded and the pellets were washed in fresh KB to remove residual traces of iron before being resuspended in 1 ml KB. Aliquots (60 µl) were then used to inoculate KB-Fe and KB-DP/T microcosms, which were inspected for growth and VM biofilm formation after 24 h.

**Cellulose detection and quantification.** Qualitative assessments of cellulose expression were made by comparing the Congo red (CR) staining of colonies grown on plates containing 0.001 % (w/v) CR (Sigma). Cellulose was also detected in plates containing 10 µg Calcein ml⁻¹ (Fluorescent Brightener 28; Sigma) and viewed using an AlphaImager HP System with a ML-26 UV transilluminator (GRI). A Leica DMR fluorescence microscope was used to view cellulose after staining with Calcein (Spiers et al., 2003). Cellulose levels were measured using a modified CR-binding assay (Spiers et al., 2003). Microcosms were gently mixed and CR was added to 0.001 % (w/v). After 2 h, the samples were remixed and a 1 ml aliquot was removed to determine the initial A₄₉₀. The samples were then centrifuged at 16000 g for 1 min to pellet the biofilm material, and the A₄₉₀ of the supernatant containing unbound CR was determined. CR binding was determined from the difference between A₄₉₀ measurements.

**Biofilm strength, attachment and manipulation.** Biofilm strength, as measured by the maximum deformation mass (MDM), was determined by carefully placing 0.0115 g glass balls onto the centre of biofilms until the biofilm broke or sank to the bottom of the microcosm (Spiers et al., 2003; Spiers & Rainey, 2005). Attachment to the glass vials in the meniscus region was measured by crystal violet staining (A₅₇₀), as described previously (Spiers et al., 2003). Both MDM and attachment were standardized for cell density using OD₆₀₀ measurements. The involvement of attachment and buoyancy in the localization of the biofilm was assessed by slowly adding KB to mature VM biofilm microcosms 1 cm below the surface over 2 h and observing whether the biofilm moved upwards with the liquid level (which was adjusted by 5 mm), flooded but remained in place or sank to the bottom of the microcosm vial. Aliquots (60 µl) of water or 10 mM FeCl₃ were similarly added to VM biofilm microcosms and allowed to equilibrate for 2 h before MDM assay to determine whether iron affected biofilm strength.

**Biofilm rheology.** Various parameters of the VM biofilm were measured by controlled-stress amplitude sweep tests using a rotational rheometer (MARS rheometer, Thermo Scientific fitted with 35 mm diameter parallel plates (PP35) and running HAAKE.

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**Table 2. P. fluorescens SBW25 and mutant strains used in this study**

<table>
<thead>
<tr>
<th>Strain</th>
<th>Description and genotype</th>
<th>Reference</th>
</tr>
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<tbody>
<tr>
<td>JB01</td>
<td>Mutant overexpressing cellulose, SBW25 <em>ppl</em>-wss, Km&lt;sup&gt;R&lt;/sup&gt;</td>
<td>Spiers <em>et al.</em> (2002)</td>
</tr>
<tr>
<td>PBR840</td>
<td>Mutant that does not express pyoverdine, SBW25 Δ<em>pvdL</em></td>
<td>Moon <em>et al.</em> (2008)</td>
</tr>
<tr>
<td>SM-13</td>
<td>Mutant that does not express cellulose, SBW25 <em>wscB</em>-mini-Tn5, Km&lt;sup&gt;R&lt;/sup&gt;</td>
<td>Spiers <em>et al.</em> (2002)</td>
</tr>
<tr>
<td>ViscA&lt;sup&gt;+&lt;/sup&gt;</td>
<td>Mutant that does not express viscosin, SBW25 <em>visCA</em>-::TnMod-Okm, Km&lt;sup&gt;R&lt;/sup&gt;</td>
<td>de Bruijn <em>et al.</em> (2007)</td>
</tr>
</tbody>
</table>
RheoWin 3 software. Biofilm samples were from independent KB-Fe microcosms incubated for 48 h. Liquid media were removed from the microcosms by pipette before the biofilm was carefully poured onto the MARS plates and excess liquid drained off with a paper towel. Samples were placed onto the bottom plate and the gap between plates was closed to 0.2 mm. An oscillating stress ramp was applied at an angular frequency of 0.5 Hz over a stress range of 0.001–1 Pa to cover the linear viscoelastic region and flow point. Measurements were made at 20 ± 0.1 °C, which was controlled with an integrated peltier thermal control unit. In an oscillating rheology test, the time-dependent phase-shift between applied torque and angular displacement can be used to determine numerous rheological parameters (Mezger, 2006). The energy required for deformation is defined by the complex shear modulus (G'). In viscoelastic materials, G' comprises elastic (i.e. reversible) energy defined by the storage modulus (G''), and viscous (i.e. irreversible) energy defined by the loss modulus (G''). The ratio of viscous to elastic energy (G'':G') defines the loss factor (tan δ). This can be used to differentiate liquids (tan δ >1) from gels or solids (tan δ <1) (Mezger, 2006). The resistance to flow is defined as viscosity, with the zero-point viscosity (η0) recorded here. Zero-point indicates that this was measured in the linear viscoelastic region, where η0, G' and G'' do not change with the applied stress level. Above a critical stress level, all rheological parameters described above will decrease, resulting in the eventual breakdown of the material with the flow point occurring when G' = G''.

Surface tension measurements. Surface tension was determined at 20 °C using 5 ml samples of vortexed KB-Fe microcosms. Measurements were made with a Kruess K100 Mk2 tensiometer (Krüss) using an SV23 Al/PTFE conical sample vessel and platinum testing rod, following the surface and interfacial tension method with the Krüss Standard Rod and SFT Rod configuration template settings.

Statistical analysis. Statistical analysis was undertaken using JMP 7.0 (SAS Institute). Data were assumed to have normal distributions. Differences between means were tested by ANOVA, Tukey–Kramer honestly significant difference (HSD) and t tests, which assumed unequal variances. Data are presented as mean ± SE, where appropriate.

RESULTS

Preliminary description of the SBW25 VM biofilm

Wild-type SBW25 growing in static KB microcosms containing 1 μM FeCl₃ (KB-Fe) produced a VM-class biofilm within 24 h, consisting of a large, viscous mass located at the A–L interface with little apparent attachment to the vial walls at the meniscus region (Fig. 1). In contrast, SBW25 cultures grown in iron-chelated KB (KB-DP/T) microcosms showed no sign of biofilm production after 24 h, despite reaching a similar final population density as determined by OD₆₀₀ measurements. Early tests indicated that no biofilm was produced in standard KB microcosms, confirming previous observations of SBW25 behaviour in comparison with WS biofilm formation (Spiers & Rainey, 2005). However, subsequent tests with KB made with water from a different source suggested that low levels of iron contamination were sufficient to result in partial VM biofilm formation. Therefore, all subsequent experimentations compared the non-biofilm iron-chelated KB-DP/T control with the biofilm-inducing KB-Fe. When disturbed by gently shaking the vial, the VM biofilm sank slowly to the bottom, whilst vortexing destroyed all structure and left no visible clumps of material remaining. The relative weakness of the VM biofilm was confirmed by maximum deformation mass measurements, whereby only 0.023–0.047 g of glass balls was able to be supported by the biofilm before breaking, in contrast to the more robust PC-class WS biofilm, which can support ~0.36 g (Spiers & Rainey, 2005). Extracellular cellulose fibres were observed by fluorescence microscopy in VM biofilm samples stained with Calcofluor, suggesting that cellulose provided the main structural (matrix) element of this weak biofilm (Fig. 2). In addition, colonies recovered from the VM biofilm showed no significant difference in morphology from those of standard KB microcosms. This is in contrast to the biofilm-forming mutants WS and CBFS, which produce colonies that are morphologically distinct from those of the wild-type SBW25, suggesting that the VM biofilm results from physiological induction by iron rather than from mutation.

Induction of the SBW25 VM biofilm

We examined the effect of FeCl₃ concentration on the induction and development of the SBW25 VM biofilm via measurements of growth, biofilm strength and attachment. As expected, SBW25 growth was found to increase with the addition of iron in the range of 0–50 μM FeCl₃ (Fig. 3a); thus, measurements of biofilm strength and attachment were standardized for cell density differences. Biofilm strength, as determined by the MDM assay, was found to increase with increasing FeCl₃ concentration (Fig. 3b), demonstrating a positive response to the presence of iron; a 1.85 ± 0.185 × increase in MDM was found between biofilms containing 1 μM FeCl₃ and biofilms grown in KB-Fe microcosms.

![Fig. 1. A novel VM-class biofilm is induced in P. fluorescens SBW25 by iron when incubated in static KB microcosms. SBW25 cultures in 6 ml KB liquid microcosms show signs of growth, including increased turbidity and siderophore expression near the top of the liquid column, within 24 h at 18–22 °C. Shown on the left is a KB-DP/T microcosm in which all free iron is chelated; on the right is a KB-Fe microcosm with 1 μM FeCl₃ added, in which a VM biofilm has developed at the A–L interface.](Image)
grown with 0.1 and those grown with 50 μM FeCl₃ (t test, DF₂₉,5, P<0.0001), though this assay did not account for growth differences. This increased SBW25 twitching might account for the decrease in levels of VM biofilm attachment in response to iron, as in PAO1, twitching cells do not form cell clusters or go on to develop biofilms.

A mutant (PBR840) deficient in the production of the iron-scavenging siderophore pyoverdine (PVD), a central component in SBW25 iron homeostasis (Moon et al., 2008), was also found to be able to form VM biofilms with either FeSO₄ or FeCl₃, suggesting that the induction mechanism is independent of this iron-uptake pathway. We further tested the specificity of iron in biofilm induction, and found that several other compounds were able to induce the VM biofilm, including 5 μM CuSO₄, 25 μM MnCl₂ and 25 μM Pb(NO₃)₂, indicating that induction is not iron-specific and may be a common response to a wider range of metals. However, of the metals tested, induction was not seen with 0.1–50 μM ZnCl₂, and in addition, the MgCl₂ in KB was found not to be necessary for induction in the presence of iron.

We examined SBW25 VM biofilm cells and derived cultures for the presence of iron-sensitive biofilm-forming mutants, which might provide an alternative explanation for the development of VM biofilms. We tested 50 randomly selected and independently isolated colonies from VM biofilms and found that none produced biofilms in KB-DP/T microcosms, but all produced biofilms in KB-Fe microcosms. Similarly, VM biofilm inocula only produced VM biofilms in KB-Fe microcosms and not in KB-DP/T microcosms. If mutation was responsible for VM biofilm formation in KB-Fe microcosms, then we would have expected to be able to isolate mutants that formed VM biofilms independently of iron in subsequent KB-Fe microcosms. These observations support our conclusion that VM biofilm formation is most probably a non-specific physiological response by SBW25 to iron in static KB microcosms. Nonetheless, iron can be used to induce this biofilm and we have used it as a representative metal to allow further investigation of the VM biofilm.

**Cellulose has a structural role as the matrix element of the VM biofilm**

Cellulose expression by SBW25 has previously been demonstrated by staining samples with Calcofluor and observation by fluorescence microscopy (Spiers et al., 2003; Spiers & Rainey, 2005). Using the same technique, we identified the presence of large masses of extracellular cellulose fibre in VM biofilm samples (Fig. 2a). We were able to detect low levels of cellulose expression by SBW25 grown in KB, confirming earlier observations of unregulated, basal expression (Spiers et al., 2003), but were not able to detect cellulose in SBW25 samples grown in KB-DP/T. Cellulose levels, as measured by the CR-binding assay, were 1.2 times greater in KB-Fe microcosms containing VM biofilms than in KB-DP/T microcosms.
which did not contain biofilms (t test, \(DF_{11.43}, t_{10.81}, P<0.0001\)). Finally, we also examined cellulose expression in colonies grown on plates containing Calcofluor (Fig. 2b), which again demonstrated that iron induced SBW25 to express cellulose.

Although iron clearly induced the expression of cellulose by SBW25 in the VM biofilm, it was not clear whether cellulose was necessary for VM biofilm formation (in the sense that cellulose provided the biofilm matrix and was largely responsible for the biofilm structure). In order to determine this, we investigated VM biofilm formation using two genetically defined SBW25 mutants. We first determined that the cellulose-deficient strain SM-13 was unable to form a VM biofilm in static KB-Fe microcosms. OD\(_{600}\) measurements confirmed that SM-13 population sizes were similar to those of SBW25 in KB-Fe, indicating that reduced cell numbers could not explain the lack of biofilm formation. Therefore, cellulose production is essential for VM biofilm formation. These findings are further confirmed by an examination of JB01, a strain which constitutively expresses cellulose, and has previously been reported to produce a very weak non-WS-like biofilm (Spiers et al., 2002). In KB-DP/T, JB01 produced a biofilm visually indistinguishable from and of similar strength to the SBW25 VM biofilm in KB-Fe, but the JB01 biofilm in KB-Fe was 4.5-fold stronger than the VM biofilm in KB-Fe (Tukey–Kramer HSD, \(q^*_{2.521}, \alpha_{0.05}\)), suggesting either that JB01 expressed more cellulose in KB-Fe than in KB-DP/T, or that the presence of extracellular iron resulted in a stronger biofilm by promoting extra matrix connectivity. The former is supported by a comparison of Calcofluor-stained colonies, which suggests that JB01 grown on KB-Fe plates expresses more cellulose than either JB01 on KB-DP/T or SBW25 on KB-Fe, which express similar amounts of cellulose (Fig. 2b), whilst the latter is addressed below.

**Iron promotes matrix connectivity in the VM biofilm**

We examined whether iron can promote interactions between cellulose fibres to increase matrix connectivity. SBW25 is known to express small amounts of cellulose in KB (Spiers et al., 2003), and it is possible that this basal level of cellulose is cross-linked by iron to produce the VM biofilm without requiring further cellulose expression, which iron has been shown to induce (notwithstanding the fact that cellulose expression was undetectable in KB-DP/T as determined by Calcofluor fluorescence microscopy). We tested this possibility by adding 100 μM FeCl\(_3\) to non-biofilm-forming cultures of SBW25 and SM-13 KB-DP/T microcosms, and in neither case did the cultures condense to form a biofilm or gel-like mass at the bottom of the vial within 2 h. However, the addition of 100 μM FeCl\(_3\) to mature VM biofilms significantly increased biofilm strength after 2 h by 1.7-fold, compared with VM biofilms to which water-only additions were made (t test, \(DF_{12.99}, t_{2.078}, P=0.0580\)). These results support the hypothesis...
that iron can promote extra matrix connectivity within existing biofilms, and results in stronger biofilms.

**Rheometry of the VM biofilm**

Increasingly, flow-cell biofilms have been directly examined by rheological measurements in order to define their rheological (structural) characteristics [for a brief introduction and a review of rheology, see Janmey & Schliwa (2008) and Janmey et al. (2007), respectively]. As these biofilms develop in a steady flow of nutrients, of key interest has been how the biofilm structure resists the pressure of the liquid moving across the surface of the biofilm (the pressure acting on one surface is known as `shear`). Flow-cell biofilms appear to have the general characteristics of a viscoelastic structure (Hall-Stoodley et al., 2004) which is able to absorb the energy of low levels of shear by slight deformation without catastrophic failure (i.e. breaking into pieces), and release the stored energy by recovery when the shear is reduced. Above a critical shear (known as the flow point), the structure begins to flow like a viscous fluid. Below this point, rheological behaviour is constant in what is known as the linear viscoelastic region.

This region, a structure can be characterized by several parameters, such as the storage (G'), loss (G'') and shear (G_d) moduli, loss factor (tan δ), flow point and zero shear viscosity (η0) (see Methods for further information), which can be used to determine whether a structure has a viscoelastic nature.

We determined that the VM biofilm was also a viscoelastic solid, with tan δ < 1 at 0.47 ± 0.03. As expected with such structures, the elastic portion of viscoelasticity (G') decreased with increasing shear stress, indicating shear-thinning behaviour, and liquid flow occurred at 0.028 ± 0.004 Pa. The VM biofilm was found to have a G' of 0.75 ± 0.16 Pa and η0 of 0.24 ± 0.05 Pa s⁻¹, which are indicators of structural stability and resistance to deformation.

**Localization of the VM biofilm to the A–L interface**

The low levels of attachment of the SBW25 VM biofilm to the vial walls and the ready sinking after gentle disturbance raise the question of how the SBW25 VM biofilm is localized to the A–L interface of the KB microcosm. Here we considered four possibilities: buoyancy (where the mass of the biofilm is less than that of the displaced KB), bacterial cell hydrophobicity (resulting in the cells having a strong preference to be at the surface rather than being submerged), weak meniscus attachment (as has already been observed, Fig. 3c) and surface interactions (where other interactions of biofilm components with the surface contribute to the localization of the biofilm). The SBW25 VM biofilm was more dense than KB medium (0.47 ± 0.09 g ml⁻¹ compared with 0.303 g ml⁻¹) and no significant difference in relative hydrophobicity was found between biofilm samples from KB-Fe microcosms and SBW25 cultures grown in KB-DP/T (t test, DF=5.182, t=−0.7164, P=0.5047), indicating that buoyancy and hydrophobicity do not contribute to the localization of the SBW25 VM biofilm to the A–L interface. In contrast, we could demonstrate that the weak attachment was important by increasing the liquid level by 5 mm in mature SBW25 VM biofilm microcosms. Rather than being moved upwards, the biofilms were flooded by a layer of KB, indicating that they were maintained in place by attachment to the vial walls. However, the flooded biofilms clearly sagged in the middle (Fig. 4), suggesting that some form of surface interaction was also involved in the localization of the biofilm to the A–L interface.

Possible surface interactions might be revealed by altered A–L surface tension (γ), as γ is altered by adsorption layers formed by surface-active agents such as amphipathic compounds and surfactants (Van Hamme et al., 2006). SBW25 is known to produce the surfactant viscosin (de Bruijn et al., 2007), which is not required for VM biofilm formation, as the viscosin-deficient mutant ViscA⁻ was found to produce an SBW25 VM-like biofilm in KB-Fe (data not shown). We determined the γ of vortexed SBW25 VM biofilms and VM-like biofilms from ViscA⁻, and of static cultures of SM-13 in KB-Fe microcosms (Fig. 5) (although all measurements were made using KB-Fe, we noted that γ of sterile KB-Fe was 1.06-fold that of KB-DP/T (t test, DF=5.26, t=2.08, P=0.0001). These data indicate that a significant reduction of γ (~19.5 mN m⁻¹) could be attributed to viscosin, as expected, and a smaller reduction (~3.8 mN m⁻¹) to cellulose, demonstrating that cellulose can act as a surface-active agent, and might enable an interaction between the A–L interface and the SBW25 VM.
biofilm. These investigations suggest that the weak levels of attachment and possible surface interactions can explain the localization of the VM biofilm to the A–L interface.

**DISCUSSION**

Although many pseudomonads are known to produce biofilms, *P. fluorescens* SBW25 is the first to be reported to produce submerged flow cell and static microcosm A–L biofilms, including two PC-class biofilms (WS and CBFS) and the VM-class biofilm reported here. The SBW25 VM biofilm is induced by the supplementation of KB static liquid microcosms with iron, as well as several other metals, and in contrast to the WS and CBFS biofilms, our results indicate that the SBW25 VM biofilm is not the result of mutation. Instead, it appears to be the result of a non-specific response to metals, including copper, iron, lead and manganese, and we have used iron as a representative metal to allow further investigation of the VM biofilm. In SBW25, it has been demonstrated that a number of different metal ions are also able to induce the transcription of a putative gene involved in copper efflux (Zhang & Rainey, 2007); thus, such non-specificity of metal ion induction is not new. Furthermore, it has been found that iron induces cellulose expression and also contributes to the overall strength of the biofilm, most likely by enhancing matrix interconnectivity.

The mature SBW25 VM biofilm is a very fragile structure, which gives the impression of being a body of viscous liquid, rather than a discrete physical structure. However, rheological testing confirms that the VM biofilm is a viscoelastic structure rather than a viscous liquid. The physical robustness of the VM biofilm can be compared with those of both the WS and the CBFS biofilms using the direct measurements of load determined by MDM assay, which suggest that the VM biofilm is an order of magnitude weaker than these two PC-class biofilms. The shear modulus (G*) of the VM biofilm (0.75 Pa) can also be compared with that of submerged flow-cell biofilms: G* varies considerably over 0.1–1000 Pa (Hall-Stoodley et al., 2004) and *P. aeruginosa* biofilms have been reported with values of 1–280 Pa (Stoodley et al., 2002), indicating that the VM biofilm is one of the least robust biofilms yet investigated. Similarly, the shear stress needed to produce viscous flow in biofilms can be compared: the VM biofilm begins to flow at 0.028 Pa, whereas in *Staphylococcus aureus* and *P. aeruginosa* flow-cell biofilms, this occurs at 0.26 and 950 Pa, respectively (Körstgens et al., 2001; Rupp et al., 2005).

Although β(1→4)-linked polymers such as cellulose tend to be rigid, and cross-connections often result in robust gels, some poorly connected exopolysaccharides produce viscous liquids instead (Sutherland, 2001). In the WS, cellulose is the major matrix component, but it is the interaction of the partially acetylated cellulose with attachment factor, LPS and the bacterial cells which results in a robust biofilm (Spiers & Rainey, 2005). The fragility of the VM biofilm might therefore be explained by the lack of attachment factor, which changes the relative hydrophobicity of SBW25 cells, thereby aiding recruitment to the A–L interface, adds strength to the biofilm, and provides a means of attachment to the meniscus region of the microcosm vial.

The fragility of the VM biofilm and its globular, viscous appearance suggest that this biofilm is more similar to slimes and mucoidal colonies than the classical well-attached and robust flow-cell biofilms. Such weak structures are known to be important in plant colonization and pathogenesis, e.g. *Agrobacterium* colonization of root hairs, *Ralstonia*, *Xanthomonas* and *Xylella* infections of plant vasculature, and *Erwinia* and *Pseudomonas syringae* colonization of leaf surfaces and mesophyll tissue (Ramey et al., 2004; Danhorn & Fuqua, 2007). These weak structures are probably maintained by internal cohesion resulting from interconnections between matrix components rather than by resistance to external forces (Klapper & Dockery, 2006). The colonization of liquid-filled cavities and surfaces subject to low flow rates and shear might only require VM-biofilm-like physical resilience and not the enhanced robustness of flow-cell or WS and CBFS biofilms. SBW25 shows a fitness advantage when able to form a VM biofilm in KB-Fe microcosms compared with the cellulose-deficient mutant SM-13 (A. J. Spiers & A. Koza, unpublished observations), confirming that this fragile
biofilm is ecologically advantageous in a static liquid microcosm [WS and CBFS mutants have a similar fitness advantage in static liquid microcosms (Spiers et al., 2002; Gehrig, 2005), whilst cellulose expression disadvantages the WS on agar plates (Spiers, 2007)].

The relatively poor levels of attachment of the SBW25 VM biofilm to the meniscus region raise the question of how this biofilm localizes to the A–L interface. In both the CBFS and WS, enhanced levels of attachment and a generally hydrophobic nature suggest that a combination of suspension and surface-tension floating (after Fidalgo et al., 2006) maintain these PC-class biofilms at the A–L interface. Although the weak levels of attachment in the VM biofilm are important, the sagging of the biofilm observed after it was flooded with additional KB suggests that surface interactions also contribute to the localization of the biofilm. The amphiphilic nature of extracellular polysaccharides (EPS) means that they can have a surface-active effect, and an additional role for EPS in biofilms has been posited in interface interactions (Flemming et al., 2007). Surface tension measurements indicated that the partially acetylated cellulose expressed by SBW25 in VM biofilms has a measurable surface activity, in agreement with findings for purified cellulose derivatives, which can alter the $\gamma$ of water to 37–63 mN m$^{-1}$ (Persson et al., 1999). It is therefore possible that cellulose fibres mediate interactions between the main biofilm mass below the surface and the A–L interface itself to help maintain the biofilm at the surface.

The ability of $P$. fluorescens SBW25 to produce both PC- and VM-class biofilms underlies the ecological advantage of being able to produce an A–L interface biofilm, and reflects the plasticity of bacterial responses to environmental challenges. In the case of the WS and VM biofilms, cellulose is utilized to produce substantially different biofilms. In the WS, the molecular mechanism that results in cellulose and attachment factor expression and biofilm formation is reasonably well understood. Preliminary investigation of the VM biofilm has shown that the response regulator WspR involved in the WS phenotype (Spiers et al., 2002, 2003) is not required, and that transcription of the wss cellulose synthase operon is not upregulated by iron, whilst de novo transcription–translation is needed, suggesting the involvement of unidentified genes in VM biofilm formation (A. J. Spiers & A. Koza, unpublished data). Further research is also needed to determine whether SBW25 responds to increasing levels of iron (and other metals) by producing more cellulose, or whether the effect of the metal is a simple on/off response.

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