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The biosurfactant viscosin transiently stimulates n-hexadecane mineralization by a bacterial consortium

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Abstract Pseudomonas produces powerful lipopeptide biosurfactants including viscosin, massetolide A, putisolvin, and amphisin, but their ability to stimulate alkane mineralization and their utility for bioremediation have received limited attention. The four Pseudomonas lipopeptides yielded emulsification indices on hexadecane of 20–31 % at 90 mg/l, which is comparable to values for the synthetic surfactant Tween 80. Viscosin was the optimal emulsifier and significantly stimulated n-hexadecane mineralization by diesel-degrading bacterial consortia but exclusively during the first 2 days of batch culture experiments. Growth of the consortia, as determined by OD$_{600}$ measurements and quantification of the alkB marker gene for alkane degradation, was arrested after the first day of the experiment. In contrast, the control consortia continued to grow and reached higher OD$_{600}$ values and higher alkB copy numbers during the next days. Due to the short-lived stimulation of n-hexadecane mineralization, the stability of viscosin was analyzed, and it was observed that added viscosin was degraded by the bacterial consortium during the first 2 days. Hence, viscosin has a potential as stimulator of alkane degradation, but its utility in bioremediation may be limited by its rapid degradation and growth-inhibiting properties.

Keywords Alkanes · Biodegradation · Lipopeptides · Pseudomonas

Introduction

Pollution with petroleum hydrocarbons is widespread and represents a problem for the environment as well as human health. Petrol and diesel contain a high amount of aliphatic hydrocarbons, in particular C$_{14}$ to C$_{20}$ alkanes (Liang et al. 2005; Stroud et al. 2007). The potential to degrade alkanes has been identified in a great variety of bacteria including Proteobacteria as well as Actinomycetales (van Beilen and Funhoff 2007). The enzyme alkane hydroxylase, AlkB, is responsible for the first step in degradation of C$_{5}$–C$_{16}$ alkanes (Ji et al. 2013), and the alkB gene has been widely used as a marker for the microbial alkane degradation potential due to its widespread occurrence among degraders (Perez-de-Mora et al. 2011; Jurelevicius et al. 2013).

Microbial degradation of alkanes is often limited by the low bioavailability of these hydrophobic pollutants. Microorganisms employ two general strategies to access hydrophobic substrates and hence circumvent the limiting bioavailability. First, they can rely on biosurfactant-mediated solubilization. Biosurfactants are amphipathic molecules with varying structures produced by a diverse group of bacteria (Van Hamme et al. 2006). They can facilitate interactions between the bacteria and alkanes by generating emulsions or by pseudosolubilization of the alkane into hydrophilic micelles (Perfumo et al. 2010; Ward 2010). Second, bacteria can access alkane drops by direct cell contact and even form biofilms on the hydrophobic substrate (Perfumo et al. 2010). Biosurfactants also play a role in the direct access strategy by increasing the cell surface hydrophobicity of the degrading organism, thereby increasing adherence to the hydrophobic substrate (Hommel 1990; Bouchez-Naitali et al. 1999; Zhang and Miller 1995; Kumar et al. 2008; Al-Tahhan et al. 2000; Perfumo et al. 2010; Ward 2010; Djeridi et al. 2013). Biosurfactants may even affect the direct access strategy in a more indirect manner, as they play several roles in biofilm...
formation, and in some cases have been shown to dissolve established biofilms (Tribelli et al. 2012; Whyte et al. 1999; Ward 2010; Rivardo et al. 2009; Kuiper et al. 2004).

Various studies have been conducted on addition of biosurfactants such as rhamnolipids (Lawniczak et al. 2013) and the Bacillus lipopeptide surfactin (Moran et al. 2000; Olivera et al. 2000; Whang et al. 2008) in bioremediation processes of petroleum hydrocarbon/alkane pollutions. Often, biodegradation is stimulated by these biosurfactants (Moran et al. 2000; Kang et al. 2010), but inhibitory effects have also been observed and may be caused by toxicity of the biosurfactant, by formation of biosurfactant micelles that offer limited access to the substrate or by destruction of degrading biosurfactant, by formation of biosurfactant micelles that offer limited access to the substrate or by destruction of degrading biofilms (Van Hamme and Ward 2001; Zeng et al. 2011; Chrzanowski et al. 2011; Kuiper et al. 2004). The use of biosurfactants in bioremediation is however still hampered by lack of detailed understanding of the impact of biosurfactants on degrader activity and performance in natural environments.

It is often highlighted that biosurfactants are biodegraded easier than chemical surfactants (Mulligan 2005), but fast biodegradation of an exogenously added biosurfactant may not only be advantageous, as fast degradation may limit its potential for usage in bioremediation. However, the stability of biosurfactants is rarely investigated in alkane biodegradation studies (Frank et al. 2010).

Most studies of biosurfactant-facilitated alkane degradation have focused on single strains or mixtures of a few strains. These studies have improved understanding of the detailed mechanisms of biosurfactant-mediated hydrocarbon uptake but even revealed that such mechanisms are often strain-specific (Thavasi 2011; Ward 2010). Thus, the effects of biosurfactants added to degrader consortia are still lacking attention to improve realism of the obtained conclusions. Bacteria in a consortium probably use different hydrocarbon accession strategies, and Owssianik et al. (2009) suggested that addition of biosurfactants to consortia stimulates biodegradation performed by the degraders taking up solubilized hydrocarbons. On the other hand, bacteria using the direct substrate accession mode may be negatively affected by biosurfactants.

Pseudomonas produces several powerful lipopeptide biosurfactants with low critical micelle concentrations (CMCs) (Raaijmakers et al. 2010). These lipopeptides, including the compounds viscosin, massetolide A, putisolvin, and amphisin, have primarily been investigated for their antimicrobial properties and their effects on biofilm formation (Raaijmakers et al. 2010). However, the ability of Pseudomonas lipopeptides to stimulate mineralization of hydrophobic hydrocarbons and their utility for bioremediation have so far received very limited attention. The aims of the current study are (1) to provide information on the ability of Pseudomonas lipopeptides to solubilize the model alkane, n-hexadecane; (2) to determine the impact of the lipopeptide viscosin on mineralization of n-hexadecane and growth by a bacterial consortium; and (3) to obtain information about the stability of viscosin in the presence of the n-hexadecane-degrading consortium.

Materials and methods

Isolation of a diesel-degrading bacterial consortium from soil

Diesel-degrading bacteria were isolated from soils polluted with hydrocarbons (Aabenraa, Denmark). Initially, bacteria were extracted from the soils as previously described by Johnsen and Karlson (2005) with a few adjustments. Suspensions of 0.50 g soil and 4.5-ml 2 mM sodium pyrophosphate solution (Na$_4$P$_2$O$_7$·10H$_2$O) (pH=8.0) were mixed for 30 min on a gyroratory shaker. The suspensions were streaked on Bushnell Haas (BH) (Sigma-Aldrich) agar plates with 0.1 % sterile filtered diesel (Shell) and 0.1 % nystatin (ICN Biomedicals Inc.) to inhibit fungal growth. Plates were incubated for 2 weeks at room temperature. Colony material from the plates was washed off by 0.9 % NaCl and used as inoculum for a culture in liquid BH medium with 0.1 % sterile filtered diesel and 0.1 % nystatin. The culture was incubated for 13 days at room temperature with stirring, whereupon the culture (diesel-degrading consortium) was stored at −70 °C as 1-ml aliquots of a 50 % v/v glycerol/bacterial suspension.

Purification of lipopeptide biosurfactants

The lipopeptides viscosin, massetolide A, putisolvin, and amphisin were purified from the producing strains Pseudomonas fluorescens SBW25 (Thomson et al. 1995), P. fluorescens SS101 (de Souza et al. 2003), Pseudomonas putida 267 (Tran et al. 2008) and Pseudomonas sp. DSS73 (Sørensen et al. 2001), respectively, essentially as described by de Souza et al. (2003). Briefly, the strains were cultivated on King’s B agar plates in darkness at 28 °C for 1 day, before being transferred to 20 °C and incubated for another 3 days. Colony material was suspended in demineralized water (Milli-Q, Millipore; referred to as Milli-Q water hereafter) and homogenized by shaking. Cells and supernatant was separated two times by centrifugation at 4,700 rpm for 20 min at 4 °C on a Sigma 3-18K centrifuge (Sciequip). The supernatant was acidified to pH 2.0 with 1 M HCl and left overnight on ice for a precipitate to form. The solution including precipitate was centrifuged in sterile centrifuge tubes for 27 min at 7,000 rpm and 4 °C in a Sigma 3-18K centrifuge. The supernatant was discarded, and the precipitate was washed four times with Milli-Q water at pH 2.0. The precipitate was dissolved in Milli-Q water, and pH was adjusted to 8.0 with 1 M NaOH to fully dissolve the precipitate. The solution was
lyophilized, and the purity of the lipopeptide preparations was qualitatively analyzed by high-performance liquid chromatography.

High-performance liquid chromatography

High-performance liquid chromatography (HPLC) analysis was carried out using a Waters Alliance series 2695 system and a Waters model 996 photodiode array detector (www.waters.com). Chromatographic analysis of the lipopeptides followed the protocol by Nielsen and Sørensen (2003) with minor modifications. Briefly, a Hypersil BDS C18 column (100 × 4.6 mm and 3 μm particle size) (www.therm Scien t i fic .com) was used for separation of the lipopeptides. Solvents were HPLC-grade acetonitrile (solvent A) and 0.1% o-phosphoric acid (solvent B), mixed in a linear gradient of 15 to 100% solvent A from 0 to 40 min, and of 100 to 15% solvent A between 40 and 44 min. The flow rate was 1 ml/min, and the column temperature was 40°C. The injected sample volume was 10 μl. The lipopeptides were monitored at wavelengths of 190 to 250 nm. For quantification of viscosin in cultures of the diesel degrading consortium, a wavelength of 220 nm was applied and viscosin concentrations were calculated using purified viscosin as a standard. Handling of chromatographic data was performed using Waters Empower 2 software.

Emulsification of n-hexadecane by lipopeptide biosurfactants

The emulsification index of viscosin, amphisin, massetolide A, and putisolvin, as well as the synthetic surfactant Tween 80 (Applichem), was determined using the assay described by Cooper and Goldenberg (1987). The surfactants were dissolved in Milli-Q water (pH 7.3) at 20 and 90 mg/l. Solutions (5 ml) were aliquoted into disposable pyrex tubes and overlaid with 5 ml n-hexadecane (Sigma-Aldrich) before vortexing at high speed for 2 min. Tubes were left for 24 h at room temperature before measuring the height of the emulsified layer and the total height of the sample. The emulsification index (E24h) was estimated as height of the emulsified layer/total height of sample × 100%.

n-Hexadecane mineralization and growth of the bacterial consortium

The effect of viscosin on n-hexadecane mineralization by the degrading consortium isolated in this study was analyzed by a 14C-n-hexadecane mineralization assay. Prior to mineralization experiments, the consortium was inoculated in BH medium with 0.1% diesel at 28°C on a shaker with 150 rpm. After 24 h, a 0.1-ml aliquot was used to establish a culture in BH medium with 0.1% n-hexadecane, which was incubated under the same conditions and used as inoculum for the mineralization experiments.

The mineralization assays were carried out for 14 days in infusion flasks containing 20-ml BH medium with 100 ppm n-hexadecane (Sigma-Aldrich), 3.15 nCi [l-14C] hexadecane at a specific activity of 55 mCi/mmol (http://www.arc-inc.com/), and 100-ppm trace element solution (Kragelund and Nybroe 1994). Viscosin was added to a final concentration of 90 mg/l. Flasks were inoculated with the consortium preculture to obtain a starting concentration of approximately 107 cells/ml. 14CO2 was collected in glass tubes with 1-M NaOH solutions, and radioactivity was measured on a Beckman LS 1801 liquid scintillation counter (Beckman Instruments, Inc.). Data were corrected for background activity (typical value of 30 DPM). Growth was determined as optical density at 600 nm (OD600) measured by an UV-mini 1240 spectrophotometer (Shimadzu). For HPLC analysis, samples of 0.5 ml were removed and stored at −20°C until analysis was carried out.

DNA extraction

Samples for DNA extraction (1 ml) were collected from above experiments in RNase/DNase-free tubes. Cell pellets were obtained by removing 900 μl of the supernatant, and the pellets were immediately flash frozen in liquid nitrogen and stored at −70°C. Prior to extraction, 100 μl of lysozyme (1 mg/ml, Sigma-Aldrich) in Tris-EDTA buffer (Sigma-Aldrich) were added to the pellet and left for 20 min at room temperature. For DNA extraction, the AllPrep DNA/RNA mini kit (Qiagen, USA) was used according to the manufacturer’s protocol. The final DNA extractions were stored at −70°C until further processing.

Quantification of the alkB genes by qPCR

The copy numbers of alkB genes were quantified using qPCR. The qPCR was carried out using a MX3000P® qPCR system (Stratagene). The alkB genes were amplified using primers alkB-f (5′-AYACIGCICAYGARCTIGCAYAA-3′) and alkB-r (5′-GCRTGRTGRTCIGARTGICGYTG-3′) (Perez-de-Mora et al. 2011). This primer set amplifies alkB genes from a range of Gram-positive as well as Gram-negative bacteria without an obvious specificity for any bacterial group, and the obtained PCR product has the length of approximately 550 bp. Amplification of the fragment was carried out in 20-μl reactions using the Brilliant Master mix SYBR® GREEN which included the following: PCR buffer, dNTPs, 5 mM MgCl2, Sure start Taq polymerase, and SYBR Green 1 dye (Agilent Technologies, Santa Clara, CA). Further, reactions contained 0.8 μl of each primer (10 pmol/μl), 2 μl bovine serum albumin (New England Biolabs), 1 μl MgCl2 (50 mM) (Finnzymes), 3.4 μl dH2O, and 2 μl DNA template. All DNA
extracts were diluted tenfold before they were used in PCR reactions. The setup of the touchdown PCR program consisted of the following: initial denaturation at 95 °C for 10 min followed by 5 cycles of 45 s at 95 °C, 1 min at 62 °C (reduced stepwise to 57 °C), and 45 s at 72 °C. This was followed of 40 cycles of 45 s at 72 °C, 45 s at 95 °C, 1 min at 57 °C, and 45 s at 72 °C (Perez-de-Mora et al. 2011). The PCR program was ended with 20 min at 72 °C. The standard used was a tenfold serial dilution of \( \text{alkB} \) genes from \( Pseudomonas putida \) GPo1, and it ranged from \( 10^3 \) to \( 10^7 \) copies per qPCR reaction. The standard curve was linear with a \( R^2 > 0.99 \) with a reaction efficiency of the real-time PCR of approximately 75 %.

Data analysis

Comparison of two data sets was performed by unpaired two-tailed \( t \) test. Analysis of concentration correlation was done with linear regression analysis. Correlation analysis between different dependent variables was done using PAST statistics (http://folk.uio.no/ohammer/past/). All experiments were carried out with triplicate samples and with at least one independent repetition of each experiment. Data are presented as mean±SD. \( P <0.05 \) is selected as cutoff for statistical significance.

Results

Emulsification of n-hexadecane by purified lipopeptides

Lipopeptide preparations were initially analyzed by HPLC to ensure purity of compounds used for subsequent experiments. Figure 1 shows the chromatograms obtained for the lipopeptides viscosin, massetolide A, putisolvin, and amphisin. They were all characterized by dominating peaks complemented by a few minor peaks eluding slightly earlier

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**Fig. 1** HPLC chromatograms obtained from preparations of viscosin, massetolide A, putisolvin, and amphisin purified from \( P. \) fluorescens SBW25, \( P. \) fluorescens SS101, \( P. \) putida 267 and \( Pseudomonas \) sp. DSS73, respectively. \( AU \) absorbance units.
of, or later, than the major peak. The analysis documented that pure lipopeptides were obtained as the additional peaks in the chromatograms represent derivatives of the major lipopeptides generated as a result of flexibility of the synthetases involved in lipopeptide synthesis (Stachelhaus et al. 1999) as discussed below.

All the *Pseudomonas* lipopeptides and the commercial synthetic surfactant Tween 80 emulsified n-hexadecane with emulsification indices of 6–24 % at 20 mg/l and 20–31 % at 90 mg/l (Fig. 2). The emulsification indexes for all compounds were dependent on the concentration, with higher values at 90 mg/l than at 20 mg/l ($P<0.05$, except from viscosin with a $P$ value of 0.055). Viscosin and amphisin had comparable emulsification powers as the synthetic surfactant Tween 80, while massetolide and putisolvin were significantly weaker emulsifiers both at 20 and 90 mg/l ($P=0.000451$ and 0.0089 for massetolide; $P=0.000331$ and 0.0177 for putisolvin). Viscosin was selected for subsequent studies, as it was the biosurfactant with the highest ability to emulsify n-hexadecane.

Effect of viscosin on n-hexadecane mineralization and growth of a bacterial consortium

The effect of viscosin on n-hexadecane mineralization and growth of the diesel-degrading bacterial consortium was studied at a concentration of 90 mg/l. The consortium treated with viscosin showed a significantly higher n-hexadecane mineralization until day 2 compared to the non-treated consortium ($P=0.027$) (Fig. 3a). However, this stimulatory effect was only short lived, and from day 3, there was no significant difference between the viscosin-treated consortium and the control. After 14 days, the control and the viscosin-treated consortium had mineralized 37.3 % ($\pm9.4 \%$) and 41.3 % ($\pm9.1 \%$) n-hexadecane, respectively.

Growth of the bacterial consortium was initially assessed by OD$_{600}$ measurements. Figure 3b shows that the control and the viscosin-treated consortia initially grew comparably and reached an OD$_{600}$ of ca. 0.3 on day 1. Thereafter, growth of the viscosin-treated consortium was arrested while the control consortium continued to grow for three additional days and reached an OD$_{600}$ of approximately 0.55. At the end of the experiment, growth yield (final OD$_{600}$) of the viscosin-treated consortium was significantly lower than of the control ($P<0.05$).

To complement OD$_{600}$ measurements, we quantified alkB gene copy numbers to determine the population dynamics of bacteria harboring an alkane degradation potential in the consortia. A comparable initial increase in abundance of alkB genes was observed for both consortia during the first day (Fig. 3c), while the amount of alkB genes was significantly higher in the control than in the viscosin-treated consortium after 2 days ($P=0.0028$). In the viscosin-treated consortia, the alkB gene numbers remained constant, while those in the controls decreased after day 2 so that no significant difference was observed after 14 days ($P=0.08$). Hence, the abundance of the alkB genes showed the same dynamics as the growth pattern observed by OD$_{600}$ measurements for both the control and the viscosin-treated consortia.

To assess the ability of viscosin to stimulate n-hexadecane mineralization within the first 2 days in more detail, the mineralization per time unit normalized by the bacterial biomass expressed as OD$_{600}$ is accentuated in Fig. 3d. The viscosin-treated consortium degraded significantly more ($P<0.05$) n-hexadecane per biomass unit than the control in the early stage of the experiment. During the remaining time of the experiment, the treated and the non-treated consortium mineralized n-hexadecane at comparable rates.

In conclusion, viscosin has a positive effect on n-hexadecane mineralization in the initial phase of the experiment while overall growth is negatively affected by the biosurfactant. Furthermore, assessment of bacterial growth by measurements of OD$_{600}$ and alkB copy numbers provides comparable estimates of viscosin impact on growth dynamics.

Degradation of viscosin

The short-lived stimulation of n-hexadecane mineralization exerted by viscosin led us to investigate the stability of the biosurfactant in the experiment. Figure 4 shows the concentration of viscosin over 14 days as quantified by HPLC. The initial amount (90 mg/l) of viscosin could not be detected after the first 48 h. At day 6, samples were re-spiked with 60 mg/l of viscosin. The viscosin added on day 6 was not detected on day 7. Hence, we conclude that viscosin is rapidly degraded by members of the bacterial consortium.
Emulsification of n-hexadecane

Viscosin, massetolide A, amphisin, and putisolvin represent different groups of *Pseudomonas* lipopeptides. The viscosin-group compounds viscosin and massetolide A both consist of nine amino acids, while members of the amphisin group have 11 amino acids in their peptide part. Finally, putisolvin represents lipopeptides with 12 amino acids and a cyclization that differs from that of the viscosin- and amphisin-group compounds (Raaijmakers et al. 2010). All four lipopeptides are powerful biosurfactants able to lower the surface tension of water to about 30 mN/m (Saini et al. 2008; de Souza et al. 2003; Krujte et al. 2009). These compounds are synthetized by non-ribosomal peptide synthetases, which display substrate flexibility so that an individual strain produces a major compound plus a number of structural analogues in minor amounts.

Discussion

Emulsification of n-hexadecane

Fig. 3 The effect of viscosin on n-hexadecane mineralization and cell growth of a diesel degrading consortium. For all panels, results for consortia treated with viscosin at 90 mg/l are symbolized by squares, whereas results for control consortia are shown by circles. a Hexadecane mineralization; b growth of the bacterial consortia measured as OD<sub>600</sub>; c the dynamics of alkB gene copies; d calculated amount of n-hexadecane mineralized per day normalized by OD<sub>600</sub> units. Note that the y-axis in d has two different intervals. Error bars represent 1 SD with n=3. Error bars not shown are smaller than the symbols.

Fig. 4 The concentration of viscosin during experiments as quantified by HPLC. Dotted line represents time of spiking with 60 mg/l of viscosin. Error bars represent 1 SD with n=3. Error bars not shown are smaller than the symbols.
ment harboring introduced surfactin has previously observed in situ in a natural environ-
et al.1999). Further, 90 mg/l of the related lipopeptide biosurfactant-mediated substrate transfer (Bouchez-Naitali et al.1999). Hence, this bacterial group might be negatively affected by viscosin in the current experiments.

Degradation of viscosin

The short-lived stimulation of n-hexadecane mineralization by viscosin addition could be due to rapid degradation of viscosin by bacteria in the consortium. In spite of its rapid degradation, viscosin irreversibly inhibits the overall growth of the
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