Pseudomonas aeruginosa recognizes and responds aggressively to the presence of polymorphonuclear leukocytes

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Polymorphonuclear neutrophilic leukocytes (PMNs) play a central role in innate immunity, where they dominate the response to infections, in particular in the cystic fibrosis lung. PMNs are phagocytic cells that produce a wide range of antimicrobial agents aimed at killing invading bacteria. However, the opportunistic pathogen Pseudomonas aeruginosa can evade destruction by PMNs and thus cause persistent infections. In this study, we show that biofilm cells of P. aeruginosa recognize the presence of attracted PMNs and direct this information to their fellow bacteria through the quorum sensing (QS) signalling system. The bacteria respond to the presence of PMNs by upregulating synthesis of a number of QS-controlled virulence determinants including rhamnolipids, all of which are able to cripple and eliminate cells of the host defence. Our in vitro and in vivo analyses support a 'launch a shield' model by which rhamnolipids surround the biofilm bacteria and on contact eliminate incoming PMNs. Our data strengthen the view that cross-kingdom communication plays a key role in P. aeruginosa recognition and evasion of the host defence.

INTRODUCTION

Evading elimination by the host defence is a crucial virulence capacity of pathogens, enabling them to cause persistent infections, especially in immunocompromised patients (Kharazmi, 1991). Among known mechanisms are inhibition of the production of antimicrobial substances, antimicrobial degradation, inhibition of the chemotaxis of polymorphonuclear neutrophilic leukocytes (PMNs) and induction of PMN apoptosis (Allen et al., 2005; Bortolussi et al., 1987; Kharazmi et al., 1984a, b). The biofilm lifestyle dominates in many chronic bacterial infections (Bjarnsholt et al., 2008; James et al., 2008; Saye, 2007; Yang et al., 2008). Biofilms consist of sessile aggregates of bacteria embedded in a self-made extracellular polymeric matrix. Important hallmarks of the biofilm mode of growth are tolerance to the highest deliverable doses of antibiotics (Donlan & Costerton, 2002; Mah et al., 2003; Stewart & Costerton, 2001) and resistance to the action of the host defence system (Bjarnsholt et al., 2005). It has been suggested that during colonization and subsequent formation of biofilms Pseudomonas aeruginosa keeps expression of virulence factors and other antigenic determinants at a minimum. This stealthy approach is carefully controlled by intercellular communication denoted quorum sensing (QS) (Fuqua et al., 1994). As the local cell density attains

Abbreviations: AHL, N-acylhomoserine lactone; BAL, bronchoalveolar lavage; CF, cystic fibrosis; C4-HSL, N-butanoyl-L-homoserine lactone; 3-oxo-C12-HSL, N-3-oxododecanoyl-L-homoserine lactone; DAPI, 4',6-diamidino-2-phenylindole; LDH, lactate dehydrogenase; PI, propidium iodide; PMN, polymorphonuclear leukocyte; PNA FISH, peptide nucleic acid fluorescence in situ hybridization; PQS, Pseudomonas quinolone signal; QS, quorum sensing.

The microarray data discussed in this publication have been deposited in NCBI’s Gene Expression Omnibus (GEO, http://www.ncbi.nlm.nih.gov/geo/) and are accessible through GEO Series accession number GSE6769.

Two supplementary tables are available with the online version of this paper.
quorum size and biofilm areas expand, the production of secreted virulence factors escalates, which through a series of immune-activating events attracts PMNs. In vitro, PMNs are capable of penetrating and eliminating QS-deficient *P. aeruginosa* biofilms by phagocytosis but incapable of eliminating QS-proficient biofilms (Aaron et al., 2002; Bjarnsholt et al., 2005; Jesaitis et al., 2003). Instead of eliminating the bacteria, the PMNs are destroyed by the encounter with biofilm bacteria. It was reported previously that purified *P. aeruginosa* rhamnolipids could destroy PMNs by necrosis (Jensen et al., 2007). Like the majority of virulence factors, antibiotic tolerance and extracellular matrix DNA, rhamnolipid synthesis is delicately controlled by QS (Brint & Ohman, 1995). We speculated that rhamnolipids might function as a protective mechanism and significantly contribute to the increased resistance of biofilms to phagocytic cells. Intriguingly, Morici et al. (2007) showed that AlgR represses rhamnolipid production in *in vitro* biofilms. These authors demonstrated that AlgR exerts its action directly on the rhlA promoter and that a ΔalgR biofilm produces increased amounts of rhamnolipids. Moreover, it was found that AlgR represses the RhlR-controlled QS cascade in biofilms, including the *Pseudomonas* quinolone signal (PQS) system (Morici et al., 2007). We therefore set out to investigate the mechanisms involved in biofilm-associated protection against attracted PMNs.

**METHODS**

**Bacterial strains.** The *P. aeruginosa* PAO1 wild-type used for the experiments was obtained from the *Pseudomonas* Genetic Stock Center (http://www.pseudomonas.med.ecu.edu, strain PAO00001). This PAO1 isolate has served as DNA source for the *Pseudomonas* Genome Project (http://www.pseudomonas.com) and, subsequently, as template for design of the Affymetrix *P. aeruginosa* GeneChip. The isogenic ΔlasΔrhlR QS mutant was constructed by Bjarnsholt et al. (2005).

The rhlA mutant, isogenic to the PAO1 wild-type, was obtained from the University of Washington, Seattle, WA, USA. The ΔpqsA mutant was constructed by Aendekerk et al. (2005) and obtained from the laboratory of Paul Williams, University of Nottingham, Nottingham, UK.

**Growth of biofilms.** Biofilms were grown at 37 °C in continuous-culture silicone tubing, as described by Hentzer et al. (2003, 2005), perfused with ABT minimal medium supplemented with 0.5% (w/v) glucose (Clark & Maaløe, 1967; Pamp & Tolker-Nielsen, 2007). The medium was supplemented with antibiotics where appropriate. Unless otherwise stated, all strains were incubated in growth medium at 37 °C. Due to the need for a large quantity of cells for the transcriptomic analysis, biofilms were grown in a 12 cm long silicone flow tube (inner diameter 0.8 cm, total volume 6 ml) instead of traditional flow channels. The flow tubes were connected to a medium reservoir with silicone tubing of smaller dimension, with the inclusion of a peristaltic pump creating the liquid flow. A bubble trap was included between the flow tube and the pump, and the effluent was collected in a waste flask that was connected via silicone tubing to the distal end of the flow tube. Overnight cultures of PAO1 wild-type or the ΔlasΔrhlR mutant were diluted 100-fold and 6 ml was injected into each channel. To allow cell attachment the flow into the tube was halted for 1 h, by clamping both the influent and effluent tubes. The medium flow was set to 23 r.p.m. (60 ml h⁻¹) using a Watson Marlow 2055 pump. The relatively high flow rate was maintained to ensure that planktonic bacteria were washed out of the setup.

**PMN treatment of biofilms.** The biofilms were allowed to grow and develop in the biofilm flow tubes for 3 days before challenge with PMNs. Fresh PMNs from human volunteers were isolated as described by Bjarnsholt et al. (2005). Prior to PMN injection the flow was halted. Ten million PMNs were resuspended in 6 ml Krebs–Ringer buffer supplemented with 10 mM glucose and injected into the flow tubes. The concentration of PMNs was found by microscopy to be approximately 1 PMN per 1000 bacterial cells. PMNs and biofilm were incubated for 2 h. After incubation the fluid inside the tube was gently collected and the attached biofilm cells were mechanically removed and collected in either 6 ml RNAlater (Ambion) for transcriptome analysis or 200 μl ethyl acetate for rhamnolipid detection.

**Dynorphin A treatment of biofilms.** The biofilms were allowed to grow and develop in the biofilm flow tubes for 3 days before challenge with 100 μM dynorphin A (1-17). Prior to dynorphin A injection the flow was halted. Dynorphin A (1-17) (AnaSpec) was resuspended in 6 ml Krebs–Ringer buffer, supplemented with 10 mM glucose, to a final concentration of 100 μM and injected into the flow tubes. Dynorphin A and biofilm were incubated for 30 min or 2 h. After incubation the fluid inside the tube was gently collected and the attached biofilm cells were mechanically removed and collected in either 6 ml RNAlater for transcriptome analysis or 200 μl ethyl acetate for rhamnolipid detection.

**Quantification of rhamnolipid.** The biofilm was collected in 200 μl ethyl acetate and sonicated following evaporation. The biomass was redissolved in methanol for mass spectrometry analysis.

A standard curve for rhamnolipid B [concentration vs total ionization current (TIC)] was calculated from LC-ESI-MS data. To minimize potential differences in ionization levels of rhamnolipid between the samples the rhamnolipid standards used for calculating the concentration curve were analysed immediately prior to, as well as after the samples undergoing analysis. The TIC level was determined on the [M + NH₄]⁺ 247 ion at 668.4, over the 7 s during which rhamnolipid B was eluted.

**HPLC-MS analysis.** HPLC-MS analysis was performed with an Agilent 1100 series HPLC connected to a micromass LCT oxA TOF MS. The concentration of rhamnolipid in untreated culture was set as index 100.

In the analysis the total rhamnolipid concentration was derived from the six major rhamnolipids, with the following masses [M + NH₄]⁺: 668.4, 694.4, 696.4, 522.4, 548.4 and 550.4. These correspond to C10-C10-Rha-Rha, an unidentified C10-C12A-Rha-Rha, C10-C12-Rha-Rha, and the respective monorhamnose derivatives.

**Genechip microarray analysis of biofilms.** The RNAlater-suspended biofilm cells were sonicated three times for 10 s to disperse clumps from the biofilm. Four times 2 ml of RNAlater-suspended cells were pelleted and subsequently lysed with 100 μl of 1 mg lysozyme ml⁻¹ at room temperature for 13 min. Total RNA was isolated with RNeasy Mini kits (Qiagen) and contaminating chromosomal DNA was removed by RQ1 RNase-free DNase treatment. cDNA was synthesized from 12 μg total RNA in 1 × first-strand buffer containing 100 ng μl⁻¹ random primers (Invitrogen Life Technologies), 10 mM DTT, 0.5 mM dNTPs, 0.5 U μl⁻¹ SUPERase In (Ambion) and 25 U μl⁻¹ SuperScriptIII (Invitrogen Life Technologies). After cDNA synthesis, RNA was removed by 1 M NaOH at 65 °C for 30 min and the reaction was...
neutralized with 1 M HCl. Four micrograms of cDNA was fragmented with 0.8 U DNase I (Amersham Pharmacia Biotech) in 1 × OnePhorAll Buffer (Amersham Pharmacia Biotech). Terminal labelling of fragmented cDNA was done using the Enzo BioArray Terminal Labelling kit with Biotin-ddUTP (Enzo Diagnostics). Biotinylated cDNA was hybridized to a GeneChip P. aeruginosa Genome Array and scanned according to standard GeneChip protocols (Affymetrix). The wild-type PMN exposure was performed as three individual experiments [once in the Parsek lab (PMN donor Rasmussen) and twice in the Givskov lab (PMN donor Alhede)]. The ΔlasRAhrR mutant PMN exposure experiment was performed as two individual experiments. The transcriptome analysis was performed with ArrayStar software (DNASTAR). Performing a t-test on the dataset, genes were recorded as being (significantly) differentially expressed if the absolute value of the fold change was ≥ 3 and the P-value was ≤ 0.05. The relaxed fold change criterion was chosen because of the assumed heterogeneous nature of biofilm. It is likely that subpopulations of bacteria within the biofilm have gene expression patterns substantially different from those of the bulk of the cells in the biofilm. Inherent to the harvesting method and the microarray technique, gene expression patterns in such subpopulations will be averaged with the bulk population in the analysis. In order to allow such subpopulation gene expression patterns to be recorded in the analysis we chose a less stringent cut-off value.

**Real-time PCR analysis.** Quantitative real-time PCR was used for validation of the microarray results.

**Primer design.** Primers (see Supplementary Table S1, available with the online version of this paper) for each gene in the real-time PCRs were designed using Integrated DNA Technologies Primer Quest software (http://www.idtdna.com). Briefly, the amplicon sizes ranged from 80 to 200 bp and the primer melting temperatures were designed for 60 °C, with a melting temperature difference of less than 2 °C for each primer pair. The primer sequences were also subjected to BLAST analysis against the P. aeruginosa PAO1 genome to eliminate the possibility of non-specific binding.

**Real-time PCR.** Each real-time PCR mixture (final volume, 20 μl) contained 9 μl cDNA, 10 μl SYBR Green qPCR Master Mix 2x (Fermentas) and 500 nM of each forward and reverse primer. Real-time PCR was performed with a chromo4 (MJ Research) using the following cycling parameters: initial denaturation at 95 °C for 10 min followed by 40 cycles of 95 °C for 15 s, 60 °C for 30 s and 72 °C for 30 s.

The data were analysed using the efficiency-corrected relative quantification method (Pfaffl, 2001). To ensure correctness of the quantification we normalized the expression of the target genes to the reference gene (including housekeeping genes) was within 5% from the treated sample. To allow accurate analysis we decided that the PCR efficiency of each gene (including housekeeping genes) was within 5% from the treated condition to the non-treated. Accordingly, the average efficiencies of each gene in this study were very similar for the conditions compared, allowing accurate analysis.

**PMN killing in biofilm.** To monitor the effect of biofilms on PMNs, the biofilms including PMNs were harvested and resuspended in 200 μl PBS, vortexed and gently centrifuged followed by sterile filtering (0.22 μm). Then the lactate dehydrogenase (LDH) activity was measured using a Cytotoxicity Detection kit (Roche). LDH is a stable cytoplasmic enzyme present in all cells, which is rapidly released into the cell culture supernatant upon damage of the plasma membrane (Weyermann et al., 2005). Two hundred microlitres of reaction mixture was added to 200 μl supernatant and incubated for 30 min at room temperature. The absorbance was measured at 490 nm and 590 nm. High, low and negative controls were included in all experiments: high, biofilm including PMNs harvested in 1% Triton; low, non-exposed PMNs only; negative, water. The number of dead PMNs was calculated as a percentage relative to the high control.

The above method was verified once by FACS analysis. The biofilms including PMNs were harvested and resuspended in 400 μl PBS and stained with 5 μl FITC-conjugated CD15 antibody (Becton Dickinson). Next, 1 μl of 20 mM propidium iodide (PI) (P-4170 Sigma) was added to monitor integrity of the cells and release of DNA. This staining enabled us to quantify the dead PMNs by FACS. Fluorescence intensity of the samples was measured on a FACS Calibur (Becton Dickinson) using cellquest software.

**Lung infection of BALB/c mice by intratracheal instillation of alginate-embedded P. aeruginosa.** Female BALB/c mice were purchased from M&B Laboratory Animals at 10–11 weeks of age. The mice were of equal size, and were maintained on standard mouse chow and water ad libitum for a minimum of 1 week prior to challenge. The National Animal Ethics Committee, Denmark, authorized all animal experiments. The mouse experiments were performed as described by Pedersen et al. (1990).

**Bacteriology in bronchoalveolar lavage (BAL) fluid.** Exposed tracheas of anaesthetized mice were cannulated with a 22 gauge catheter (OPTIVA* 2; Johnson & Johnson Medical). BAL was performed by flushing five times with 1.5 ml ice-cold PBS without Ca2+ and Mg2+. The BAL fluid was stored on ice until staining for necrotic PMNs. The mean recovery of BAL fluid was 1.1 ml (CV 13%).

**Quantitative lung bacteriology.** For colony counting, the exposed lungs were isolated in 5 ml 0.9% NaCl, and homogenized on ice. A serial dilution of the lung homogenate mixed with BAL fluid was performed, and dilutions were plated on blue agar plates (State Serum Institute, Copenhagen, DK), which are selective for Gram-negative bacilli (Høiby, 1974).

**Necrotic and apoptotic PMNs in mouse BAL fluid.** PMNs were stained with an Annexin V-FITC Apoptosis Detection kit I (556747, BD Biosciences Pharmingen), using a modification of the preparation supplied by the manufacturer. Two hundred microlitres of BAL fluid was equilibrated by centrifugation with 2.5 ml cold 1 × binding buffer (BD Biosciences Pharmingen) at 350 g for 7 min at 5 °C. For discrimination between necrotic and apoptotic PMNs, the pellet was mixed with 100 μl 1 × binding buffer with 2.5 μg PI ml−1, annexin V-FITC component (1:40), and the PMN phenotypic surface marker monoclonal allophycocyanin (APC)-conjugated rat anti-Ly-6G antibody (clone RB6–8C5, BD Biosciences Pharmingen) (1:50) and incubated for 15 min at room temperature in the dark. The incubation was terminated by addition of 400 μl 1 × binding buffer and the samples were analysed by flow cytometry.

**PNA FISH.** Deparaffinized tissue sections were analysed by fluorescence in situ hybridization (FISH) using peptide nucleic acid (PNA) probe (Stender, 2003). A mixture of Texas-red-labelled P. aeruginosa-specific PNA probe, a fluorescein-labelled Staphylococcus aureus-specific PNA probe and a fluorescein-labelled or Texas-red-labelled universal bacterium PNA probe in hybridization solution (AdvanDx) was added to each section and hybridized in a PNA FISH
workstation at 55 °C for 90 min covered by a lid. The slides were washed for 30 min at 55 °C in Wash Solution (AdvantDx), mounting medium with 4',6-diamidino-2-phenylindole (DAPI) was applied, and a coverslip was added to each slide. The entire PNA FISH procedure took approximately 2.5 h. Slides were read using a fluorescence microscope equipped with an FITC, a Texas red, a dual FITC/Texas red and a DAPI filter.

RESULTS

To investigate the importance of rhamnolipid production for PMN viability in vivo, we infected lungs of mice with equal amounts of rhamnolipid-proficient (PAO1 wild-type) and -deficient (rhlA mutant) P. aeruginosa strains. Our alginate bead model of infection features introduction of bacteria in the form of aggregates, and previous microscopic investigations have revealed that the bacteria organize in biofilm aggregates surrounded by multiple PMNs similar to those in the lungs of cystic fibrosis (CF) patients and chronic infected wounds (Wu et al., 2000). After 24 h, flow-cytometry-based live–dead stain analysis (see Methods) showed that the BAL fluid obtained from the group of mice infected with the rhlA mutant contained twofold more viable PMNs (P<0.0001, t-test) compared with the PAO1 wild-type-infected group (Fig. 1a). Overnight, we also found one order of magnitude higher numbers of c.f.u. in the lungs of wild-type-infected mice compared with a two orders of magnitude reduction in those infected with the rhlA mutant (P<0.0001, Mann–Whitney test) (Fig. 1b). PMNs are summoned to the lungs upon infection and have a high turnover rate. However, with respect to wild-type bacteria, the present 24 h ‘snapshot’ measurements suggest that rhamnolipid production (Fig. 1c) may account for the elimination of a significantly higher fraction of the attracted PMNs in contrast to the rhlA-deficient bacteria, which become eliminated by the action of the PMNs. This view was strengthened by in vitro studies of biofilms in flow tubes. Measurement of the LDH activity of in vitro biofilms exposed to freshly isolated PMNs revealed that a large fraction of the PMNs exposed to the PAO1 wild-type biofilm lysed during the period of the experiment, whereas no PMNs lysed after exposure to the rhlA mutant (Fig. 2c).

In line with Morici et al. (2007), we were only able to detect very low amounts of rhamnolipid in the in vitro biofilm of P. aeruginosa. We then hypothesized that the rhamnolipid content might rise and spike in the biofilm under conditions of PMN exposure. In order to prove this hypothesis, we measured the amount of rhamnolipids after 2 h of exposure to freshly isolated PMNs. Interestingly, when comparing with non-exposed biofilms, we found a significant increase in rhamnolipid concentration (P<0.003, t-test) in the cell fraction of the harvested biofilms (Fig. 2a). As expected, we did not find any detectable amounts of rhamnolipids in the rhlA-deficient strain before or after exposure to the PMNs (Fig. 2a). In addition, we found a more than 2000-fold higher rhamnolipid concentration in the biofilm cell paste than in the surrounding fluid (P<0.0001, t-test), showing that the de novo-synthesized rhamnolipids stick to the biomass rather than being released to the fluid (Fig. 2b). We therefore propose that PMNs become eliminated by contact with the biofilm aggregates, in accordance with our observations that microscopic inspections cannot

![Fig. 1. Intratracheal instillation of alginate-embedded P. aeruginosa in BALB/c mice. (a) After 24 h of infection there is a significantly higher proportion of dead PMNs in the BAL fluid of BALB/c mice infected with the PAO1 wild-type infected mice (●) compared to the corresponding rhlA mutant-infected mice (▲). (b) There is a significant difference (P<0.0001, Mann–Whitney test) in bacteriology after 24 h in the lungs of mice infected with PAO1 wild-type and the corresponding rhlA mutant (▲). The horizontal bars represent mean percentage dead PMNs. Each group contained 12 mice at the day of sacrifice (P<0.0001, t-test). (b) There is a significant difference (P<0.0001, Mann–Whitney test) in bacteriology after 24 h in the lungs of mice infected with PAO1 wild-type and the corresponding rhlA mutant (▲). Immediately after infection four mice were sacrificed to estimate the inoculum size (at 0 h). The horizontal bars represent median c.f.u. numbers. (c) Rhamnolipid concentration in BAL fluid from mice infected with PAO1 wild-type (●) and the corresponding rhlA mutant (▲). The horizontal bars represent median rhamnolipid concentration.
demonstrate the presence of intact PMNs in close contact with the top layers of the in vitro biofilms (Fig. 3c).

Numerous microscopic investigations of P. aeruginosa-infected ex vivo tissue samples from the lungs of CF patients and chronic wounds (Fig. 3a, b) gave the impression that PMNs may gather in the periphery but never reach inside the biofilm cell aggregates. Furthermore, the biofilm aggregates are surrounded by blue-stained smears, which may represent DNA leaking out of nearby, lysed PMNs. Biofilm-associated rhamnolipids therefore appear to be a feasible explanation for biofilm resistance to the cellular immune defence.

Taken together, the present study suggests that the biofilm bacteria can sense the presence of PMNs and respond accordingly. Consequently, we should be able to see changes in the transcriptomic profile of the exposed bacteria in vitro. By means of Affymetrix GeneChips we calculated absolute expression values obtained from PMN-exposed biofilms compared to the corresponding values of a non-exposed control. The changes in gene expression were calculated as simple fold changes. In line with our rhamnolipid measurements, we found that among 27 other genes there was a significant (P<0.03 and P<0.01) upregulation of the rhlA and rhlB genes during PMN exposure (Table 1), with very low expression under conditions of no exposure. We validated the change in transcription of rhlA, pqsA and rpoD found by the GeneChips by RT-PCR and obtained comparable results (See Supplementary Table S2).

Rhamnolipid-dependent PMN killing appears to be QS regulated (Jensen et al., 2007). When comparing with data presented by Hentzer et al. (2003) and Rasmussen et al. (2005), 85% of the PMN-responsive genes are also regulated by QS. In particular, those virulence factors encoded by phz, rhlAB and lecB are able to cripple and eliminate eukaryotic cells (Allen et al., 2005; Jensen et al., 2007; Sonawane et al., 2006). These genes have been shown to be controlled by the PQS branch of the QS system.
The biosynthetic pqs operon is also upregulated during PMN exposure, suggesting an increase in PQS signal production. Concurrently, we found in the biofilm set-up that the pqsA mutant is unable to respond by increasing rhamnolipid production upon exposure to PMNs (Fig. 2a). The PQS signal is known to stimulate the RhlR-C4-HSL-controlled lower part of the QS hierarchy through PqsR and PqsE. It is therefore likely that the cells in a P. aeruginosa biofilm channel the PMN response by means of the PQS branch of the QS system. In support of this is the finding that a QS-deficient (ΔlasRΔrhlR) biofilm showed a completely different response, i.e. no upregulation of QS-controlled genes and virulence factors (Table 1).

Comparison with a previous analysis showed that 70% (19 of 27) of the present PMN-upregulated genes were also upregulated in a ΔalgR mutant biofilm studied by Morici et al. (2007) (genes also upregulated in a ΔalgR mutant biofilm are indicated in Table 1). Since AlgR is known to repress the PQS-controlled genes in biofilms, it is tempting to speculate that a cue from the PMNs overrides AlgR repression and thereby induces the observed virulent response. Evidence has been provided by Zaborina et al. (2007) that the opioid and endogenous κ-agonist dynorphin A induces production of PQS and phenazines and this induction requires functional QS systems. The group demonstrated that 100 μM dynorphin A (1-17) is required to enhance the production of virulence factors in a planktonic P. aeruginosa culture (Zaborina et al., 2007). Since immune cells are capable of synthesis and release of opioids at sites of inflammation (Cabot et al., 2001; Chadzinska et al., 2005; Przewlocki et al., 1992) it is an appealing hypothesis that these peptides are in fact the signals by which P. aeruginosa biofilms recognize the presence of incoming and attacking PMNs.
**Table 1.** GeneChip transcriptomic analysis of 3-day-old *in vitro* *P. aeruginosa* biofilms exposed to PMNs for 2 h

By comparison with unexposed biofilms, we found that 28 genes were significantly (>2-fold change and 3-fold regulated) differentially expressed. Results are based on the mean of three separate experiments. Of these, 85% are QS controlled according to Hentzer et al. (2003); these QS-controlled genes are identified in the ‘QS controlled’ column. Genes found to be AlgR repressed in biofilms by Morici et al. (2007) are indicated in the ‘AlgR repressed’ column. We also included a QS-deficient Δ*rhl*Δ*lasR* biofilm in the transcriptome analysis. The response to PMN exposure was completely different from that of the isogenic QS-proficient strain, with only two genes being significantly upregulated.

<table>
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<th>Probe set ID</th>
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<td>PA4210</td>
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<td>Δ<em>rhlΔlasR</em></td>
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We decided to challenge this hypothesis by exposing a 3-day-old biofilm to 100 μM dynorphin A (1-17) for 30 min or 120 min. After exposure we measured the rhamnolipid concentration in the biofilm as previously described. In comparison to the non-treated biofilm no induction of rhamnolipid was observed (Fig. 4). Microarray analysis of the harvested biofilms revealed 225 and 13 genes, respectively, being differentially (>3-fold) regulated following 30 min and 120 min of dynorphin exposure. Consistent with the rhamnolipid measurement, we were unable to detect any induction of PQS-controlled transcription, including the *rhlAB* genes. These data indicate that dynorphin A is not the cue from the PMNs that activates the PQS system and makes the bacteria launch the protective shield.

**DISCUSSION**

*P. aeruginosa* uses QS communication systems to trigger the onset of virulence factor production. The production of virulence factors contributes to activation of the immune system and PMNs will be summoned to the sites of infection. In the present study we show that the presence of PMNs triggers the *P. aeruginosa* biofilm to respond aggressively by producing a QS-dependent cocktail of toxic components rather than protective mechanisms such as those involved in oxidative stress (Chang et al., 2005; Small et al., 2007). This corroborates the previous finding that freshly isolated PMNs are unable to develop an oxidative burst and phagocytose *in vitro* *P. aeruginosa* biofilm (Bjarnsholt et al., 2005).

Among the upregulated substances are rhamnolipids, which stick to the biofilm, forming a deadly shield surrounding the biofilm cells. Approaching PMNs are consequently killed by contact and the bacteria persist. This is further supported by the results of Kownatzki et al. (1987), who found that sputum from *P. aeruginosa*-infected CF patients contained rhamnolipid and correlated high concentrations with poor clinical state.

Why is rhamnolipid production simply not ‘on’ at all times? This may be due to the fact that rhamnolipid-induced necrosis is not limited to PMNs or macrophages...
but affects all eukaryotic cells surrounded by a plasma membrane. We propose that too much rhamnolipid may cause inflammation and host tissue damage to levels beyond those favourable for the persistence of \textit{P. aeruginosa} and therefore a tight regulation must take place at all times. Furthermore, the QS signals 3-oxo-C12 HSL and PQS have been reported to change the maturation pattern of stimulated dendritic cells away from a proinflammatory T-helper type 1 directing response, thereby decreasing the antibacterial activity of the adaptive immune defence (Skindersoe \textit{et al}., 2009). The present study supports a model by which cross-kingdom-based communication contributes significantly to immunomodulation and evasion that facilitate the infective properties of \textit{P. aeruginosa}.

This model suggests the potential for harnessing the power of small molecules that block bacterial QS systems as future antimicrobial medicines. A comparison with data from Hentzer \textit{et al.} (2003) showed that 77\% of the genes induced by exposure to PMNs can be significantly repressed by the QS blocker furanone C-30. This experimental drug is capable of blocking the aggressive anti-PMN response. Consequently, it restores the proper action of the attracted PMNs and strongly promotes bacterial clearance in animal models of infection (Christensen \textit{et al}., 2007; Hentzer \textit{et al}., 2003). Further study of interactions between \textit{P. aeruginosa} and components of the innate and adaptive immune system will provide valuable information for the design of novel antimicrobials aimed at stimulating the efficiency of and restoring an adequate host defence.

ACKNOWLEDGEMENTS

This work was supported by grants from the Danish Research Council for Technique and Production (FTP) and the German Mukoviscidose ev to M. G. T. B. has received financial support from the Carlsberg Foundation and Lundbeck Foundation ‘The implication of bacterial ev to M. G. T. B. has received financial support from the Carlsberg Foundation and Lundbeck Foundation ‘The implication of bacterial communication on biofilm persistence in an in vivo body infection model.’

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Edited by: P. Cornelis