Residence in biofilms allows *Burkholderia cepacia* complex (Bcc) bacteria to evade the antimicrobial activities of neutrophil-like dHL60 cells

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One sentence summary: A study of the influence 'Burkholderia cepacia complex' biofilms have on the behavior and fate of neutrophil-like dHL60 cells, also addressing the influence of those cells on the biofilm.

ABSTRACT

Bacteria of the *Burkholderia cepacia* complex (Bcc) persist in the airways of people with cystic fibrosis (CF) despite the continuous recruitment of neutrophils. Most members of Bcc are multidrug resistant and can form biofilms. As such, we sought to investigate whether biofilm formation plays a role in protecting Bcc bacteria from neutrophils. Using the neutrophil-like, differentiated cell line, dHL60, we have shown for the first time that Bcc biofilms are enhanced in the presence of these cells. Biofilm biomass was greater following culture in the presence of dHL60 cells than in their absence, likely the result of incorporating dHL60 cellular debris into the biofilm. Moreover, we have demonstrated that mature biofilms (cultured for up to 72 h) induced necrosis in the cells. Established biofilms also acted as a barrier to the migration of the cells and masked the bacteria from being recognized by the cells; dHL60 cells expressed less IL-8 mRNA and secreted significantly less IL-8 when cultured in the presence of biofilms, with respect to planktonic bacteria. Our findings provide evidence that biofilm formation can, at least partly, enable the persistence of Bcc bacteria in the CF airway and emphasize a requirement for anti-biofilm therapeutics.

Keywords: cystic fibrosis; host-pathogen interaction; innate immunity

INTRODUCTION

People with cystic fibrosis (CF) generate a dehydrated and unusually viscous pulmonary mucous which, in turn, diminishes mucociliary clearance from their airways (Boucher 2004). This allows for inhaled microorganisms to remain in situ and colonize the airway, transiently or permanently, depending in part on their ability to subvert the immune response.

Among the many opportunistic microorganisms which can colonize the airways of people with CF are members of the *Burkholderia* genus of Gram-negative bacteria, particularly the 18 closely related species currently considered part of the *Burkholderia cepacia* complex (Bcc) (Vandamme and Dawyndt 2011; Peeters et al. 2013). Presently, the incidence of Bcc bacteria in CF patients is approximately 2.6% in the USA and 3.8% in the UK (CFF 2012; UKCFR 2013), with *B. cenocepacia* and *B. multivorans* strains comprising the majority of isolates (Drevinek and Mahenthiralingam 2010).

Though less common colonizers of CF patients than *Pseudomonas aeruginosa*, Bcc bacteria are disproportionately highly associated with morbidity and mortality amongst sufferers...
(Tablan et al. 1987; Jones et al. 2004), owing to the bacteria’s induction of an acute-onset lung deterioration with associated septic bacteraemia, termed ‘cepacia syndrome’ (Isles et al. 1984), as well as its correlation with poorer post-operative outcomes following lung transplantation (Alexander et al. 2008; Meachery et al. 2008). Pathogenic strains of Bcc species are also commonly multidrug resistant and most display an ability to form biofilms in vitro (Caraher et al. 2007), which further reduce their sensitivity to antibiotics.

Over the duration of pulmonary colonization, P. aeruginosa displays a number of adaptive phenotypes which aid in its persistence (Sousa and Pereira 2014). Among these adaptations is the formation of a biofilm, largely comprised of secreted exopolysaccharide (EPS) and DNA, often detected in the CF airway as non-surface-attached bacterial aggregates (Singh et al. 2000; Srimulu et al. 2005).

Biofilm formation by P. aeruginosa is known to facilitate their evasion of the antimicrobial actions of neutrophils. The biofilm can impede migration of leukocytes at their exofacial surface (Jesaitis et al. 2003), while mucoid P. aeruginosa is resistant to phagocytosis by neutrophils relative to non-mucoid counterparts (Cabral, Loh and Speert 1987), by virtue of cell surface alteration which renders the bacterium unrecognizable.

A similar situation may prevail for Burkholderia, a close relative of Pseudomonas; mucoid CF isolates of B. cenocepacia display reduced adhesion to neutrophils relative to similar, non-mucoid strains (Conway et al. 2004). Indeed, biofilm-deficient mutants of P. aeruginosa are more susceptible to phagocytosis by neutrophils than their biofilm-competent counterparts (Bjarnsholt et al. 2005).

Following a period of antimicrobial activity, neutrophils undergo secondary necrosis and shed their cellular components which become part of the exopolymyleric matrix of the biofilm (Parks et al. 2009), serving to further enhance its biomass while preserving the antibiotic resistance characteristic of the biofilm (Caceres et al. 2014). Indeed, clinical observations of P. aeruginosa biofilm aggregates highlight their close association with large quantities of, mostly non-viable, neutrophils (Bjarnsholt et al. 2009).

In studies of the CF airways, Bcc bacteria have predominantly been observed either as singular, planktonic cells or surviving within epithelial cells or leukocytes (Sajjan et al. 2001; Lamothe et al. 2007; Schwab et al. 2014), rather than biofilm-like aggregates. However, the heterogeneity of the airways makes it clear that prohibitively comprehensive sampling of the airways would be required to definitively conclude that Bcc biofilms are absent or present. Accordingly, other investigators suggest that Bcc biofilms exist in the airways of CF patients (Ciofu et al. 2015).

Given the utility of the biofilm to P. aeruginosa in evading neutrophil antimicrobial activity and the relatedness of Burkholderia to Pseudomonas, we hypothesize that the formation of a biofilm is important to the Bcc species’ resistance to neutrophil-mediated killing, considering that they are capable of persistence in CF patients’ airways, despite the continuous presence of large numbers of neutrophils in the lungs of CF patients. This report provides support for this hypothesis by elucidating the nature of Bcc species’ interaction with neutrophil-like dHL60 cells.

We have investigated the bacteria’s interaction with the cells from two broad standpoints, namely from the perspective of cell functionality and survival in the presence of biofilm-dwelling Bcc bacteria and the influence of the presence of the cells on the formation and development of biofilms by Bcc species.

We report that chemotaxis, migration and secretion of chemokines are diminished in the cells, while their viability is reduced following culture in the presence of Bcc biofilms, with respect to planktonic bacteria. Concurrently, the biofilm formed by the bacteria is enhanced during this coculture. Our findings contribute to our understanding of the respective effects that neutrophils and Bcc biofilms have on one another in the CF airway and emphasize the utility of the biofilm to Bcc bacteria as a virulence trait in CF.

**EXPERIMENTAL PROCEDURES**

**Bacterial strains**

Bcc bacteria had been obtained from the Belgian Co-ordinated Collections of Microorganisms (BCCM/LMG) and were stored at −20°C. The strains used were B. multivorans LMG 13010, B. dolosa LMG 18941 and B. cenocepacia K56–2 (LMG 18863), each of which had been isolated from CF patients prior to deposition with the BCCM. For use during this study, bacteria were maintained on B. cepacia selective agar (Henry et al. 1997) and cultured in LB medium (Sigma) at 37°C.

**Cell culture**

The promyelocytic cell line, HL60, was obtained from the European Collection of Cell Cultures (Health Protection Agency Culture Collections) and cultured in IMDM (Lonza) supplemented with 10% (v v−1) FBS (Sigma) at 37°C with 5% CO2. Cells were differentiated by culture in the presence of 1.25% (v v−1) DMSO (Sigma) for 7 days prior to use (Collins et al. 1979). Differentiation was confirmed by means of flow cytometry for the expression of neutrophil-specific markers, CD66 and CD11b (Owaki et al. 1991).

**Assessment of biofilm biomass**

Bacteria (1 × 10⁶ cfu ml−1) were cultured in the absence or presence of viable dHL60 cells (1 × 10⁷ cells ml−1) or an equivalent amount of whole-cell lystate in 100 μl LB in wells of 96-well plates at 37°C for up to 72 h. Lysates were prepared by repeated freeze–thaw cycles at −80°C with mechanical disruption through a fine gauge syringe needle, so as to avoid the use of detergents. Lysis was confirmed by visual inspection of discrete 20 μl aliquots using a light microscope at 400× total magnification. DNA was extracted from dHL60 cells using the DNeasy blood and tissue kit (Qiagen), as per the manufacturers’ protocol. Wells were washed three times with deionized water and biofilm biomass was stained by addition of 125 μl of an aqueous 1.25% (w v−1) crystal violet solution (Sigma) for 45 min. Wells were then washed further three times and biofilm-bound crystal violet was solubilized in 200 μl 95% (v v−1) ethanol/0.05% triton X-100 (Sigma). Crystal violet was quantified spectrophotometrically at 590 nm using a Varioskan microtiter plate reader (Thermo Scientific).

In parallel, dHL60 cells, or lysates thereof, were cultured in triplicate wells in the absence of bacteria. These wells were similarly subjected to crystal violet staining and revealed negligible adsorption of cellular material (mean OD₅₉₀ = 0.102 ± 0.14; n = 3).

**Assay of IL-8 secretion**

Bacteria were cultured in LB to mid-exponential phase of growth and inoculated (1 × 10⁶ cfu ml−1) into triplicate wells of 24-well plates prior to incubation at 37°C for 24, 48 or 72 h. Biofilms were washed with sterile, pre-warmed PBS and dHL60 cells (1 × 10⁷ cells ml−1) were added in IMDM. Planktonic bacteria, cultured to
mid-exponential growth in LB, were discretely diluted to each of 1 × 10^6 or 1 × 10^7 cfu ml⁻¹ in IMDM and inoculated separately into triplicate wells of 24-well plates in the presence of dHL60 cells. Supernatants were recovered following 3 h of coculture at 37 °C and stored at −20 °C. Levels of IL-8 in 1:10 dilutions of supernatants were determined by sandwich ELISA. ELISA was conducted using the OptiELISA IL-8 ELISA Set (BD Biosciences) according to the manufacturers’ protocol.

Enumeration of bacteria residing within biofilms

Bacteria residing within the biofilms cultured in wells of microtiter plates were enumerated according to the method described by Van den Driessche et al. (2014). Briefly, bacteria (1 × 10^6 cfu ml⁻¹) were cultured in 100 μl LB in triplicate wells of 96-well plates at 37 °C for 24, 48 or 72 h. Wells were then rinsed three times with Ringers solution, 100 μl of Ringers solution was added and biofilm was disrupted by repeated pipetting followed by sonication in an ultrasonic bath (Branson Ultrasonics Corp.) for 5 min. Contents of wells were then subjected to a series of 10-fold dilutions in Ringers solution. Approximately 20 μl aliquots of these dilutions were plated in triplicate and plates were incubated at 37 °C for 48 h. Colonies formed on each plate were then enumerated where possible. The experiment was conducted in triplicate.

Confocal microscopy

For observation by confocal microscopy, bacteria were transfected, by triparental mating (de Lorenzo and Timmis 1994), with plasmid, pBAH 8 (Huber et al. 2002), which encoded mutated green fluorescent protein (GFPmut3) (Cormack, Valdivia and Falkow 1996). The donor strain used for triparental mating was Escherichia coli MT102, which had been cultured in the presence of 10 μg ml⁻¹ gentamycin (Sigma). The helper strain used was E. coli HB 101, having the plasmid pRK600, which had been cultured in the presence of 10 μg ml⁻¹ chloramphenicol (Sigma). Donor and helper strains were kindly donated by Prof. Leo Eberl (Institute of Plant Biology, University of Zurich, Switzerland).

Transfected bacteria were cultured under continuous flow of minimal nutrients as described in detail elsewhere (Weiss Nielsen et al. 2011). Briefly, bacteria (1 × 10^6 cfu per chamber) were inoculated in LB into triplicate flowcell chambers and allowed to adhere to the glass slide component of the chamber in the absence of medium throughput for 45 min. Following this, bacteria were incubated at 37 °C under a flow of 0.1 ml min⁻¹.

CellTracker Red fluorescent cytoplasmic dye (Molecular Probes) was used to stain dHL60 cells (4 × 10^5 cells) following the manufacturers’ protocol 30 min prior to addition to flowcell chambers in the absence of media throughput. Confocal micrographs were generated using an Olympus FV-1000 microscope with a 60× oil immersion lens (Olympus). Computer-generated surfaces were created using Imaris software (Bitplane).

Assessment of cell death

Bcc bacteria were cultured to mid-exponential phase of growth and diluted to 1 × 10^6 cfu ml⁻¹ in LB for inoculation into triplicate wells of 24-well plates. Plates were cultured at 37 °C for 24, 48 or 72 h. Wells were then rinsed twice with pre-warmed, sterile PBS prior to addition of 1 × 10^7 dHL60 cells ml⁻¹ in IMDM. Bacteria were separately cultured to mid-exponential growth phase in LB and diluted to 1 × 10^5 cfu ml⁻¹ in IMDM immediately prior to addition of dHL60 cells (1 × 10^7 cells ml⁻¹).

Supernatants were also prepared from discrete cultures of planktonic or biofilm-dwelling bacteria (24, 48 and 72 h) by careful removal of liquid without disturbance of biofilm, centrifugation at 4000 × g for 10 min and filtration through syringe filters having a pore size of 0.22 μm. dHL60 cells were then inoculated into supernatants in triplicate wells. Plates were then incubated for 3 h at 37 °C.

Additional cells were discretely incubated for 3 h in the presence of known inducers of apoptosis. Actinomycin D (2 μg ml⁻¹) was used to induce apoptosis for assessment of caspase-3 activation. Staurosporine (15 μM) was used as an inducer of apoptosis during assessment of Mcl-1 expression to avoid interference from the mRNA transcription inhibitory property of actinomycin D.

Quantification of caspase-3 activation was performed using the caspase-3/7 colorimetric assay kit (Biovision) according to the manufacturers’ instructions.

For flow cytometry, cells were removed, washed with Annexin V binding buffer and stained with annexin V-FITC and propidium iodide (BD Biosciences). Flow cytometry was performed on a FACSCalibur instrument (BectonDickinson).

Gene expression analysis by RT-qPCR

The relative expression of IL-8 or Mcl-1 mRNA transcripts was determined for dHL60 cells (1 × 10^7 cells ml⁻¹) which had been incubated at 37 °C in triplicate wells of 24-well plates in the presence of Bcc bacteria for 3 h. The bacteria had been cultured planktonically to mid-exponential phase and diluted to 1 × 10^6 cfu ml⁻¹ in 1 ml IMDM or inoculated at 1 × 10^6 cfu ml⁻¹ into triplicate wells of 24-well plates prior to culture as biofilms for 24, 48 or 72 h. Biofilms were rinsed twice with sterile PBS prior to addition of dHL60 cells in 1 ml IMDM.

Total RNA was isolated using the RNeasy kit (Qiagen), followed by removal of residual DNA by use of the TURBO DNA-free kit (Ambion). Reverse transcription was carried out using the Superscript® VILO cDNA synthesis kit (Invitrogen). Quantification of mRNA was conducted on an ABI 7300 real-time PCR system (Applied Biosystems) using Power SYBR® green PCR master mix (Applied Biosystems). Primer sequences were either obtained from PrimerBank or were designed using Primer-BLAST software (NIH); TBP Primer-Bank ID: 285026518c3, Mcl-1 PrimerBank ID: 11386165a2, SDHA forward sequence: 5′-GGGCAACAGCAGACAT-3′, SDHA reverse: 5′-ACTGTTGGCCACGCCCTT-3′, IL-8 forward: 5′-ACACTGGCCAAACACAGAAAA-3′, IL-8 reverse: 5′-ATTCTCAGCCCTCTTCAAAACTCTC-3′. Primers were obtained from Invitrogen. Cq values and average efficiencies per gene were determined by LinRegPCR (Ruijter et al. 2009). Differential gene expression between cell populations was then estimated using REST 2009 software (Qiagen) (Pfaffl, Horgan and Dempfle 2002; Hellemans et al. 2007), normalized against the geometric mean expression of the reference genes TBP and SDHA, previously validated as stable in neutrophils exposed to LPS (Ledderose et al. 2011). The geometric mean of the fold changes and their respective standard errors were calculated (Bengtsson et al. 2005) and means were then subjected to log₂ transformation (Rieu and Powers 2009).

Statistical analysis of data

Parametric data were subjected to Student’s two-independent-sample t-test (Student 1908). Comparisons of multiple discrete populations were conducted by one-way analysis of
Figure 1. dHL60 cells do not enter established B. multivorans LMG 13010 and B. dolosa LMG 18941 biofilms. Bcc bacteria (1 × 10^6 cfu ml^-1), harboring a GFP-expression plasmid, were inoculated into flowcells and cultured under continuous throughput of defined, minimal medium at 37°C for 72 h. To the flowcell was added 4 × 10^5 dHL60 cells ml^-1, which had been stained with 5 μM Celltracker Red cytoplasmic dye (Molecular Probes), 30 min prior to image collection. Composite confocal micrographs of sections through the z-axial plane of biofilms formed by (A) B. dolosa LMG 18941 and (D) B. multivorans LMG 13010 were then generated, with biofilms proximal to the viewer. (B) Enlargement of a single micrograph of the area demarcated by white box in A showing a phagocytosing cell. (E) Enlargement of the area demarcated by white box in D to highlight (F) a transverse section corresponding to image E illustrating a cell resting above the biofilm. Using the data presented in A and D, discrete surfaces were overlaid onto the detected fluorescence intensities for each channel using Imaris software (Bitplane) in order to illustrate the spatial relationship between (C) B. dolosa and (G) B. multivorans biofilms, and dHL60 cells.

RESULTS
The Bcc biofilm is a physical barrier to dHL60 cell migration
Neutrophils have been shown to migrate through staphylococcal biofilms and to phagocytose the biofilm-dwelling bacteria (Leid et al. 2002; Günther et al. 2009a,b; Meyle et al. 2010). However, Jesaitis et al. (2003) have shown that neutrophils are incapable of migration into established P. aeruginosa biofilms, hence limiting their ability to phagocytose and eradicate the biofilm-dwelling P. aeruginosa bacteria. Thus, the failure of neutrophils to eradicate colonizing Bcc bacteria in the CF airway may be due, at least in part, to their inability to effectively gain access to the bacteria residing in biofilms. To determine whether this was the case, Bcc bacteria were cultured in flowcell chambers in order to generate biofilms amenable to visual inspection. dHL60 cells were then added and their position assessed by confocal microscopy. By 30 min post-addition of dHL60 cells, they demonstrated a rounded morphology and were residing superficially on the luminal surface of the biofilm matrix (Fig. 1A). In most cases, bacteria had swarmed around the cells and the cells had internalized bacteria in some cases (yellow–green coloration, Fig. 1B). A lack of lateral movement or ingress was apparent, with cells residing superficially on the biofilm exterior (Fig. 1C, F and G). This was particularly evident for B. multivorans biofilms of high biomass (Fig. 1D), suggesting that the biofilm physically impedes the migration of the cells into the biofilm.

The Bcc biofilm masks the bacteria from being recognized by dHL60 cells
Pulmonary inflammation in CF is largely driven by excessive activation of infiltrated neutrophils, correlating with high levels of proinflammatory cytokines and chemokines (Bonfield et al. 1995). As such, we sought to determine whether the biofilm
promoted an excessive secretion of neutrophil-recruiting and -activating IL-8. Surprisingly, we observed a significant suppression in the amount of IL-8 secreted by dHL60 cells in response to Bcc biofilms, relative to planktonic Bcc bacteria, for each of the three species investigated (P < 0.001; Fig. 2 A–C). This suggests that the biofilm is masking the bacteria within from being recognized by the dHL60 cells. Furthermore, the amount of IL-8 secreted in response to the biofilms showed little intraspecific variation regardless of whether the biofilm was 24, 48 or 72 h old. This implies that by 24 h post-inoculation, the Bcc biofilm is sufficiently robust to diminish recognition of the bacteria by the dHL60 cells. This finding was corroborated by our observation of diminished induction of IL-8 expression in dHL60 cells cultured with biofilms of *B. multivorans* or *B. dolosa* bacteria, with respect to those cells that were cultured in the presence of planktonic bacteria (Fig. 2 D and E).

In order to ascertain whether numbers of bacteria residing within biofilms influenced the observed IL-8 secretion, the numbers of colony-forming units present in the wells at 24, 48 and 72 h under matching conditions were determined by biofilm disruption and bacterial enumeration by spread plate method. Overall, mean cfu per well was 2.57 ± 1.54 × 10^7 cfu. Quantities observed did not correlate significantly with either bacterial species or duration of culture (logistic regression, P = 0.659).

Hence, the IL-8 secreted by dHL60 cells upon exposure to Bcc biofilms was markedly less than would be expected in response to an equivalent quantity of planktonic bacteria.

**Exposure to biofilms enhances necrosis in dHL60 cells**

The proinflammatory nature of neutrophils in responding to invading bacteria is resolved in large part by their ingestion by macrophages. Neutrophils to be ingested are recognized by cell-surface markers of apoptosis which themselves are products of phagocytosis-induced cell death (PICD) (Coxon et al. 1996; Flannagan et al. 2014). CF airways, however, are marked by a continuous inflamed state, indicating this process is not occurring correctly.

Hence, we sought to ascertain whether biofilm-dwelling Bcc bacteria disrupted the normal onset of apoptosis in dHL60 cells. Assay of the activation of caspase-3 revealed that dHL60 cells, which had been cultured in the presence of Bcc biofilms established 24, 48 or 72 h previously, were undergoing apoptosis and that the extent of apoptosis in the cell population was proportional to the maturity of the biofilm to which they were exposed (Fig. 3A), while planktonic bacteria induced little caspase-3 activation during the course of coculture. This outcome may be explained by increased numbers of bacteria yielding a corresponding increase in phagocytosis and, hence, PICD. However, this supposition was not corroborated by our findings following enumeration of bacteria within our cultured biofilms. While biofilms harbored 2.57 ± 1.54 × 10^7 cfu—markedly higher than the experimental inoculum of planktonic bacteria—there was no consistent correlation between biofilm maturity and bacterial count for any of the species tested.
Figure 3. Culture in the presence of Bcc biofilms induces cell death in dHL60 cells. Bacteria ($1 \times 10^6$ cfu ml$^{-1}$) were cultured at 37°C for 0, 24, 48 or 72 h. Wells were then washed with sterile PBS and dHL60 cells ($1 \times 10^7$ cells ml$^{-1}$) were added. Discrete dHL60 cell populations were also cultured in the absence or presence of 2 ug ml$^{-1}$ actinomycin D or 15 μM staurosporine. Plates were incubated at 37°C for 3 h prior to harvesting of cells. Cells were lysed and assayed for caspase-3 activity (A). Columns represent the mean caspase-3 activity of triplicate dHL60 cell populations from each of three independent experiments, relative to the mean uninduced control population. Bullets indicate statistically significant differences with respect to uninduced controls ($^*P < 0.05$, $^{**}P < 0.01$, $^{***}P < 0.001$, one-way ANOVA). Asterisks denote significant differences between biofilm age for a given species ($^{*}P < 0.05$, $^{**}P < 0.01$, $^{***}P < 0.001$). Cells from each population were then isolated by centrifugation and resuspended in binding buffer in the presence of annexin V-FITC and propidium iodide. Each cell population was analyzed by flow cytometry. Representative scatterplots are presented for dHL60 cells which had been cultured with (B–E) supernatants derived from planktonic or biofilm-dwelling populations of B. multivorans or (F–I) the planktonic or biofilm-dwelling B. dolosa bacteria themselves. The quantity of viable, apoptotic or necrotic cells following each culture condition is also presented for (J) supernatants and (K) planktonic or biofilm-dwelling bacteria. RNA was isolated from cells and converted to cDNA; cDNA was then analyzed semiquantitatively by qPCR (L–N). Data represent the log$_2$-transformed geometric means ($\pm$ SE) of triplicate populations from each of three independent experiments. Bullets indicate significance differences mediated by biofilms vs planktonic bacteria ($^*P < 0.05$, $^{**}P < 0.01$, $^{***}P < 0.001$, Kruskal–Wallis test), asterisks indicate significance mediated by biofilms vs one another ($^*P < 0.05$, $^{**}P < 0.01$, $^{***}P < 0.001$).
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Figure 4. The impact of dHL60 cells on the formation of biofilms by Bcc bacteria. Bacteria (1 × 10⁵ cfu) were cultured in 96-well plates in the absence or presence of (A–C) dHL60 cells (1 × 10⁶ cells), (D–F) an equivalent quantity of whole-cell lysate thereof, or (G–I) DNA derived from an equivalent number of cells, for up to 72 h. In some cases, additional dHL60 cells or cellular material were added 24 h after initial inoculation. Biofilm density was assessed after 24, 48 or 72 h by crystal violet staining. Asterisks indicate significant differences between normal biofilm biomass and biofilm formed in the presence of dHL60 material (∗*P < 0.05, ∗∗∗P < 0.001, two independent sample t-test). Bullets indicate significant differences comparison between normal biofilm biomass and biomass after two additions of dHL60 material (rP < 0.05, rP < 0.01, rrrP < 0.001). Crosses indicates significant differences between biofilm formed in the presence of dHL60 material vs biomass after two additions of dHL60 material (†P < 0.05, ††P < 0.001). Data are expressed as the mean (± SD) of four biological replicates from each of three independent experiments.

However, we then used flow cytometry to discern the extent and progression of cell death to secondary necrosis. This demonstrated that substantial proportions of the cell populations which had been cultured with well-established B. dolosa LMG 18941 static biofilms (72 h maturity) had progressed to necrosis by 3 h post-inoculation (Fig. 3I). This outcome was not elicited by supernatants derived from the medium in which biofilms had been cultured (Fig. 3E), suggesting that the dHL60 cell death was not mediated by a soluble factor, but rather a factor associated with the biofilm itself or the nature of the interaction.

Furthermore, the loss of the cells’ viability during culture with Bcc biofilms coincided with the loss of Mcl-1 expression, required for neutrophil survival (Fig. 3L–N).

Coculture of Bcc bacteria with dHL60 cells results in increased biofilm formation

Considering the advantage that biofilm residence confers on Bcc bacteria in limiting neutrophil antimicrobial activities, we sought to ascertain whether dHL60 cells had an impact on the extent of biofilm formed by Bcc bacteria. To determine whether dHL60 cells had an effect on the extent of biofilm formation and development by Bcc species, bacteria were cultured in the presence of dHL60s for up to 72 h. Staining of the biofilm biomass revealed that B. dolosa biofilms possessed significantly greater biomass following culture in the presence of dHL60s than in their absence (P < 0.001; Fig. 4B). This outcome was further pronounced when B. dolosa LMG 18941 bacteria were cultured in the presence of whole-cell lysates of dHL60 cells (P < 0.001; Fig. 4E), a finding mirrored by B. cenocepacia bacteria (Fig. 4F). This suggests that, while viable dHL60s kill some proportion of the bacterial population, once they have become necrotic and disintegrated, their cellular components become incorporated into the bacterial EPM.

Culture of the bacteria with DNA purified from dHL60 cells highlighted that this alone could enhance the quantity of biofilm biomass (Fig. 4H), though this effect was not as pronounced. Burkholderia multivorans LMG 13010 bacteria displayed significantly increased biomass with respect to controls following 48 h of culture after two additions of viable dHL60 cells, as well as...
following addition of DNA (Fig. 4A and G). However, these levels of biomass were no longer evident 72 h post-inoculation (Fig. 4A, D and G). This could be due to a destabilized proportion of the biofilm, comprised of cellular components, detached during analysis.

**DISCUSSION**

Clinical treatment of the microbial cause of pulmonary exacerbations in CF patients is confounded by the biofilm phenotype of the colonizing bacteria. When treatment is based on antibiotic susceptibility testing of biofilms formed by patients’ bacterial isolates, the mean time to pulmonary exacerbations is extended (Keays et al. 2009).

*Pseudomonas aeruginosa* biofilms, and their relevance to overcoming neutrophil-mediated immunity, have been the subject of a number of studies, owing to the high prevalence of the pathogen in the airways of people with CF. However, there is a relative lack of similar studies concerning Bcc, despite the disproportionate mortality associated with them. In the present study, we provide further evidence supporting the assertion that the biofilm is important for bacterial persistence in the CF airway and that this is in part due to the disruption of neutrophil antimicrobial activities.

Neutrophils, stimulated by microbially associated molecular patterns, produce IL-8 which serves to gather and further activate the neutrophil population at the site of colonization (Schröder et al. 2006b). We have demonstrated, however, that, when encountering Bcc biofilms, dHL60 cells display a diminished IL-8 secretory response (Fig. 2). This lack of self-reflexive activation may favor the bacteria by virtue of limiting the proinflammatory, antimicrobial character of the cells.

That similar levels of IL-8 secretion were observed for each of the three species, regardless of the age of the biofilm tested, reflects two complementary scenarios; first, an exofacial moiety itself induces a specific IL-8 response, possibly an EPS. Support for this concept can be derived from the study reported by König, Ceska and König (1995), which looked at the ability of *P. aeruginosa* to stimulate IL-8 release from neutrophils, as well as the ability of alginate from two *P. aeruginosa* strains to induce IL-8 release. They observed that alginate from the bacteria induced a statistically significant, but 10-fold lesser, release than did the bacteria themselves. Hence, though alginate is distinct from the EPS produced by Bcc bacteria, they may be similarly immunogenic.

Secondly, the type of pathogen recognition receptor (PRR) which becomes activated has a bearing on the type of IL-8 response (Schröder et al. 2006a). The authors of that study noted that, while LPS stimulation prompted de novo synthesis of IL-8 by neutrophils in conjunction with secretion from intracellular stores, agonism of CD66b resulted only in secretion of stored IL-8. Hence, Bcc EPS may be recognized by such a PRR, whose down-stream signaling fails to prompt IL-8 transcription.

dHL60 cells, which had been cultured in the presence of planktonic Bcc bacteria, exhibited 6.5–9.4-fold upregulation in expression of IL-8 mRNA transcript. Notably, while the dHL60 cells did, for the most part, display an induction of IL-8 expression when cultured with Bcc biofilms, the extent of the increase in IL-8 transcript synthesis was diminished with respect to that seen during planktonic coculture. This apparent loss of IL-8 induction mirrors our finding that secretion of IL-8 by dHL60 cells is greatly reduced during coculture with Bcc biofilms relative to planktonic Bcc bacteria.

In contrast to our observations, neutrophils have been reported to secrete comparable amounts of IL-8 when exposed to either planktonic *P. aeruginosa* PA01 bacteria or PA01 bacteria dwelling in a biofilm of 18 h maturity (Fuxman Bass et al. 2010). This may be due to the ability of *P. aeruginosa* extracellular DNA (eDNA) to stimulate IL-8 release from neutrophils, doing so in a TLR9-independent fashion (Trevani et al. 2003; Alvarez et al. 2006), given that *Pseudomonas* eDNA is often a component of their biofilms. Bcc bacteria vary in the extent to which their DNA forms part of the biofilm (Messiaen, Nelis and Coenye 2014); as such, it may not be as prevalent or immunogenic as that of *P. aeruginosa*.

There may be a failure by the dHL60 cells to encounter the cognate ligands for their PRRs. Such ligands may however be obscured by the surrounding EPS. We have shown that the ability of the cells to enter into the biofilm is compromised (Fig. 1), while the same compromised migration has been shown for neutrophils invading *P. aeruginosa* biofilms (Jesaitis et al. 2003; Bjarnsholt et al. 2005).

The implication, therefore, of our findings, and those of others highlighted here, would be that the biofilm would physically mask the bacteria within from being recognized by the dHL60 cells. Exposure of some proportion of the bacterial population at the surface of the biofilm would be likely, but this can be accounted for by the finding that under most conditions in our study the expression or secretion of IL-8 was observable.

Purified EPS has been shown to be a potent tactile stimulus of motility for neutrophils, which migrated into soft agarose suffused with EPS, even ignoring FMLP gradients (Hänisch et al. 2008). This was not specific chemotaxis, as Boyden chamber assays failed to demonstrate specific directional chemotaxis toward EPS in solution, but rather an enhanced motility which coincided with increased surface expression on the neutrophils of the cellular adhesion-associated protein, CD66b.

Jesaitis et al. (2003), however, reported that neutrophils were rendered ostensibly non-motile once deposited on *P. aeruginosa* biofilms, a finding which mirrors that of Hänisch and colleagues for biofilms rather than EPS alone. Movement of neutrophils on *Staphylococcus aureus* biofilms has also been shown to be limited, with greater migration being observed at the periphery of the larger biomass (Günther et al. 2009b). Our study of dHL60-Bcc biofilm interactions revealed the same phenomenon when the cells were inoculated onto established biofilms (Fig. 1).

When added to 6-day-old *S. aureus* biofilms, neutrophils were able to take up adjacent components of the biofilm over 60 min of observation, indicating that they were able to phagocytose the material (Günther et al. 2009b). The antimicrobial activity of the stationary neutrophil, then, is limited to its immediate vicinity, while more mature biofilms suffered lesser phagocytosis-mediated losses of biomass.

As we have shown for *B. multivorans* and *B. dolosa* biofilms, and as has been discussed for other biofilm-forming species, neutrophils are not well able to penetrate into the biofilm. They can be seen to internalize components of the biofilm from the periphery of the overall extracellular polymeric matrix, but whether this behavior could translate to an ‘outside-in’ form of biofilm clearance in vivo is unclear. We have observed that dHL60s remained largely stationary once adhered to the EPM. We also describe increased cell death associated with the cell’s encounter with Bcc biofilms, which would limit their ability to ingress into the biofilm. While compositional differences between our microtiter plate and flowcell-cultured biofilms are possible, the formation of a substantial biomass remains common between them. These findings suggest that neutrophils
may be capable only of erosion of the external face of a biofilm; thus, the biofilm can function as an effective defensive barrier for the bacteria within.

Phagocytosis of bacteria induces cell death in the neutrophil, with these apoptotic neutrophils being themselves phagocytosed by macrophages (Kobayashi et al. 2003; Zhang et al. 2003). The purpose of these events is presumably minimization of potential tissue damage which would arise from the action of proteases released from the decaying cell, as apoptosis allows for their controlled decomposition in macrophages without degradation of the cell membrane. Accordingly, PICD instigates a macrophage-directed signal encouraging phagocytosis of the neutrophil (Scannell and Flanagan 2007).

Neutrophils derived from healthy individuals undergo PICD when challenged with viable, planktonic B. cenocepacia isolates, as would be expected (Bylund et al. 2005). In the present study, we provide evidence that Bcc biofilms are also capable of inducing cell death in dHL60 cells (Fig. 3A) but that this is skewed toward necrosis in a biofilm age-dependent fashion over the duration of study (Fig. 3). Increasing age of the biofilm did not correlate directly with a corresponding increase in bacteria dwelling within. This suggests that another facet of the biofilm which emerges during its development may promote cell death in the dHL60 cells.

Induction of necrosis would result in the failure of effecrocytosis—neutrophil ingestion by macrophages—combined with the uncontrolled release of host tissue-damaging proteases, with a resulting proinflammatory state, as is observed in people with CF, who endure chronic pulmonary inflammation (Elizur, Cannon and Ferkol 2008).

Pursuant to the advantage of evading phagocyte attention, P. aeruginosa expresses toxins capable of inducing cell death in neutrophils (Dacheux et al. 2000; Xu et al. 2012). For example, biofilm-associated P. aeruginosa rhamnolipid has been demonstrated to induce necrosis in neutrophils in vitro (Jensen et al. 2007) and to prevent the bacteria’s clearance in a murine pulmonary infection model (van Gennip et al. 2009), resulting from the onset of neutrophil cell death following their inoculation onto Pseudomonas biofilm (van Gennip et al. 2012). Hence, biofilm-derived rhamnolipids enable the biofilm to act as a defensive barrier for Pseudomonas; as such, their production is upregulated in response to the presence of neutrophils via quorum sensing (Alhede et al. 2009).

The epidemic strain, B. cenocepacia J2315, has been shown to secrete a hemolytic lipopeptide with an ability to induce apoptosis in neutrophils (Hutchison, Poxtton and Govan 1998), though the prevalence of hemotoxins across the Burkholderia genus is limited (Carvalho et al. 2007). Hence, the Bcc strains studied here may not produce a compound with specific antineutrophil potency equivalent to that of either rhamnolipids or pyocyanin.

Pseudomonas aeruginosaa-secreted pyocyanin accelerates neutrophil cell death by inducing lysosomal membrane permeabilization with apoptosis ensuing through loss of Mcl-1 and induction of caspase-3 (Bianchi et al. 2008; Prince et al. 2008).

Notably, it has been shown that neutrophil-derived antimicrobial peptides interfere with phagocytosis by neutrophils (Voglis et al. 2009). Neutrophils for which phagocytosis is disrupted undergo cell death alternative to PICD, which may lead to their secondary necrosis within the CF airway milieu. The combination of this and the deliberate induction of necrosis by P. aeruginosaa and possibly other species would lead to the uncontrolled release of neutrophil antimicrobial peptides in a self-propagating cascade. This situation, then, would negate the antimicrobial influence of neutrophils and facilitate persistent infection.

Pseudomonas aeruginosaa bacteria are capable of biofilm formation in vitro despite the presence of viable neutrophils (Walker et al. 2005; Parks et al. 2009; Robertson et al. 2011; Caceres et al. 2014). In those and other studies, culture of P. aeruginosaa bacteria in the presence of human neutrophils resulted in greater biofilm formation than that formed by the bacteria alone due to the incorporation into the biofilm of exogenous DNA and protein (Chiang et al. 2013; Watters et al. 2014). No equivalent study concerning Bcc bacteria has yet been reported. Hence, we examined the effect of the bacteria with viable dHL60 cells, whole-cell lysates thereof or purified DNA.

We have shown that, despite an initial reduction in biofilm formation evident for B. multivorans LMG 13010 (Fig. 4A), culture of Bcc bacteria in the presence of viable dHL60 cells leads to an increase in biofilm biomass, while increased biomass was consistently observed for B. dolosa (Fig. 4B). This enhancement of biofilm biomass was further pronounced when bacteria were cultured in the presence of dHL60 cell lysate (Fig. 4E). It is likely that the presence of the lysate stimulated the bacteria to produce more EPS, macromolecules derived from the dHL60 cells adhered to the existing EPS or a combination of these events occurred.

Differing from either of these species, B. cenocepacia K56–2 bacteria demonstrated relatively static biomass. This may be due to the lesser biofilm-forming ability of this strain, with respect to the other two strains in this study, based on our findings during this study and in agreement with previous findings by Caraher et al. (2007), supported by data of other studies which report similar levels of biofilm biomass using the crystal violet assay (Aubert, Flannagan and Valvano 2008).

This provides a tentative explanation for our observations of differing outcomes between this strain and B. dolosa LMG 18941 and gives cause to speculate that secretion of greater quantities of EPS subsequently gives rise quantities of biofilm which are greater still, when formed in the presence of HL60 or dHL60 cells, with the EPS perhaps acting as an adhesive scaffold.

Many commonalities exist between the formation of biofilm by B. multivorans and B. cenocepacia bacteria. They share, for example, a dependence on the production of poly-β-1, 6-N-acetyl-D-glucosamine in order to produce a cohesive biofilm (Yakandawala et al. 2011). DNA is also a component of B. cenocepacia biofilms, wherein it is associated with the DNA-binding protein, DNABII (Novotny et al. 2013). However, the species have recently been shown to diverge in the prevalence of DNA in their respective biofilms (Messiaen, Nelis and Coenye 2013). In that study, B. multivorans LMG 13010 biofilms were reported to contain ∼1.5 μg ml⁻¹ DNA per 10⁸ cfu following 24 h of culture, while B. cenocepacia LMG 16656 bacteria (a CF isolate) contained only ∼0.15 μg ml⁻¹ DNA/10⁶ cfu. Whether this disparity could explain the observed difference in biomass between the biofilms of our strains when cultured with dHL60 cells is unclear.

The presence of extracellular DNA may be a requirement for initiation of biofilm formation by P. aeruginosaa (Hitchcurch et al. 2002). Inclusion by P. aeruginosaa of exogenous DNA appears, at least in part, to be driven by pyocyanin, through stimulation of DNA release from the bacteria themselves (Das and Manefield 2012; Das et al. 2013), as well as from neutrophils via induction of apoptosis. Indeed, such fortuitous or autologous lytic mechanisms are common in populations of many bacteria whose biofilms comprise DNA (Montanaro et al. 2011).

The DNA observed in the work of Messiaen et al. (2014) was autologously derived; we have provided evidence in our study
Figure 5. The paradigm of neutrophil recruitment to the CF airway. The airway epithelium, stimulated by microbial antigens, secretes proinflammatory agents and chemoattractants which recruit neutrophils in an attempt to clear the colonizing microorganisms. (1) Following bacterial challenge, neutrophils transmigrate into the airway lumen, where they encounter dehydrated, mucus-rich ASL (green). Segregated from the neutrophil, the bacteria adapt to their environment, for example by generation of an extracellular polymeric matrix. (2) The neutrophil encounters bacteria in a complex setting, including EPM polysaccharides and attempts to phagocytose the invading microorganism. (3) In a healthy airway, the neutrophil will then undergo phagocytosis-induced cell death (PICD) and be cleared by macrophage; however, in CF, phagocytosis is frustrated by the biofilm and neutrophil undergoes apoptosis followed by necrosis. Alternatively, necrosis may be directly induced by the invading pathogen. (4) PICD and (5) pro-resolving signaling are thereby inhibited, so the airway cannot return to a healthy state. (6) Where the neutrophils have not been cleared, they become necrotic and disintegrate; this releases antimicrobial peptides (AMPs) which (7) propagate further inflammation and (8) reinforce the EPM, leading, ultimately, to fibrotic lesions and lung damage.
formation. The necrotic cells then become incorporated into the EPM. These findings are in accord with those of Walker et al. (2005), who studied the effect of neutrophils on biofilm formation and development by P. aeruginosa PA01. Similarly to our observations, they reported increased formation of biofilm by PA01 in the presence of neutrophils.

Thus, if Burkholderia can remain viable while in the presence of neutrophils, then this would establish a self-perpetuating cycle of transient, ineffectual phagocytosis of invading Bcc member species while simultaneously amplifying the background inflammation and leukocyte recruitment (Fig. 5). Indeed, as discussed, biofilm formation can reduce phagocytosis by neutrophils, as well as reduce the efficacy of antimicrobial agents against the biofilm-dwelling microorganisms and may be an important factor in ensuring the persistence of infection in CF.

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