Intracellular survival of *Burkholderia cepacia* complex isolates in the presence of macrophage cell activation

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Strains of the *Burkholderia cepacia* complex have emerged as a serious threat to patients with cystic fibrosis due to their ability to infect the lung and cause, in some patients, a necrotizing pneumonia that is often lethal. It has recently been shown that several strains of the *B. cepacia* complex can escape intracellular killing by free-living amoebae following phagocytosis. In this work, the ability of two *B. cepacia* complex strains to resist killing by macrophages was explored. Using fluorescence microscopy, electron microscopy and a modified version of the gentamicin-protection assay, we demonstrate that *B. cepacia* CEP021 (genomovar VI), and *Burkholderia vietnamiensis* (previously *B. cepacia* genomovar V) CEP040 can survive in PU5-1.8 murine macrophages for a period of at least 5 d without significant bacterial replication. Furthermore, bacterial entry into macrophages stimulated production of tumour necrosis factor and primed them to release toxic oxygen radicals following treatment with phorbol myristoyl acetate. These effects were probably caused by bacterial LPS, as they were blocked by polymyxin B. Infected macrophages primed with interferon gamma produced less nitric oxide than interferon-gamma-primed uninfected cells. We propose that the ability of *B. cepacia* to resist intracellular killing by phagocytic cells may play a role in the pathogenesis of cystic fibrosis lung infection. Our data are consistent with a model where repeated cycles of phagocytosis and cellular activation without bacterial killing may promote a deleterious inflammatory response causing tissue destruction and decay of lung function.

Keywords: *Burkholderia cepacia*, macrophage activation, intracellular survival, cystic fibrosis

INTRODUCTION

Cystic fibrosis (CF), occurring at a frequency of 1:2500 live births, is the most common inherited lethal disease of Caucasian individuals. The product of the CF gene is a chloride channel found in both secretory and absorptive epithelia (di Sant’Agnese & Davis, 1976). Abnormal chloride transport across epithelial cells leads to pancreatic exocrine insufficiency, male and female infertility, and lung disease (Bye et al., 1994; di Sant’Agnese & Davis, 1976). The highly viscous mucus in the airways of CF patients may impair normal mucociliary clearance mechanisms, thus allowing colonization by opportunist micro-organisms (Gilligan, 1991; Govan & Deretic, 1996). Proteoglycans showing increased sulfation are components of the mucus in the airways, and may also interact with bacteria or their products to further increase mucus viscosity (Cheng et al., 1989; Gilligan, 1991). In addition, high sodium chloride concentrations in the airway surface fluid from CF patients have been found to inhibit the activity of bactericidal peptides secreted by epithelial cells (Smith et al., 1996). The combination of all these alterations is thought to contribute to recurrent and chronic respiratory tract infections that are the major cause of morbidity and mortality in CF disease (Bals et al., 1999).

Abbreviations: CF, cystic fibrosis; DCF, dichlorofluorescein; IFN-γ, interferon gamma; PMA, phorbol myristoyl acetate; PMB, polymyxin B; TNF, tumour necrosis factor.
Lung infection in CF individuals exhibits a characteristic age-related pattern involving colonization and intermittent exacerbations caused by *Staphylococcus aureus* and *Haemophilus influenzae* in infancy and early childhood, followed almost invariably in adolescence by *Pseudomonas aeruginosa* (Gilligan, 1991; Govan & Deretic, 1996). However, more recently and in parallel with an improved life expectancy of CF patients (FitzSimmons, 1996), the spectrum of microbial pathogens has expanded to also include *Burkholderia cepacia*, which currently can be isolated from as many as 40% of CF patients (Govan & Deretic, 1996). Colonization and infection by *B. cepacia* has become a major concern for these patients due to three salient features. First, although most CF patients harbouring *B. cepacia* show a chronic non-aggressive infection associated with periods of acute exacerbations, up to 20% exhibit an accelerated clinical deterioration referred to as the cepacia syndrome. This condition is characterized by a rapid reduction of pulmonary function associated with sepsis and high mortality (Govan & Deretic, 1996; Isles et al., 1984). Second, there is compelling evidence indicating that *B. cepacia* can be transmitted from patient to patient (Govan et al., 1996). Third, the innate multi-drug resistance of *B. cepacia* has made effective antimicrobial treatment very problematic. *B. cepacia* is a soil bacterium, metabolically versatile, and taxonomically complex. Several different species can be distinguished by polyphasic taxonomy and PCR-based genetic analysis among isolates from clinical and environmental sources. Since the members of each group are genotypically distinct but have phenotypic traits shared among groups, they are referred to as genomovars. Up to six genomovars are known to occur in CF patients; these isolates are referred to as the *B. cepacia* complex (Vandamme et al., 1997). Of these, genomovar III strains occur in approximately 50% of CF cases colonized with *B. cepacia* and they are more commonly, but not exclusively, associated with the cepacia syndrome (Vandamme et al., 1997). Members of the other genomovars occur at lower frequencies.

Recent work in various laboratories suggests that *B. cepacia* isolates may be able to survive intracellularly. Burns et al. (1996) reported that a clinical isolate of *B. cepacia* could enter and replicate in the A549 human epithelial carcinoma cell line. Also, *B. cepacia*-like micro-organisms are endosymbiotically associated with arbuscular mycorrhizal fungi, residing within cytoplasmic vesicles (Perotto & Bonfante, 1997). More recently, Marolda et al. (1999) have shown that several members of the *B. cepacia* complex are able to survive within various types of free-living amoeba of the genus *Acanthamoeba*. It is well established that several microorganisms surviving within amoebae, such as *Legionella pneumophila*, *Mycobacterium avium*, *Chlamydia pneumoniae* and *Listeria monocytogenes*, are also able to survive and replicate within macrophages (Brown & Barker, 1999). Therefore, we investigated whether *B. cepacia* complex strains are capable of surviving intracellularly within macrophages, since this property could contribute to its pathogenicity in CF individuals. In the present study, we demonstrate that two clinical isolates of the *B. cepacia* complex can survive for an extended period of time, as compared to *Escherichia coli*, in a murine macrophage cell line. We also show that bacterial survival occurs despite macrophage activation following phagocytosis.

**METHODS**

**Bacterial strains, cell lines and reagents.** Clinical isolates of *B. cepacia* from CF patients were obtained from the Vancouver collection, and assigned to individual genomovars by P. Vandamme (Laboratory of Microbiology, University of Gent, Belgium) and E. Mahenthiralingam (Department of Pediatrics, University of British Columbia, Canada). *B. cepacia* strain CEP021 belongs to the newly defined genomovar VI (P. Vandamme, personal communication). This strain is a non-epidemic isolate obtained from a CF patient in Philadelphia in 1986 that had a different ribotyping profile than the Philadelphia epidemic strain (J. J. Lipuma, personal communication). *B. vietnamiensis* (formerly *B. cepacia* complex genomovar V) strain CEP040, originally described as *B. cepacia* PC315 (J. Burns, personal communication) was recovered at the time of autopsy from sputum of a CF patient with necrotizing pneumonia, and is the same isolate used by Burns et al. (1996) to demonstrate invasion and survival in an epithelial cell line. Cultures were grown at 37 °C in Luria–Bertani (LB) medium. *B. cepacia* isolates were examined for antibiotic susceptibility to determine their suitability for the intracellular invasion assay (see below). Isolates were grown on plates containing 100 µg gentamicin ml⁻¹, 100 µg kanamycin ml⁻¹, 100 µg ceftazidime ml⁻¹ (Eli Lilly), 100 µg streptomycin ml⁻¹ or 100 µg tetracycline ml⁻¹. The strains of *E. coli* HB101(pHP45Ω), resistant to penicillin and streptomycin, and of *Listeria monocytogenes* 42 were from our laboratory and the E. G. D. Murray culture collection (Dept Microbiology and Immunology, University of Western Ontario, Canada), respectively. Murine macrophage cell lines P51-1.8 (ATCC TIB-63) and J774A.1 (ATCC TIB-67), and L929 fibroblasts (ATCC CCL-1) were maintained in RPMI 1640 medium (Life Technologies) supplemented with 10% fetal bovine serum (Life Technologies), 50 U penicillin ml⁻¹ and 50 U streptomycin ml⁻¹. All chemicals and reagents were from Sigma unless indicated otherwise.

**Macrophage cell invasion assay.** The gentamicin-protection assay (Elingshorst, 1994), as modified by Burns et al. (1996), was used to quantify intracellular bacteria. P51-1.8 macrophages were seeded at a density of 7.5 x 10⁴ cells well per well and allowed to grow overnight. Bacteria grown to mid-exponential phase were washed, diluted in tissue culture medium, and then added to macrophages at an m.o.i. of 1–5 bacteria per macrophage. Infected monolayers were centrifuged at 500 g for 5 min and incubated for 2 h at 37 °C in 5% CO₂, followed by three washes with PBS to remove non-ingested extracellular bacteria. Fresh medium containing 1 mg ceftazidime ml⁻¹ and 500 µg kanamycin ml⁻¹ was added to kill the remaining extracellular bacteria, and cultures were incubated for another 2 h. At this time, antibiotics were removed by washing with PBS, and an aliquot from the final wash was plated to determine the number of any remaining extracellular bacteria. Macrophages were lysed by adding Triton X-100 to a 1% final concentration. Lysates were serially diluted and plated following the procedure described by Jett et al. (1997). The number of intracellular bacteria was calculated from the difference between colony counts in the PBS wash and those in
the lysate. For long-term incubation experiments, parallel sets of cultures were maintained with and without kanamycin (25 µg ml⁻¹). In both cases, prior to sampling, cultures were treated with 1 mg ceftazidime ml⁻¹ and 300 µg kanamycin ml⁻¹ for 2 h, to further minimize the number of extracellular bacteria recovered (Burns et al., 1996). At each time point, macrophages were removed from the dish by treatment with trypsin, counted and then lysed to quantify intracellular organisms as described above.

**Fluorescence and electron microscopy.** Infected and uninfected macrophages grown on coverslips were stained with acridine orange and counterstained with crystal violet as previously described by Miliotis (1991). Briefly, 1 ml 0.01% acridine orange solution in PBS was added to each well for 45 seconds. Wells were washed with PBS, followed by the addition of 1 ml 0.05% crystal violet in 0.15 M NaCl for 1 min. After washing with PBS, coverslips were removed, inverted onto glass slides and the samples examined with a Zeiss Axioskop fluorescence microscope with excitation at 488 nm from an Argon-Helium laser. Vital staining with acridine orange is a sensitive method for detecting ingested bacteria and also allows differentiation between live and dead intracellular bacteria (Rost, 1995), as dead bacteria fluoresce red (metachromasy) and live bacteria fluoresce green (orthochromasy). Transmission electron microscopy was performed as previously described by Jones et al. (1996) and samples were visualized using a Phillips 300 transmission electron microscope located at the University of Guelph Scanning and Transmission Electron Microscope Facility.

**Flow cytometry.** Light scattering from PU5-1.8 cells infected with bacteria was used to determine the size of the cell population using flow cytometry (Salzman et al., 1990). Infected cultures were incubated for 72 h, after which time the macrophages were collected, washed in PBS and fixed overnight in 4% formaldehyde. Fixed cells were washed and resuspended in 300 µl PBS. Flow cytometry was performed with a FACSscan flow cytometer (Becton Dickinson) and the mean values of forward scatter (FSC), representing cell size, and side scatter (SSC), representing cell granularity, were determined utilizing CellQuest software.

**Tumour necrosis factor (TNF) production.** Live or heat-killed (80 °C for 30 min) bacteria were added to macrophage cultures at a ratio of 10 bacteria per macrophage. The levels of biologically active TNF in supernatants collected after the 2 h antibiotic treatment were determined by the use of a standard L929 killing assay (Martin & Dorf, 1990). Supernatants from macrophages treated with 100 ng LPS ml⁻¹, which was prepared and purified in our laboratory from cultures of *Shigella flexneri* (Yao et al., 1992), were used as a positive control for TNF production. Each experiment was conducted in quadruplicate. Briefly, 1 x 10⁴ L929 cells were seeded in each well of a 96-well plate and incubated overnight at 37 °C in 5% CO₂. Fifty microlitres of macrophage supernatants (serially diluted in RPMI 1640) were added to L929 cell cultures, followed by the addition of 50 µl RPMI 1640 medium containing 5 µg actinomycin D ml⁻¹. Cultures were incubated for 18 h and the viability assessed using the modified MTT [3-(4,5-dimethylthiazol-2-yl)2,5-diphenyltetrazolium bromide] assay as described by van de Loosdrecht et al. (1991). TNF-α activity in culture supernatants was quantified by comparison with standard murine recombinant TNF-α, with 1 U TNF-α being defined as the amount of TNF causing 50% L929 cell death. To ascertain whether TNF-α induction was caused by *B. cepacia* LPS, experiments were conducted with polymyxin B (PMB), which was added to the macrophages at final concentrations of 5, 50 and 100 µg ml⁻¹ 1 h prior to infection.

**Measurement of respiratory-burst activity.** Priming of PU5-1.8 macrophages for superoxide release upon phorbol myristate acetate (PMA) stimulation following bacterial phagocytosis was determined by using the fluorescent probe dichlorofluorescein (DCF) and flow cytometry (Bass et al., 1983). This method is based on the metabolism of 2,7'-dichlorofluorescin diacetate, which, after being taken up by macrophages, is subsequently hydrolysed by esterases to non-fluorescent, membrane impermeable DCF. Stimulation of the respiratory burst within cells results in the rapid oxidation of DCF to a fluorescent form by the action of peroxides in a process accelerated by the presence of intracellular peroxidases. Thus, DCF fluorescence is proportional to peroxide production. All reagents used in these experiments were obtained from Beckman-Coulter (CellProbe DCFH, PMA- Oxidative Burst). Macrophages were infected with bacteria at an m.o.i. of approximately 10 bacteria per macrophage, and treated with antibiotics as indicated above. Uninfected macrophages treated and untreated with 100 ng *S. flexneri* LPS ml⁻¹, served as positive and negative controls, respectively. Macrophages were harvested at 36 h post-infection, and levels of superoxide release after 30 min following stimulation with PMA, as well as prior to stimulation, were determined by fluorescence-activated cell sorting analysis. Experiments were also conducted with PMB as indicated above for determination of TNF.

**Measurement of nitrite.** Interferon gamma (IFN-γ) is known to prime macrophages to release NO upon further stimulation, increasing their microbialid properties (Stuehr & Marletta, 1985, 1987). One set of macrophage cultures was treated with 50 U IFN-γ ml⁻¹ 24 h prior to infection. Live or heat-killed...
bacteria were added to macrophage cultures at 10–20 bacteria per macrophage, followed by antibiotics as previously described. After the initial 2 h antibiotic treatment, macrophages were washed with Hank’s balanced salt solution and the medium replaced with 200 µl fresh medium containing 25 µg kanamycin ml⁻¹ and 100 µM PTIO (2-phenyl-4,4,5,5-tetramethylimidazolin-1-oxyl 3-oxide; ICN). Nitrite production was determined using the Griess reagent as described by Amano & Noda (1995). Absorbance was measured at 540 nm using an ELISA plate reader spectrophotometer, and nitrite concentrations were quantified from a standard curve with NaNO₂. NO production by IFN-γ-primed macrophages treated with LPS was determined as a positive control.

**Statistics.** Data were analysed by single factor ANOVA. Results were considered significant at $P < 0.01$.

**RESULTS**

**B. cepacia CEP021 and B. vietnamiensis CEP040 survive within PU5-1.8 macrophages**

Removal of extracellular bacteria was an obligatory step prior to the quantification of intracellular bacteria. Thus, we used B. cepacia complex strains susceptible to antibiotics that do not penetrate the macrophage cell membrane, such as ceftazidime and kanamycin. We found that the majority of genomovar III isolates examined were resistant to 500 µg ceftazidime ml⁻¹ and all aminoglycosides tested, making it impossible to assess their intracellular survival. In contrast, B. cepacia CEP021 and B. vietnamiensis CEP040 were sensitive to 100 µg ceftazidime ml⁻³ and 100 µg kanamycin ml⁻³. A similar combination of antibiotics has been used previously by Burns *et al.* (1996) to show entry and intracellular replication of strain CEP040 in epithelial cells. Also, we have recently shown that strains CEP021 and CEP040 can survive intracellularly within *Acanthamoeba* (Marolda *et al.*, 1999). Therefore, both bacterial strains were used in the experiments described in this study.

The PU5-1.8 macrophage cell line was chosen based on a previous report indicating that these cells possess immunocytochemical characteristics of mature macrophages (Nibbering & Van Furth, 1988). Phagocytosis and intracellular survival of CEP021 was followed using acridine orange–crystal violet staining. As a control, macrophages were infected with *E. coli* HB101(pPH45Ω). Both *E. coli* HB101 and CEP021 cells were visible within the cytoplasm of infected macrophages at 2 h and 24 h post-infection (Fig. 1a, b). At these times, 3–5% of macrophages were infected with either CEP021 or *E. coli*. At 48 h and 72 h post-phagocytosis, *E. coli* HB101 cells were no longer visible within infected macrophages (Fig. 1c), suggesting that intracellular bacteria were killed and degraded. In contrast, CEP021 cells remained clearly visible as clusters of cells and also as individual bacteria scattered within the cytoplasm (Fig. 1d). A few bacteria were dead, as they displayed red fluorescence, but the great majority (more than 80%) exhibited green fluorescence, indicating they were still viable. This was supported by the fact that intracellular *B. cepacia* were highly motile, suggesting that they remained metabolically active. In some experiments, the observation period was extended...
to 96 h and 120 h, with similar results. Therefore, we concluded that *B. cepacia* CEP021 resisted macrophage bactericidal mechanisms and thus survived intracellularly for a prolonged period of time. Pretreatment of macrophages with cytochalasin D reduced bacterial internalization to less than 0.25% with respect to untreated controls. Internalization of bacteria was largely unaffected when macrophages were pretreated with colchicine (data not shown). Thus, internalization to less than 0.25% was not restricted to PU5-1.8 cells. Identical results were obtained using a macrophage cell line of different lineage, J774A.1, with serum. Similar results were obtained using a PU5-1.8 macrophage does not require opsonization of recovered intracellular bacteria also increased. Based on these results, we standardized the bacterial inoculum – 10⁸ bacteria were used with 10⁶ macrophages per well. Macrophages were incubated with bacteria for 2 h and then treated for another 2 h with ceftazidime and kanamycin as indicated in Methods. The experiment was done in triplicate. The number of intracellular bacteria was calculated from the subtraction of extracellular bacteria from the total bacteria recovered after lysis of macrophages.

### Table 1. Kinetics of invasion of PU5-1.8 macrophages by CEP021

<table>
<thead>
<tr>
<th>Bacterial inoculum (cfu)</th>
<th>m.o.i.</th>
<th>Intracellular bacteria ± SD*</th>
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<tbody>
<tr>
<td>9 × 10⁶</td>
<td>0.009</td>
<td>2.01 × 10⁷ ± 0.51 × 10⁷ (22.2)</td>
</tr>
<tr>
<td>9 × 10⁷</td>
<td>0.09</td>
<td>6.48 × 10⁵ ± 0.32 × 10⁵ (7.2)</td>
</tr>
<tr>
<td>9 × 10⁵</td>
<td>0.9</td>
<td>2.92 × 10⁵ ± 0.96 × 10⁵ (3.2)</td>
</tr>
<tr>
<td>9 × 10⁶</td>
<td>9</td>
<td>5.08 × 10⁴ ± 0.76 × 10⁴ (6.4)</td>
</tr>
<tr>
<td>9 × 10⁷</td>
<td>90</td>
<td>1.82 × 10³ ± 0.53 × 10³ (2)</td>
</tr>
</tbody>
</table>

*Numbers in parentheses indicate the percentage of the inoculum recovered after lysis of macrophages.

To further elucidate whether intracellular replication occurs we used a modified antibiotic protection assay (Burns et al., 1996). A combination of 1 mg ceftazidime ml⁻¹ and 500 µg kanamycin ml⁻¹ was used to kill extracellular bacteria in all invasion experiments. Table 1 reveals that as the inoculum size increased, the number of recovered intracellular bacteria also increased. Based on these results, we standardized the bacterial inoculum to approximately 10⁸ micro-organisms per ml, and used an m.o.i. of 1–10 bacteria per macrophage. Under these conditions, extracellular bacteria represented less than 6% of the total bacteria recovered from the cell lysate. During long-term incubation experiments in the presence of kanamycin, the levels of extracellular bacteria remained roughly constant, indicating that the antibiotic prevented the growth of residual extracellular bacteria. This was confirmed by the lack of visible extracellular bacteria as determined by direct microscopic observation of infected cultures. However, over the course of 72 h, the number of intracellular bacteria decreased by ~ 3 log units, strongly suggesting that there is no intracellular bacterial replication (Fig. 3a). It is possible that during long-term incubations, the antibiotic can

**Fig. 3.** Quantification of intracellular bacteria in long-term incubations. After 2 h infection with CEP021, PU5-1.8 macrophages were treated with kanamycin and ceftazidime for 2 h. Fresh medium with 25 µg kanamycin ml⁻¹ (a) or without kanamycin (b) was added and incubations continued for 24, 48 and 72 h. Both intracellular (black bars) and extracellular (white bars) micro-organisms were quantified at 2, 24, 48 and 72 h following the standard 2 h incubation with ceftazidime and kanamycin (see Methods). Error bars represent the standard error of three values.

**Quantification of intracellular bacteria**

To further elucidate whether intracellular replication occurs we used a modified antibiotic protection assay (Burns et al., 1996). A combination of 1 mg ceftazidime ml⁻¹ and 500 µg kanamycin ml⁻¹ was used to kill extracellular bacteria in all invasion experiments. Table 1 reveals that as the inoculum size increased, the number of recovered intracellular bacteria also increased. Based on these results, we standardized the bacterial inoculum to approximately 10⁸ micro-organisms per ml, and used an m.o.i. of 1–10 bacteria per macrophage. Under these conditions, extracellular bacteria represented less than 6% of the total bacteria recovered from the cell lysate. During long-term incubation experiments in the presence of kanamycin, the levels of extracellular bacteria remained roughly constant, indicating that the antibiotic prevented the growth of residual extracellular bacteria. This was confirmed by the lack of visible extracellular bacteria as determined by direct microscopic observation of infected cultures. However, over the course of 72 h, the number of intracellular bacteria decreased by ~ 3 log units, strongly suggesting that there is no intracellular bacterial replication (Fig. 3a). It is possible that during long-term incubations, the antibiotic can
penetrate macrophages and prevent growth of intracellular bacteria. For this purpose, we also conducted phagocytosis experiments without antibiotic during long-term incubations. In these cases, microscopic examination of cultures at 2 h and 24 h post-infection revealed no extracellular bacteria, but by 48 h and 72 h, extracellular bacteria were readily visible. Fig. 3 (b) shows that between 2 h and 72 h, the number of intracellular bacteria increased 12-fold (1-1 log units), while extracellular bacteria increased 157-fold (2-2 log units). Extracellular bacteria never represented more than 1-8% of the total bacteria recovered at any given time point, suggesting that greater than 98% of the total bacteria recovered were of extracellular origin. However, the presence of extracellular bacteria in the tissue culture medium, especially at 48 h and 72 h post-infection, indicates that the initial antibiotic treatment did not efficiently kill all extracellular bacteria. Similar observations were made by Burns et al. (1996) with strain CEP040 in the epithelial cell line A549. Conceivably, extracellular bacteria escaping killing by antibiotics administered at the beginning of the experiment may enter the macrophages at later times, and contribute to the intracellular fraction recovered at 48 h and 72 h. Thus, the apparent increase in the number of intracellular bacteria is most likely an artefact due to the presence of residual extracellular bacteria not effectively killed by the initial treatment with ceftazidime and kanamycin. From all these experiments, we conclude that intracellular bacteria do not replicate, although it is not possible to rule out a low level of replication. In a control phagocytosis assay using the intracellular pathogen *Listeria monocytogenes*, we detected more than two doublings in intracellular micro-organisms over the course of a 6 h invasion experiment (data not shown), demonstrating that PU5-1.8 macrophages can support the replication of intracellular bacteria.

**Bacterial infection causes activation of macrophages**

Under our experimental conditions, the viability of macrophages at the various time points during phagocytosis assays exceeded 95%, as determined by trypan blue exclusion. However, following bacterial infection macrophages appeared to undergo morphological changes involving enlargement of the cytoplasm and vacuolization (Fig. 4a, b). At 96 h and 120 h after phagocytosis, most of the cells were floating in the
Activated macrophages are known to produce a number of cytokines including TNF-α (Wewers, 1997). Thus, we investigated production of TNF-α by PU5-1.8 macrophages infected with bacteria as a marker for cell activation. The concentration of TNF-α in macrophage culture supernatants 2 h post-phagocytosis was determined using a bioassay based on the killing of TNF-α-sensitive L929 cells. A basal level of 0.04 ± 0.009 U TNF-α was spontaneously produced by PU5-1.8 macrophages, as previously reported for this cell line (Biragyn & Nedospasov, 1995). However, following treatment with LPS, PU5-1.8 cells produced and released significantly more TNF-α than untreated macrophages (3.36 ± 1.147 U, \( P < 0.01 \)). Culture supernatants of macrophages exposed to live bacteria had a significantly higher amount of TNF-α (67.48 ± 4.18 U, \( P < 0.01 \)) than supernatants of untreated macrophages. In contrast, heat-killed bacteria did not induce higher levels of TNF-α than those found following stimulation with LPS alone (5.94 ± 0.64 U), suggesting that production of TNF-α required viable bacteria. It has been shown by other investigators that ceftazidime causes release of biologically active LPS from the bacterial outer membrane (Dofferhoff et al., 1993; Kirikae et al., 1998). Therefore, to determine whether LPS was involved in TNF-α production, we conducted inhibition studies using PMB. PMB binds to the lipid A component of the LPS molecule, thereby preventing its interaction with its receptor and thus inhibiting TNF-α production (Lasfargues et al., 1989). Treatment with PMB at concentrations of 5, 50 and 100 μg ml\(^{-1}\) prior to infection of macrophages reduced TNF-α production by 50, 75 and 85%, respectively. Thus, we concluded that soluble LPS released by live bacteria is involved in the stimulation of macrophages following bacterial infection.

Activated macrophages also produce reactive oxygen species and nitrogen intermediates (Granger et al., 1990; Guthrie et al., 1984) that contribute to the killing of phagocytized bacteria. In a previous study, it has been shown that oxidative killing is an important component of the macrophage bactericidal activity against B. cepacia complex isolates (Speert et al., 1994). Since our microscopic studies indicated that B. cepacia CEP021, but not the E. coli control, survive within macrophages for at least 72 h, it was important to determine whether survival occurs in the presence of bactericidal oxygen.

**Fig. 5.** Flow cytometry analysis to determine the formation of superoxide by macrophage cell cultures. The graphs show the fluorescence distribution of DCF oxidation prior to (a–d) and 30 min after (e–h) stimulation with PMA. The abscissa represents the macrophage cell fluorescence intensity on a logarithmic scale (arbitrary units). The experiment included untreated macrophages (a, e), B. cepacia-infected macrophages (b, f), LPS-treated macrophages (c, g) and B. cepacia-infected macrophages pre-treated with PMB (d, h).
Table 2. Nitrite production by PU5-1.8 macrophages

<table>
<thead>
<tr>
<th>Treatment*</th>
<th>Nitrite concentration (µM) ± SD†</th>
<th>Induction factor</th>
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<tr>
<td>None</td>
<td>7.42 ± 0.83</td>
<td>1</td>
</tr>
<tr>
<td>IFN-γ</td>
<td>76.5 ± 4.26</td>
<td>10.33</td>
</tr>
<tr>
<td>LPS</td>
<td>8 ± 1.88</td>
<td>1</td>
</tr>
<tr>
<td>IFN-γ + LPS</td>
<td>110.76 ± 5.37</td>
<td>14.8</td>
</tr>
<tr>
<td>Live bacteria</td>
<td>8.65 ± 0.49</td>
<td>1</td>
</tr>
<tr>
<td>IFN-γ + live bacteria</td>
<td>52.65 ± 2.84</td>
<td>7.1</td>
</tr>
<tr>
<td>Heat-killed bacteria</td>
<td>8.88 ± 0.37</td>
<td>1</td>
</tr>
<tr>
<td>IFN-γ + heat-killed bacteria</td>
<td>79.65 ± 5.06</td>
<td>10.7</td>
</tr>
</tbody>
</table>

*IFN-γ pre-treated and untreated macrophages were exposed to heat-killed or live B. cepacia CEP021 cells. Macrophages were also treated with LPS to determine the maximum production of nitric oxide. Data were collected 24 h after phagocytosis or treatment with LPS.
†Culture supernatants were assayed for nitrite concentration using the Griess reagent. Data shown represent the mean and standard error of 8 experiments.

and nitrogen intermediates. LPS primes macrophages for enhanced release of superoxide in response to other stimuli such as PMA or N-formylated bacterial peptides (Guthrie et al., 1984; Vosbeck et al., 1990). Thus, superoxide ion production was investigated by treating infected macrophage cultures with PMA. Macrophage cultures either infected with bacteria or treated with purified LPS showed much less superoxide production than stimulated with PMA (Fig. 5f, g). In contrast, macrophage cultures treated with bacteria in the presence of PMB showed much less superoxide production (Fig. 5h), suggesting released LPS is the primary mechanism involved in the ability of B. cepacia to prime macrophages for superoxide production. This conclusion is in agreement with previous observations by other groups indicating that B. cepacia LPS is a potent inducer of phagocytes (Hughes et al., 1997; Smith et al., 1999).

To examine whether phagocytosis of B. cepacia is associated with production of nitric oxide, we measured the levels of nitrite in supernatants from infected and uninfected macrophages. In this case, macrophages were primed with IFN-γ. Irrespective of the stimulus, in the absence of priming by IFN-γ the concentration of nitrite produced by macrophage cultures was very low and virtually identical to the basal level spontaneously produced by untreated cells (Table 2). In contrast, nitrite production with IFN-γ pre-stimulation and LPS treatment increased 10- and 14-fold with respect to basal levels (Table 2). Priming with IFN-γ followed by exposure to heat-killed bacteria resulted in nitrite levels similar to those observed with IFN-γ alone, suggesting that inactivated bacteria do not stimulate any further increase in NO synthesis. In contrast, infection of primed macrophages with live B. cepacia CEP021 resulted in a significantly lower nitrite production than that attributed to IFN-γ alone (P < 0.01), which suggests that B. cepacia infection may interfere with the levels of nitrite. Since under the experimental conditions used, live B. cepacia release LPS, and LPS enhances the levels of measured nitrite (Table 2), we attributed the observed decrease to either inhibition of the NO synthesis pathway or utilization of nitrite by metabolically active intracellular bacteria.

**DISCUSSION**

*B. cepacia can survive intracellularly in the absence of significant bacterial replication*

We report in this study that two isolates of the B. cepacia complex can survive within the murine macrophages cell line PU5-1.8 for a period of several days. Direct observations with fluorescence and electron microscopy indicate that B. cepacia CEP021, B. vietnamiensis CEP040 and the E. coli control entered PU5-1.8 macrophage cells, but only B. cepacia and B. vietnamiensis remained detectable 3 d after infection. This is in accordance with previously reported results (Caron et al., 1994), indicating that survival of E. coli K-12 in U937 macrophages was limited to a maximum of 3 d. Several observations support the conclusion that intracellular B. cepacia CEP021 and B. vietnamiensis CEP040 cells are viable: i) bacteria within vacuoles were highly motile, even in some experiments where observations were extended to 120 h, suggesting they are metabolically active; ii) more than 80% of the intracellular bacteria displayed green fluorescence with acridine orange; iii) micro-organisms were recovered by culture after lysis of infected macrophages in the absence of extracellular bacteria, as observed microscopically. Therefore, we were able to show that B. cepacia CEP021 and B. vietnamiensis CEP040, but not E. coli HB101, are capable of long-term intracellular survival in a murine macrophage-derived cell line.

Experiments to demonstrate intracellular replication by the use of the kanamycin/ceftazidime protection assay consistently showed that in the presence of kanamycin during long-term incubations, there was a progressive decrease in the number of viable bacteria recovered over the course of 72 h. Several groups working with other micro-organisms such as Rhodococcus equi (Hondalus & Mosser, 1994), Klebsiella pneumoniae (Oelschlaeger & Tall, 1997) and Mycobacterium tuberculosis (Mehta et al., 1996) have reported a continual decrease in recovered intracellular bacteria despite microscopic observations suggesting bacterial replication. Although aminoglycoside antibiotics are generally thought to be unable to penetrate eukaryotic cell membranes, several studies show they can enter macrophages and inactivate intracellular bacteria (Crowle et al., 1984; Drevets et al., 1994). It has also been proposed that toxicity due to growth of intracellular bacteria causes alterations of the eukaryotic cell membrane, thereby increasing its permeability to antibiotics in the murine environment (Hondalus & Mosser, 1994). Furthermore, bacterial infection with B. cepacia complex strains could cause macrophage cell
Lysis and release of intracellular bacteria into the medium. These micro-organisms would be subsequently killed by the extracellular kanamycin and thus, despite their intracellular origin, they would not contribute to the recovered intracellular pool of bacteria. A recent study has reported that a purified lipopeptide produced by a genomovar III strain under very strict growth conditions in vitro induces apoptosis in neutrophils (Hutchinson et al., 1998). We were unable to demonstrate apoptosis in PU5-1.8 macrophages infected with the genomovar VI strain CEP021 and the B. vietnamiensis strain CEP040 (data not shown). However, the growth conditions used in these experiments are different than those of Hutchinson et al. (1998).

Increasing bacterial cell numbers, as early as 48 h after incubation, were observed in the absence of kanamycin. This experimental protocol was the same as the one used by Burns et al. (1996) to infect epithelial cells with B. vietnamiensis CEP040. Under these conditions, we observed only a 1-1 log increase of intracellular bacteria over a period of 72 h. This represents a small increase compared with bacterial growth in culture without macrophages, where the micro-organisms reach several log units within 24–36 h (Marolda et al., 1999). In a previous study of intracellular survival of B. cepacia complex strains (including CEP021 and CEP040) within amoebae, we have reported a similarly small increase in the number of intracellular bacteria and demonstrated that the principal mechanism of bacterial replication is extracellular (Marolda et al., 1999). Although in the long-term phagocytosis assays, extracellular bacteria represented no more than 1.8% of the total number of recovered bacteria, the small increase in intracellular micro-organisms could be explained by phagocytosis of bacteria undergoing replication that fail to initiate new replication cycles once they become intracellular. Taken together, our results suggest that our B. cepacia complex isolates do not replicate intracellularly in phagocytic cells, and caution is warranted when interpreting invasion experiments with B. cepacia complex strains using other cell systems.

**Presence of intracellular B. cepacia is associated with macrophage cell activation**

The change in physical size of infected macrophages, as observed by light microscopy, suggested that they might be activated. Macrophage activation, along with physical cell changes, leads to up-regulation of major histocompatibility complex class II antigens and accessory molecules such as B7.1, as well as production of various cytokines, including TNF-α and interleukin-8. Our studies suggested that live B. cepacia complex strains primed PU5-1.8 cells to produce TNF-α and superoxide. Released LPS mediated these effects, since they were inhibited by PMB, and a much lower level of stimulation was observed with heat-killed bacteria. Release of LPS from bacteria may be a consequence of antibiotic treatment with ceftazidime. Other investigators have documented significant levels of biologically active LPS in the medium of cultures of P. aeruginosa and E. coli incubated with ceftazidime (Dofferhoff et al., 1993; Kirikae et al., 1998). A similar situation may occur in the CF patient, where LPS release following antibiotic treatment, in conjunction with the ability of B. cepacia complex strains to remain viable after phagocytosis, may contribute to an enhanced inflammatory response and progression of lung disease.

NO, produced by the nitric oxide synthase, has been identified as the effector molecule for killing a range of intracellular pathogens (Lowenstein et al., 1994; Summersgill et al., 1992). When primed with IFN-γ for 24 h prior to infection, macrophages exposed to heat-killed bacteria yielded nitrite levels similar to those obtained with PU5-1.8 cells treated with IFN-γ alone. Although this could be due to a lack of LPS release by heat-killed bacteria, other possibilities should be considered. Previous studies suggested that only phagocytosis of opsonized micro-organisms can engage the intracellular signalling pathways necessary for NO production (Gross et al., 1998). Since we have shown that serum does not appear to be required for B. cepacia CEP021 entry into PU5-1.8 macrophages, lack of NO induction could be explained by bacterial entry via a non-opsonic mechanism. The decreased level of nitrite found in macrophages exposed to live B. cepacia CEP021 was unexpected, since we have shown that under our experimental conditions live bacteria release LPS, which in turn stimulates NO production. It is possible that following phagocytosis, B. cepacia complex isolates may induce the production of immunomodulatory molecules, such as prostaglandins, that downregulate NO synthesis (D’Acquisto et al., 1998; Venkataprasad et al., 1996), or alternatively, live micro-organisms utilize nitrate. Utilization of nitrate by B. cepacia has been previously documented (Gessner & Mortensen, 1990).

In summary, the results presented in this work, together with those of a previous report (Marolda et al., 1999) indicate that B. cepacia complex isolates can survive intracellularly within phagocytic cells. More importantly, survival takes place in the presence of macrophage activation, especially superoxide release. Although a previous report indicated that B. cepacia complex isolates can be killed by oxidative mechanisms while resisting non-oxidative damage (Speert et al., 1994), our data suggest that resistance to oxidation may play a role in survival of intracellular bacteria. Both catalase and superoxide dismutase are important for detoxifying toxic oxidants from cells (Hassett & Cohen, 1989). We are currently examining their occurrence and characteristics in isolates of the B. cepacia complex in relation to intracellular survival (M. D. Lefebre & M. A. Valvano, unpublished). In the context of the pathogenesis of CF lung infection, our data suggest a model whereby B. cepacia complex strains can resist phagocytic killing, while LPS released from invading micro-organisms (either spontaneously or after antibiotic treatment) at the site of infection may sensitise local phagocytic cells to amplify the inflammatory response. It has been shown by other investigators that B. cepacia...
LPS is more pro-inflammatory than LPS of *P. aeruginosa* (Hughes et al., 1997; Shaw et al., 1995; Zughai et al., 1999). Since this amplified response may fail to clear the infection by *B. cepacia*, the inflammation will perpetuate itself with subsequent cycles of phagocytosis and activation of macrophages, resulting in progressive lung tissue damage.

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