Burkholderia mallei expresses a unique lipopolysaccharide mixture that is a potent activator of human Toll-like receptor 4 complexes

Summary

Burkholderia mallei, the aetiologic agent of glanders, causes a variety of illnesses in animals and humans ranging from occult infections to acute fulminating septicaemias. To better understand the role of lipopolysaccharide (LPS) in the pathogenesis of these diseases, studies were initiated to characterize the structural and biological properties of lipid A moieties expressed by this organism. Using a combination of chemical analyses and MALDI-TOF mass spectrometry, B. mallei was shown to express a heterogeneous mixture of tetra- and penta-acylated lipid A species that were non-stoichiometrically substituted with 4-amino-4-deoxy-arabinose residues. The major penta-acylated species consisted of bisphosphorylated d-glucosamine disaccharide backbones possessing two amide linked 3-hydroxyhexadecanoic acids, two ester linked 3-hydroxytetradecanoic acids [C14:0(3-OH)] and an acyloxyacyl linked tetradecanoic acid, whereas, the major tetra-acylated species possessed all but the 3\textsuperscript{1}H11032-linked C14:0(3-OH) residues. In addition, although devoid of hexa-acylated species, B. mallei LPS was shown to be a potent activator of human Toll-like receptor 4 complexes and stimulated human macrophage-like cells (THP-1 and U-937), monocyte-derived macrophages and dendritic cells to produce high levels of TNF-\textgreek{alpha}, IL-6 and RANTES. Based upon these results, it appears that B. mallei LPS is likely to play a significant role in the pathogenesis of human disease.

Introduction

Burkholderia mallei, the aetiologic agent of glanders, is a non-motile, aerobic, facultative-intracellular Gram-negative bacillus that is primarily responsible for disease in horses, mules, donkeys and occasionally humans (Howe and Miller, 1947; Redfearn et al., 1966; Yabuuchi et al., 1992; Srinivasan et al., 2001). In horses, glanders presents as chronic or acute illnesses characterized primarily by lung involvement, ulcerative nasal and tracheal lesions and visceral abscess formation. In contrast, farcy, the cutaneous form of the disease, is characterized by the presence of nodules and ulcerative skin lesions on the extremities of the animals (Redfearn et al., 1966; Howe et al., 1971; Bartlett, 2005). The clinical progression of human glanders is similar to that observed in solipeds and may manifest as chronic or acute localized infections, acute pulmonary infections or fulminating septicaemias. Although rare, human infections are thought to be acquired via the inoculation of mucosal or cutaneous tissues with aerosols or secretions from diseased animals. In the absence of chemotherapeutic intervention human glanders is invariably fatal (Dance, 1996; Bartlett, 2005). Due to this, the Centers for Disease Control and Prevention have listed B. mallei as a category B biological threat agent (Rotz et al., 2002). Currently, there are no human or veterinary vaccines commercially available for immunization against these diseases.

Although glanders is one of the oldest of known infectious diseases, the molecular mechanisms by which B. mallei causes disease remain poorly defined (Wilson, 1981). Recently, however, a number of groups have demonstrated that quorum sensing, type three secretion system and capsular polysaccharide (CPS) mutants are attenuated in their ability to cause disease in mice, golden hamsters and/or miniature horses (DeShazer et al., 2001; Lopez et al., 2003; Ulrich and DeShazer, 2004; Ulrich et al., 2004). In addition, we have previously shown that B. mallei strains expressing rough lipopolysaccharide
Lipopolysaccharides, commonly referred to as endotoxins, are a major component of Gram-negative cell envelopes (Burns et al., 2006). The ‘barrier function’ provided by bacterial outer membranes is largely due to the presence of these molecules (Nikaido, 2003). Bacterial strains expressing a ‘smooth’ phenotype synthesize LPS antigens that are composed of three covalently linked domains: an O-antigen, a core region and a lipid A moiety (Raetz and Whitfield, 2002). Lipid A, which normally functions as the hydrophobic membrane-anchor component of LPS, is also the domain responsible for stimulating a variety of pathophysiological responses in mammals (Alexander and Rietschel, 2001; Backhed et al., 2002). In general, bisphosphorylated hexa-acylated lipid A species possessing C12 and C14 fatty acids represent the most biologically active forms of the molecule (Erridge et al., 2002; Raetz and Whitfield, 2002; Caroff and Karibian, 2003). Lipid A moieties that deviate from this pattern often demonstrate a significant decrease in endotoxic activity (Alexander and Rietschel, 2001; Backhed et al., 2001; 2003).

Host recognition of LPS antigens by mammalian cells is mediated through interactions with Toll-like receptor 4 (TLR4) complexes. These complexes, consisting of TLR4 receptors and the MD2 and CD14 coreceptors enable a variety of cell types to rapidly detect the presence of Gram-negative bacteria in host tissues (Poltorak et al., 1998; da Silva Correia et al., 2001; Miller et al., 2005). Monocytes and macrophages, for instance, when activated by LPS through TLR4 complexes secrete pro-inflammatory cytokines such as TNF-α, IL-1β and IL-6 which augment innate immune responses against invading microbes. In excess, however, these cytokines can cause multiple organ failure, shock and death (Zanotti et al., 2002). Although studies have examined some structural aspects of B. mallei LPS, none have actually reported on a potential immunopathological role for this molecule (Pitt et al., 1992; Anuntagool and Sirisinha, 2002; Woods et al., 2002). In order to address this issue, we have utilized a combination of genetic, chemical, physical and immunological approaches to further characterize the LPS expressed by this bacterial pathogen. The main objective of this study was to develop a better understanding of the potential role of endotoxin in the pathogenesis of disease caused by this organism.

Results

Characterization of purified LPS antigens

In order to obtain highly purified LPS for use in this study, endotoxin was isolated from B. mallei GM3773 (CPS mutant) as well as ATCC 23344 (wild-type strain) using a modified enzyme hot aqueous-phenol procedure. SDS-PAGE and silver stain analysis of the LPS isolated from the CPS mutant demonstrated that the preparation consisted of a mixture of both rough and smooth species. Further examination of the gel revealed no apparent structural differences between the LPS species isolated from the CPS mutant and wild-type strain (Fig. 1A). Similar results were also observed by immunoblot analysis (data not shown). Additional analysis of the samples by enzyme-linked immunosorbent assays (ELISA), however, demonstrated that while the GM3773 preparation contained no detectable levels of the CPS, the LPS isolated from ATCC 23344 was contaminated with this antigen (Fig. 1B). Based upon these findings, as well as those from the chemical and MALDI-TOF-MS analyses (see below), we opted to utilize the LPS isolated from GM3773 to determine the biological activities associated with B. mallei endotoxin.

Composition and chemical analyses of lipid A

Lipid A samples were isolated from the purified LPS preparations via mild acid hydrolysis. Glycosyl composition analysis of lipid A obtained from B. mallei GM3773 LPS confirmed the presence of d-glucosamine (GlcN) in the sample. Fatty acid composition analysis of the lipid A sample also revealed the presence of tetradecanoic (C14:0), 3-hydroxytetradecanoic [C14:0(3-OH)], hexadecanoic (C16:0) and 3-hydroxyhexadecanoic acids [C16:0(3-OH)] in the molar ratios of ~1.0:1.6:0.2:1.9

![Fig. 1. Analysis of purified LPS preparations.](image-url)
respectively. In addition, \(O\)-linked fatty acid analyses demonstrated that the C14:0, C14:0(3-OH) and C16:0 residues were ester linked indicating that the C16:0(3-OH) residues were amide linked. Similar results were also obtained for the lipid A antigens isolated from the \(B. mallei\) ATCC 23344 LPS.

MALDI-TOF-MS analyses of lipid A

The negative-ion MALDI-TOF mass spectrum of the purified \(B. mallei\) GM3773 lipid A preparation revealed a complex pattern of molecular ion peaks indicative of a heterogeneous mixture of species (Fig. 2A). Utilizing the information obtained from chemical and composition analyses in combination with the MS data, the molecular ion peaks were assigned as follows. The four main series of ion peaks identified were representative of a combination of tetra-acylated (\(m/z\) 1448 and 1579) and penta-acylated (\(m/z\) 1675 and 1806) species (Table 1). These species were also present in significant quantities as sodium adducts (\(\Delta m/z + 22\)). The ion at \(m/z\) 1448 (species 1) was consistent with a tetra-acylated bisphosphorylated GlcN disaccharide backbone possessing C14:0 and C14:0(3-OH) residues in ester linkage and two C16:0(3-OH) residues in amide linkage. The ion at \(m/z\) 1579 (species 2) was representative of species 1 modified with a 4-amino-4-deoxy-arabinose (Ara4N) residue (\(\Delta m/z + 131\)). The ion at \(m/z\) 1675 (species 3) was consistent with a penta-acylated bisphosphorylated GlcN disaccharide backbone possessing one C14:0 residue and two C14:0(3-OH) residues in ester linkage and two C16:0(3-OH) residues in amide linkage. The ion at \(m/z\) 1806 (species 4) was representative of species 3 modified with an Ara4N residue. The minor ion peak at \(m/z\) 1959 (species 5) was representative of a species 3 sodium adduct modified with two Ara4N residues. Taking into account the results of the fatty acid

![Figure 2](image)

**Table 1.** Main MALDI-TOF-MS negative-ion peaks and the proposed interpretation of the fatty acid, phosphate and carbohydrate substituents on the \(B. mallei\) GM3773 lipid A backbone.

| Lipid A species\(^a\) | Observed ion (\(m/z\)) | Calculated ion (Da) | Acyl substitution | Predicted fatty acid, phosphate and Ara4N composition |
|-----------------------|------------------------|---------------------|------------------|-----------------------------------------------|
| 1                     | 1448                   | 1445.4              | Tetra            | C14:0(3-OH), 2 \(\times\) C16:0(3-OH), C14:0, 2 \(\times\) P |
| 2                     | 1579                   | 1576.5              | Tetra            | C14:0(3-OH), 2 \(\times\) C16:0(3-OH), C14:0, 2 \(\times\) P, Ara4N |
| 3                     | 1675                   | 1671.3              | Penta            | 2 \(\times\) C14:0(3-OH), 2 \(\times\) C16:0(3-OH), C14:0, 2 \(\times\) P, Ara4N |
| 4                     | 1806                   | 1802.3              | Penta            | 2 \(\times\) C14:0(3-OH), 2 \(\times\) C16:0(3-OH), C14:0, 2 \(\times\) P, Ara4N |

\(a\). Sodium adducts (\(\Delta m/z + 22\)) are not listed.
composition analyses, the GM3773 preparation appeared to contain similar ratios of the tetra- and penta-acylated species. Similar results were obtained for LPS isolated from \textit{B. mallei} ATCC 23344 confirming that lipid A species expressed by the wild type and CPS mutant strains were identical to one another (Fig. 2B).

The positive-ion MALDI-TOF mass spectrum of the purified \textit{B. mallei} GM3773 lipid A preparation demonstrated the presence of a number of diagnostic oxonium ions (m/z 984, 956, 736 and 730) arising from the cleavage of the glycosidic linkage between the reducing and non-reducing GlcN residues of the lipid A backbone. The major oxonium ion observed at m/z 956 was consistent with the sodium adduct of a non-reducing GlcN possessing one ester linked C14:0 residue, one ester linked C14:0(3-OH) residue and one amide linked C16:0(3-OH) residue. In contrast, the minor ion observed at m/z 984 was consistent with a non-reducing GlcN possessing one ester linked C14:0 residue, one ester linked C14:0(3-OH) residue and one amide linked C16:0(3-OH) residue (Fig. 3). In addition, the major oxonium ion observed at m/z 730 was consistent with the sodium adduct of a non-reducing GlcN possessing an amide linked C16:0(3-OH) residue esterified by a C14:0 residue while the minor ion observed at m/z 736 was consistent with a non-reducing GlcN possessing an amide linked C16:0(3-OH) residue esterified by a C16:0 residue (data not shown). Based upon the results of these analyses the proposed structures of the major lipid A species identified in this study are depicted in Fig. 4.

\textbf{Burkholderia mallei LPS is a potent activator of hTLR4 complexes}

Lipopolysaccharide can activate a variety of cell types through interactions with TLR4 complexes (Fujihara \textit{et al.}, 2003). The magnitude of cell activation, however, is greatly influenced by endotoxin structure (Backhed \textit{et al.}, 2003). In order to characterize the ability of \textit{B. mallei} LPS to function as a hTLR4 agonist, HEK 293 cells expressing hMD2-CD14, hTLR2-CD14 or hTLR4-MD2-CD14 were stimulated with varying concentrations of endotoxin and the culture supernatants were assayed for IL-8. The results of these studies demonstrated that when stimulated with the \textit{B. mallei} LPS, the HEK 293-hTLR4/MD2-CD14 cells produced dose-dependent IL-8 responses that were similar to those observed when stimulated with the \textit{Escherichia coli} O55:B5 LPS control. In contrast, the HEK 293-hTLR2/CD14 cells secreted significantly lower levels of IL-8 when stimulated with identical concentrations of the LPS samples (Fig. 5). In addition, neither of the \textit{E. coli} or \textit{B. mallei} LPS preparations stimulated responses from the HEK 293-hMD2-CD14 cells at any of the concentrations tested (0.1–100 ng ml$^{-1}$; data not shown). Based upon the results of these assays, \textit{B. mallei} LPS was shown to be a potent activator of hTLR4-complexes.

\textbf{Burkholderia mallei endotoxin activates human macrophages and dendritic cells}

Antigen presenting cells (APCs) such as macrophages and dendritic cells recognize bacteria through TLRs and other pattern recognition receptors. Stimulation of APCs through these receptors activates signal transduction cascades that culminate in the production of cytokines, chemokines and other costimulatory molecules. To test the ability of \textit{B. mallei} LPS to activate human APCs,
macrophage-like THP-1 cells were stimulated with 10 ng ml\(^{-1}\) of LPS isolated from GM3773 and the culture supernatants were assayed for the production of specific cytokines and chemokines. When stimulated with GM3773 LPS, THP-1 cells produced TNF-\(\alpha\), IL-6 and RANTES responses that were similar to those detected using \(E.\ coli\) O55:B5 LPS as a control (Fig. 6). Similar trends were also observed when the macrophage-like U-937 cells were stimulated with \(B.\ mallei\) LPS (data not shown). In addition, studies demonstrated that when MDM were stimulated with \(B.\ mallei\) LPS, they produced high levels of TNF-\(\alpha\), IL-6 and RANTES (Fig. 7). Furthermore, when stimulated with \(B.\ mallei\) LPS, MDDC secreted high levels of TNF-\(\alpha\), IL-6, RANTES and IP-10 in a CD14-dependent manner (Fig. 8). Taken together, these results indicated that \(B.\ mallei\) LPS was a potent activator of human APCs and was capable of stimulating cytokine and chemokine responses that were consistent with the activation of both hTLR4-MyD88-dependent (TNF-\(\alpha\) and IL-6) and -independent (RANTES and IP-10) signalling pathways (Kawai \textit{et al}., 2001; Re and Strominger, 2001; Fitzgerald \textit{et al}., 2003; Zughaier \textit{et al}., 2005).

**Discussion**

Lipopolysaccharide antigens expressed by \textit{Burkholderia pseudomallei} and \textit{B. mallei} are typically isolated from whole cell pellets using a modified hot aqueous-phenol extraction procedure (Brett and Woods, 1996; Burtnick \textit{et al}., 2002; Brett \textit{et al}., 2003; Perry \textit{et al}., 1995). Unfortunately, the smooth LPS species expressed by these...
Fig. 8. Production of TNF-α, IL-6, RANTES and IP-10 by LPS stimulated MDDC is CD14-Dependent. Prior to stimulation, the MDDC were incubated for 30 min with an anti-CD14 antibody (black bars) or a non-specific isotype control (grey bars). The cells were then stimulated for 24 h with 10 ng ml⁻¹ of B. mallei GM3773 or E. coli O55:B5 LPS following which the culture supernatants were assayed for the production of TNF-α, IL-6, RANTES or IP-10. Unstimulated cells were included as a control. Values represent the means of two readings. The figures are representative of three independent experiments.

organisms partition into the phenol phase rather than the aqueous phase following extraction so that isolating homogeneous preparations of these molecules can be a challenge (Perry et al., 1995). For example, the high molecular weight CPS expressed by virulent isolates of B. mallei and B. pseudomallei partitions with the LPS into the phenol phase so that size exclusion chromatography is required to separate these carbohydrate antigens. In order to simplify the task of obtaining highly purified preparations of LPS for use in this study, we constructed B. mallei GM3773. By using this CPS mutant, we were able to isolate LPS antigens that were structurally identical to those expressed by the wild-type strain yet were free from any contaminating CPS. In addition, because B. mallei CPS mutants are attenuated for virulence (DeShazer et al., 2001), working with strain GM3773 minimized exposure to the virulent type strain.

Recent studies by Silipo et al. have demonstrated that the human pathogen, Burkholderia cepacia BTS7, expresses a complex mixture of tetra- and penta-acylated lipid A species that are non-stoichiometrically substituted with Ara4N residues (Silipo et al., 2005). Structural analysis of the lipid A moieties isolated from this organism indicated that the penta-acylated species with the highest mass consisted of disaccharide backbones that were substituted with two phosphoryl-arabinosamine residues [Arap4N-L-β-1-4-P-β-d-GlcP-N-(1'-6)-α-d-GlcP-N-1-P-1-[β-L-Arap4N]] and possessed two ester linked C14:0(3-OH) chains, two amide linked C16:0(3-OH) chains and an acyloxyacyl linked C14:0 residue at the C-3 position of the 2′-C16:0(3-OH) chain. In addition, studies confirmed that the tetra-acylated species with the highest mass were identical to the penta-acylated species except for the lack of the 3′-ester linked C14:0(3-OH) chain. By utilizing a combination of chemical analyses and MALDI-TOF-MS in the present study, we have been able to demonstrate that B. mallei also expresses a heterogeneous mixture of tetra- and penta-acylated lipid A species that appear to be non-stoichiometrically substituted with Ara4N residues. In addition, all of the major molecular ion peaks identified in this study were also observed during the analysis of the B. cepacia BTS7 lipid A moieties. Interestingly, however, none of the minor B. mallei lipid A species predicted to bear acyloxyacyl linked C16:0 residues at the 2′-position that were identified during our studies were shown to be expressed by B. cepacia BTS7. At present, the significance of this phenomenon remains unknown.

Previous studies indicate that lipid A antigens consisting of 1,4′-bisphosphorylated β(1′-6)-linked β-glucosamine backbones substituted with two (R)-3-acyloxyacyl and two (R)-3-hydroxyacyl residues in either the 4 + 2 (E. coli-like) or 3 + 3 (Neisseria meningitidis-like) configuration represent the optimal structures for activating human and mammalian TLR4-complexes (Alexander and Rietschel, 2001; Erridge et al., 2002; Dixon and Darveau, 2005; Zughaier et al., 2005; 2006). In addition, it has been shown that lipid A antigens deviating from these patterns tend to exhibit reduced biological activities in comparison to these molecules (Kumazawa et al., 1988; Nakatsuka et al., 1989; Takada and Kotani, 1989; Backhed et al., 2003). Consistent with these findings, Backhed et al. have demonstrated that LPS antigens isolated from wild-type strains of E. coli stimulate significantly higher levels of IL-8 from human bladder carcinoma cells (T24 cells) than penta-acylated LPS antigens isolated from waaN (lpxM) mutants (Backhed et al., 2001). Interestingly, Ernst et al. have also shown that LPS preparations obtained from Pseudomonas aeruginosa clinical isolates expressing
both penta- and hexa-acylated species stimulate more robust pro-inflammatory responses from human macrophages and endothelial cells than LPS preparations purified from clinical isolates expressing only penta-acylated species (Ernst et al., 1999; 2003; Hajjar et al., 2002). In contrast to established structure/function paradigms, however, our studies indicate that although devoid of hexa-acylated species, B. mallei LPS is a potent hTLR4 agonist and is capable of stimulating human APCS in a manner similar to E. coli LPS. Based upon these results, it appears that by optimizing the length, number and order of their lipid A fatty acids, organisms such as B. mallei and B. cepacia are able to synthesize a mixture of tetra and penta-acylated LPS antigens whose individual or combined biological properties mimic those typically associated with hexa-acylated LPS preparations (Zughaier et al., 1999; De Soyza et al., 2004; Bamford et al., 2006).

Capping of lipid A phosphates represents an important molecular mechanism utilized by Gram-negative pathogens to resist the bactericidal effects of cationic antimicrobial peptides (CAPs) produced by host innate immune responses (Vaara, 1992; Gunn et al., 1998; 2000; Gunn, 2001; Trent, 2004). By modifying lipid A phosphates with amine-containing substituents like Ara4N and phosphoethanolamine, pathogens such as Salmonella typhimurium and Yersinia pestis can lower their affinity for CAPs and cationic antibiotics by decreasing the overall negative charge of their cell surfaces (Zhou et al., 2001; Kawahara et al., 2002; Trent, 2004). In S. typhimurium, modification of lipid A moieties with Ara4N residues is governed by the PmrA/PmrB two-component system which regulates the expression of genetic loci necessary for the biosynthesis of Ara4N (Soncini and Groisman, 1996). Transcription of PmrA-regulated loci is induced by ferric iron (Fe³⁺), which is sensed by the cognate sensor-PmrB, and by low concentrations of magnesium (Mg²⁺), which requires not only PmrA and PmrB but also PhoP, PhoQ and PmrD (Groisman, 2001; Winfield et al., 2005). Consistent with current observations that B. mallei lipid A moieties are modified with Ara4N residues, bioinformatics analyses have revealed that a number of genetic loci (ugd, arrA, arrB, amC and amT) known to be involved in the biosynthesis of Ara4N and its transfer to S. typhimurium and E. coli lipid A phosphates are also present in the genome of B. mallei ATCC 23344 (Trent et al., 2001; Nierman et al., 2004). Curiously, however, although a putative pmrA homologue was identified during these analyses, no obvious pmrB, pmrD, phoP or phoQ homologues were found. Studies are ongoing in order to characterize the mechanism(s) by which B. mallei modifies lipid A moieties with Ara4N residues and the potential significance of these modifications with regards to CAP resistance.

In addition to remodelling the charge characteristics of their LPS antigens, S. typhimurium, P. aeruginosa and Y. pestis are also capable of modifying the acylation patterns of their lipid A moieties in response to environmental changes such as temperature, pH and cation concentrations (Ernst et al., 1999; 2006; Zhou et al., 2001; Kawahara et al., 2002; Rebeil et al., 2004; 2006; Tran et al., 2005). In particular, modifications resulting in the synthesis of tetra-acylated species have been shown to significantly affect virulence phenotypes as well as play an important role in modulating mammalian TLR4-dependent immune responses (Ernst et al., 1999; Hajjar et al., 2002; Backhed et al., 2003; Rebeil et al., 2004). Recently, Rebeil et al. demonstrated that when grown at 21°C, Y. pestis expressed a heterogeneous mixture of LPS antigens in which hexa-acylated species predominated, while at 37°C, the organism expressed a mixture of LPS species in which tetra-acylated forms of the antigen predominated (Rebeil et al., 2004). During the study, it was also shown that LPS preparations isolated from Y. pestis grown at 37°C stimulated significantly less TNF-α production from human peripheral blood monocytes than did LPS preparations purified from the organism when grown at 21°C. Based upon these findings it was suggested that the ability of Y. pestis to modify LPS acylation patterns in response to changing temperature may enhance its ability to survive during transitions from its flea vector to mammalian hosts. More recently, Reife et al. demonstrated that tetra-acylated LPS species expressed by Porphyromonas gingivalis are potent TLR4 antagonists and that by altering the relative amounts these antigens in its outer membrane the organism may be capable of modulating host immune responses (Reife et al., 2006). Interestingly, studies have shown that LPS preparations from a clinical isolate of Burkholderia cenocepacia expressing only penta-acylated species stimulated stronger TNF-α and IL-6 responses from U-937 cells than from LPS preparations purified from a clinical isolate of Burkholderia multivorans expressing both tetra- and penta-acylated species (De Soyza et al., 2004).

In the present study we have demonstrated that B. mallei expresses a mixture of tetra- and penta-acylated LPS species that in combination exhibit some unique biological properties. In order to fully characterize the significance of this heterogeneity, however, further studies will be required to elucidate the specific activities of these individual LPS species. Whether or not changes in environmental conditions/stimuli modulate LPS acylation patterns and biological activity also remains to be experimentally defined. Studies are ongoing to address these important issues.

**Experimental procedures**

**Bacterial strains and growth conditions**

The bacterial strains used in this study are described in Table 2. Escherichia coli strains were grown at 37°C on LB...
This study

B. mallei

et al.

the following concentrations: 10

with 4% glycerol. When appropriate, antibiotics were used at

were grown at 37°C in LBL broth or on LBL agar supplemented

were inoculated with

Lipopolysaccharide purification and lipid a isolation

PCR using the wcbB-up and MB1ori-P1 primer pair (Table 2).

plasmid into the ATCC 23344 chromosome was verified via

Chemically competent

mosomal DNA was isolated using a Wizard™ Genomic DNA

purified using a QIAprep Spin Miniprep Kit (Qiagen). Chro-

ligase products were purified using a QIAquick Gel Extrac-

Polymerase chain reaction (PCR) and restriction enzyme

Recombinant DNA techniques

DNA manipulations were performed using standard methods. Polymerase chain reaction (PCR) and restriction enzyme
digested products were purified using a QIAquick Gel Extraction Kit (Qiagen). The plasmids described in Table 2 were

transformed using standard methods.

Mutant construction

The B. mallei GM3773 CPS mutant was constructed essen-
tially as previously described (DeShazer et al., 2001; Burtnick et al., 2002). Briefly, a 547 bp fragment internal to the wcbB
allele was PCR amplified from purified B. mallei ATCC 23344 chromosomal DNA using the wcbB-if and wcbB-ir primer pair (Table 2); the BsaI/EcoRI and BsaI/NheI restriction sites
in the linker regions are underlined. The amplified product
was then cloned into pGSV3. The resulting suicide plasmid,
pGMwcbB, was mobilized into ATCC 23344 via conjugal mat ing using E. coli S17-1. Site specific recombination of the
plasmid into the ATCC 23344 chromosome was verified via
PCR using the wcbB-up and MB1ori-P1 primer pair (Table 2).

Lipopolysaccharide purification and lipid a isolation

LBL broth inoculated with B. mallei ATCC 23344 or GM3773
was incubated overnight at 37°C with vigorous shaking. Cell
pellets were subsequently obtained by centrifugation and
extracted using a modified hot aqueous-phenol procedure

Lennox (LBL) agar or in LBL broth (Difco). B. mallei strains
were grown at 37°C in LBL broth or on LBL agar supplemented
with 4% glycerol. When appropriate, antibiotics were used at
the following concentrations: 10 \mu g \text{ml}^{-1} gentamicin or 15 \mu g \text{ml}^{-1} polymyxin B for E. coli and 5 \mu g \text{ml}^{-1} gentamicin for
B. mallei. Bacterial stocks were maintained at -80°C as 20% glycerol suspensions. All studies utilizing viable B. mallei were
conducted under biosafety level three containment conditions.

Recombinant DNA techniques

DNA manipulations were performed using standard methods. Polymerase chain reaction (PCR) and restriction enzyme
digested products were purified using a QIAquick Gel Extraction Kit (Qiagen). The plasmids described in Table 2 were
purified using a QIAprep Spin Miniprep Kit (Qiagen). Chromosomal DNA was isolated using a Wizard™ Genomic DNA
Isolation Kit (Promega). Chemically competent E. coli were
transformed using standard methods.

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pellets were subsequently obtained by centrifugation and
extracted using a modified hot aqueous-phenol procedure

(Perry et al., 1995). Following extraction, the resulting phenol
and aqueous phases were combined and dialysed in distilled water to remove the phenol. The dialysates were then clari-
fied by centrifugation and concentrated by lyophilization.
The crude preparations were solubilized (20 \mu g \text{ml}^{-1}) in RD buffer [10 mM Tris-HCl (pH 7.5), 1 mM MgCl₂, 1 mM CaCl₂,
50 \mu g \text{ml}^{-1} RNase A and 50 \mu g \text{ml}^{-1} DNase I] and incubated
for 3 h with shaking at 37°C. Proteinase K was then added to
a final concentration of 50 \mu g \text{ml}^{-1} and the digests were incu-
bpated for an additional 3 h at 60°C. The enzymatic digests
were clarified by centrifugation and the supernatants filter
sterilized. LPS was then isolated from the supernatants as precipitated gels following three rounds of ultracentrifugation
at 100 000 \text{g} and 4°C. After the final spin, the gel-like pellets
were resuspended in pyrogen-free water and lyophilized. To
remove contaminating phospholipids, the lyophilized LPS
samples were repeatedly extracted with 90% EtOH. Lipid A
was isolated from the purified LPS samples via mild acid hydrolysis. Briefly, LPS was hydrolysed for 4 h at 95°C in
0.1 M NaOAc (pH 4.5). The lipid A fraction was then removed
by extraction with chloroform : methanol (2:1) and dried
under a gentle stream of air and lyophilized.

SDS-PAGE, immunoblotting and ELISA analyses

Purified samples were solubilized in 1x SDS-PAGE sample buffer and heated to 100°C for 5 min prior to electrophoresis
on 12% Express Gels (ISC BioExpress). The LPS was visu-
alyzed by silver staining as previously described (Fomsgaard et al., 1990). Immunoblot analysis was performed as previ-
ously described (Burtnick et al., 2002). ELISA were per-
formed essentially as previously described (DeShazer et al., 1998). The primary antibodies used in this study were the
O-antigen specific 3D11 monoclonal antibody (mAb; Research
Diagnostic) and the CPS specific MCA147 mAb (Reckseidler-Zenteno et al., 2005). The secondary antibodies
used were anti-mouse IgG or IgM horseradish peroxidase
conjugates (Sigma).

Glycosyl composition analysis

Per-O-trimethylsilyl (TMS) derivatives of monosaccharide
methyl glycosides produced from the acidic methanolsysis of

| Strain, plasmid or primer | Description | Source or reference |
|---------------------------|-------------|---------------------|
| E. coli TOP10             | High efficiency transformation: Gm<sup>s</sup> | Invitrogen          |
| S17-1                     | Mobilizing strain; transfer genes of RP4 integrated in chromosome: Gm<sup>s</sup>, Pm<sup>r</sup> | Simon et al. (1983) |
| B. mallei ATCC 23344     | Type strain; isolated in 1944 from a human case of glanders: Gm<sup>s</sup>, Pm<sup>r</sup> | Yabuuchi et al. (1992) |
| GM3773                    | ATCC 23344::pGMwcbB: Gm<sup>r</sup>, Pm<sup>r</sup> | This study          |
| Plasmids                  |             |                     |
| pGSV3                     | Mobilizable suicide vector: Gm<sup>r</sup> | DeShazer et al. (2001) |
| pGMwcbB                   | pGSV3 containing a 547 bp PCR fragment internal to wcbB: Gm<sup>r</sup> | This study          |
| Primers                   |             |                     |
| wcbB-if                   | 5′-CCCAACGGTCTCAATTGCCATTTGCAGGACTAGCCGCGC-3′ | This study          |
| wcbB-ir                   | 5′-CCCAACGGTCTCAGTTCAGGACGTTCCTGATTCCAGCGGCTCG-3′ | This study          |
| wcbB-up                   | 5′-ACGAAGTCCGCCGTTTGATACAGC-3′ | This study          |
| MB1ori-P1                 | 5′-GAAGATCTTTTGATCTTTTGACGG-3′ | This study          |

Table 2. Bacterial strains, plasmids and primers used in this study.
the lipid A samples were analysed by gas chromatography/mass spectrometry (GC/MS). The methyl glycosides were prepared by heating the dried lipid A samples in acidified methanol (1 M HCl) at 80°C for 18–22 h. Following this, the samples were re-N-acetylated with pyridine and acetic anhydride in order to facilitate the detection of amino sugars. The samples were then per-O-trimethylsilylated via treatment with Tri-Sil (Pierce) at 80°C for 30 min. GC/MS analysis of the TMS methyl glycosides was performed on a HP 5890 GC interfaced with a 5970 MSD using a All Tech EC-1 fused silica capillary column (30 m x 0.25 mm id).

Fatty acid analysis

Methyl ester derivatives of the fatty acids generated by acidic methanolsis of the lipid A samples, were dried, resuspended in hexane and analysed by GC/MS to determine total fatty acid content. In order to detect O-linked fatty acids, the lipid A samples were suspended in alkaline methanol (0.25 M NaOMe), incubated overnight at 37°C and then neutralized with an equal volume of 0.25 M HCl. The fatty acid methyl esters were then extracted with 0.15 volumes of chloroform and analysed by GC/MS after a partial drying under a gentle stream of dry air. In addition to using the All Tech EC-1 fused silica capillary column, GC was also performed using a Supelco SP 2380 capillary column equipped with a FID detector and a bimodal temperature gradient starting at 130°C and increasing at 4°C per minute to 190°C followed by a 1 min hold at 190°C and a 17°C per minute gradient to 260°C and 3 min hold.

Mass spectrometry

Matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF-MS) was performed in the negative-ion mode on a Voyager DE-STR mass spectrometer (Applied Biosystems). Lipid A samples in chlorofrom : methanol (2:1) were mixed on target with 2,5-dihydroxybenzoic acid (Applied Biosystems). Lipid A samples in chloroform : methanol (1 M HCl) at 80°C for 18–22 h. Following this, the samples were re-N-acetylated with pyridine and acetic anhydride in order to facilitate the detection of amino sugars. The samples were then per-O-trimethylsilylated via treatment with Tri-Sil (Pierce) at 80°C for 30 min. GC/MS analysis of the TMS methyl glycosides was performed on a HP 5890 GC interfaced with a 5970 MSD using a All Tech EC-1 fused silica capillary column (30 m x 0.25 mm id).

Cell culture

Human embryonic kidney 293 (HEK 293) cells transfected with human MD2-CD14 (HEK 293-hMD2-CD14; Invivogen), human TLR2/CD14 (HEK 293-hTLR2/CD14; Invivogen) or human TLR4/MD2-CD14 (HEK 293-hTLR4/MD2-CD14; Invivogen) were maintained at 37°C with 5% CO2 in RPMI 1640 medium (Invitrogen) supplemented with 10% HI FBS and 100 µg ml⁻¹ Normocin™ (Invivogen). To differentiate the monocytes into macrophage-like cells, the U-937 and THP-1 cells were transferred into 24-well tissue culture plates at 1 × 10⁶ cells well⁻¹ and incubated at 37°C with 5% CO2 for 36 h in the presence of 50 ng ml⁻¹ phorbol myristate acetate. Prior to stimulation with LPS, the differentiated cells were washed with Hanks balanced salt solution (Invitrogen).

Human monocyte-derived macrophage and dendritic cells

Human peripheral blood mononuclear cells were isolated from buffy coats by centrifugation through a Ficoll-Paque Plus density gradient (Amersham Pharmacia Biotech). Cells were enriched for monocytes (CD14⁺ cells) using a RossetteSep Monocyte Enrichment Kit (Stem Cell Technologies). Macrophages (MDM) were obtained by resuspending the monocytes (1 × 10⁶ cells ml⁻¹) in Mcf medium (RPMI 1640, 10% HI FBS, 50 ng ml⁻¹ recombinant human M-CSF; Peprotech) and then culturing them at 37°C with 5% CO2 for 7 days with the addition of fresh cytokine on day three. On day seven, the adherent cells were washed once with phosphate-buffered saline (PBS) and then incubated on ice for 20 min in PBS. The cells were then harvested by gentle scraping and centrifugation (500 g), resuspended in Mcf medium and transferred into 24-well tissue culture plates at 1 × 10⁶ cells well⁻¹ prior to LPS stimulation. Monocyte-derived dendritic cells (MDDC) were isolated and the CD14 blocking studies were performed as previously described (Shannon et al., 2005).

Lipopolysaccharide stimulation assays and cytokine quantification

Prior to use, the E. coli O55:B5 (Sigma) and B. mallei GM3773 LPS samples were repeatedly extracted with 90% ETOH to remove any contaminating phospholipids (Zughaier et al., 2005). LPS stocks were prepared in pyrogen free water (Cambrex) and were quantified on a w/v basis. HEK 293 cell lines were stimulated for 6 h with varying concentrations (0.1–100 ng ml⁻¹) of GM3773 or O55:B5 LPS following which the culture supernatants were harvested and assayed for IL-8 production using Human IL-8 Chemiluminescent ELISA Kits (Endogen). THP-1, U-937, MDM and MDDC cells were stimulated for 24 h with 10 ng ml⁻¹ of GM3773 or O55:B5 LPS following which the culture supernatants were harvested and assayed for TNF-α, IL-6, RANTES and IP-10 production using human SearchLight Custom Multiplex Arrays (Pierce).

Acknowledgements

We are grateful to Donald Woods for providing B. mallei ATCC 23344 and the MCA147 mAb. We thank B. Joseph Hinnebusch for critical review of the manuscript. Hua Su is acknowledged for assistance with tissue culture. This research was supported in part by the Intramural Research Program of the NIH, National Institute of Allergy and Infectious Disease and the Department of Energy-funded (DE-FG09-93ER-20097) Center for Plant and Microbial Complex Carbohydrates.
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