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Increased Phagocytosis of Mycobacterium marinum Mutants Defective in Lipooligosaccharide Production

A STRUCTURE-ACTIVITY RELATIONSHIP STUDY*

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Background: Biosynthesis and functions of Mycobacterium marinum lipooligosaccharides (LOSs) remain elusive.

Results: M. marinum mutants expressing various LOS profiles were generated and used to infect macrophages and amoebae.

Conclusion: Deep LOS mutants are more efficiently phagocytosed than those lacking only LOS-IV.

Significance: Three novel biosynthetic genes and the effect of the LOS content in modulating uptake by phagocytes are reported.

Mycobacterium marinum is a waterborne pathogen responsible for tuberculosis-like infections in ectotherms and is an occasional opportunistic human pathogen. In the environment, M. marinum also interacts with amoebae, which may serve as a natural reservoir for this microorganism. However, the description of mycobacterial determinants in the early interaction with macrophages or amoebae remains elusive. Lipooligosaccharides (LOSs) are cell surface-exposed glycolipids capable of modulating the host immune system, suggesting that they may be involved in the early interactions of M. marinum with macrophages. Herein, we addressed whether LOS composition affects the uptake of M. marinum by professional phagocytes. Mutants with various truncated LOS variants were generated, leading to the identification of several previously uncharacterized biosynthetic genes (wbbL2, MMAR_2321, and MMAR_2331). Biochemical and structural approaches allowed resolving the structures of LOS precursors accumulating in this set of mutants. These strains with structurally defined LOS profiles were then used to infect both macrophages and Acanthamoebae. An inverse correlation between LOS completeness and uptake of mycobacteria by phagocytes was found, allowing the proposal of three mutant classes: class I (papA4), devoid of LOS and highly efficiently phagocytosed; class II, accumulating only early LOS intermediates (wbbL2 and MMAR_2331) and efficiently phagocytosed but less than class I mutants; class III, lacking LOS-IV (losA, MMAR_2319, and MMAR_2321) and phagocytosed similarly to the control strain. These results indicate that phagocytosis is conditioned by the LOS pattern and that the LOS pathway used by M. marinum in macrophages is conserved during infection of amoebae.

The hallmark of all members of the Mycobacterium genus is the presence of a unique and waxy cell envelope, which plays a key role in resistance to antimicrobial agents and in the pathobiology of mycobacterial infections by modulating the host immune response as well as phagocytic cell functions (1). This cell wall comprises mainly two types of lipids: (i) the very long-chain fatty acids, mycolic acids, covalently attached to the arabinogalactan/peptidoglycan backbone (2) and (ii) a vast panoply of structurally diverse and extractable compounds (3), including lipooligosaccharide, lipomannan, and related phosphatidylinositol mannosides; glycopeptidolipids; phthiocerol dimycocerosates and related phenolic glycolipids; triacylglycerols as well as different families of trehalose-based glycolipids, such as trehalose dimycolate, sulfolipids, di-, tri-, and pentaacyltrehalose; and lipooligosaccharides (LOSs). Although the structure and the distribution of these glycolipids have been well studied in various mycobacterial species, evidence for their participation in pathogenesis and virulence is now emerging (4).

LOSs have been found in many mycobacterial species, such as Mycobacterium kansasii, Mycobacterium gastri, Mycobacterium szulgai, the Mycobacterium canetti variant of Mycobacterium tuberculosis, and Mycobacterium marinum (5–10).

Acanthamoebae

* The abbreviations used are: LOS, lipooligosaccharide; Rha, rhamnose; MeRha, methylrhamnose; Xyl, xylose; OADC, oleic acid, albumin, dextrose, catalase; SCO, single crossover; DCO, double crossover; mut, mutated; m.o.i., multiplicity of infection; PYG, peptone-yeast-glucose; Hex, hexose; deHex, deoxyhexose; OS, oligosaccharide; Pent, pentose; Car, caryophylllose; PAT, pentaacyltrehalose.
Role of LOS in Mycobacterial Phagocytosis

However, among the different species, these glycolipids exhibit considerable structural variations in the glycan core as well as in the lipid moiety. Initial work identified the presence of four major LOSs in *M. marinum* that are designated LOS-I to LOS-IV (10). Partial characterization established the structure of LOS-1 as a triacylated 3-O-Me-Rhap-(1–3)-GlcP-(1–3)-GlcP-(1–4)-GlcP-(1–1)-GlcP. This common oligosaccharide moiety is sequentially modified by additional monosaccharides, thus giving rise to more polar LOS species. In particular, an α-D-Xylp further substitutes the glycan core in LOS-II, LOS-III, and LOS-IV. In addition, LOS-II possesses a terminal α-caryophylllose residue (α-3,6-dideoxy-4-C-(D-altro-1,3,4,5-tetrahydroxyhexyl)-D-xylp-hexopyranose), whereas two α-caryophylllose units are present in LOS-III and LOS-IV (11). Caryophylllose may be substituted by the related monosaccharide of LOS-IV decorating the second caryophylllose residue (α-6-deoxy-4-C-(D-altro-1,3,4,5-tetrahydroxyhexyl)-D-galactopyranose). The terminal monosaccharide of LOS-IV decorating the second caryophylllose residue has recently been identified as an unusual α-4-amino-4,6-dideoxy-Gal-(1–1)-GlcP-GlcP-GlcP.

Partial characterization established the structure of LOS-IV (10). This common oligosaccharide further substitutes the glycan core in LOS-II, LOS-III, and LOS-IV. In particular, an α-(1–1)-GlcP-GlcP-GlcP-GlcP. The terminal monosaccharide of LOS-IV decorating the second caryophylllose residue (α-6-deoxy-4-C-(D-altro-1,3,4,5-tetrahydroxyhexyl)-D-galactopyranose). The terminal monosaccharide of LOS-IV decorating the second caryophylllose residue has recently been identified as an unusual α-4-amino-4,6-dideoxy-Gal-(1–1)-GlcP-GlcP-GlcP-GlcP.

Albeit the structures of all LOSs from *M. marinum* now appear to be well described, the knowledge about the genes involved in LOS biogenesis is far from being complete, and until recently, only a few biosynthetic genes had been identified (10, 11, 13, 15, 16). However, in a recent search of additional components of the ESX-5 secretion system, a transposon screen in *M. marinum* for impaired PE_PGRS secretion (known as substrates of ESX-5) led to the identification of several mutants with transposon insertions in the LOS biosynthetic gene cluster (14). Intriguingly, in these LOS mutants, the PE_PGRS proteins appeared to be more firmly attached to the cell surface, thus putting forward an unexpected link between LOS and PE_PGRS release (14). This study not only brought to light a large variety of LOS biosynthetic genes whose disruption led to defects ranging from LOS-IV to complete LOS deficiency but also pointed out the transcriptional regulator WhiB4, which is located outside of the LOS region, as capable of regulating LOS production by controlling expression of several LOS biosynthetic genes (14).

LOSs are highly antigenic cell surface-exposed glycoconjugates and represent useful molecules to be targeted for serotyping in a given mycobacterial species. Early studies demonstrated that rough variants of *M. kansasii* lacking all LOSs are able to induce chronic systemic infections in mice, whereas smooth variants containing LOS are rapidly cleared from the organs of infected animals (17, 18), leading to the speculation that LOSs may be regarded as virulence factors by masking other cell wall-associated virulence factors. Studies performed with *M. marinum* mutants suggested that LOSs may play a role in sliding motility and biofilm formation (15). In vitro studies indicated that purified LOSs inhibit the secretion of TNF-α in LPS-stimulated human macrophages, supporting the view that these glycolipids are key effectors susceptible to interfere with the induction of a proinflammatory response (11). In addition, it was demonstrated that the terminal monosaccharide conferred to LOS-IV important biological functions, such as stimulation of ICAM-1 and CD40 expression at the macrophage cell surface (12). That LOS-IV plays an important role in pathogenesis and/or modulation of *M. marinum* virulence was corroborated by a recent study demonstrating that a LOS-IV-deficient mutant (inactivated in wecE) showed increased virulence in infected zebrafish embryos compared with the wild-type strain (14). However, despite one study based on a restricted panel of mutants and suggesting that LOSs participate in phagocytosis of *M. marinum* by macrophages (15), our understanding and the precise contribution of LOS subspecies with respect to the early events between *M. marinum* and macrophages remain unknown. Such structure-function relationship studies are particularly difficult to assess mainly because of the limited number of isogenic LOS mutant strains and the lack of precise structural data of LOS variants.

To extend our understanding of LOS biosynthesis and to examine the potential role of LOSs in the early interactions of *M. marinum* with professional phagocytes, we selected/generated a panel of *M. marinum* mutants expressing a wide diversity of truncated LOS variants. This set of structurally defined mutants was used to decipher the role of LOSs in phagocytosis of *M. marinum* by macrophages and free-living amoebae.

**EXPERIMENTAL PROCEDURES**

**Bacterial Strains and Growth Culture Conditions—Escherichia coli** Top-10 (Invitrogen) was used as the host for cloning and was grown in LB medium. Plasmid selection and maintenance were performed using ampicillin (100 μg·ml⁻¹), hygromycin (200 μg·ml⁻¹), and kanamycin (50 μg·ml⁻¹). The plasmids used in this study are listed and described in Table 1. *M. marinum* strain M isolated from a human patient as described previously (19) and the *MmaM*: *Tn*5370 transposon library (20) were grown/maintained in Sauton's broth medium at 30 °C or on Middlebrook 7H10 agar plates containing 10% oleic acid, albumin, dextrose, catalase (OADC) enrichment and supplemented with appropriate antibiotics. For selection, kanamycin (25 μg·ml⁻¹), hygromycin (80 μg·ml⁻¹), 5-bromo-4-chloro-3-indolyl β-D-galactopyranoside (X-Gal; 50 μg·ml⁻¹), or 2% sucrose were used. For polar lipid extractions and infection experiments, *M. marinum* strains were grown at 30 °C on Sauton's agar plates containing 10% OADC enrichment and recovered by scraping off the bacterial lawn.

**Screening of M. marinum Tn Library and Determination of the Transposon Insertion Site**—We have previously screened a *M. marinum Tn* library in *Dictyostelium discoideum* aiming to identify new cell wall-defective mutants (20). During the course of this screening, a *M. marinum* mutant carrying a *Tn*5370 insertion in *MMAR_2343/papA4* was identified and shown to be defective in LOS production (13). With the aim to identify new LOS biosynthetic genes and based on the observation that the *MMAR_2343/papA4* exhibited a rough morphotype, addi-
TABLE 1
Plasmids used in this study

| Plasmid                                      | Description                                                                 | Source/Ref. |
|----------------------------------------------|-----------------------------------------------------------------------------|-------------|
| Cloning vectors                              |                                                                            |             |
| pJE1.2/blunt                                 | PCR product cloning vector, Amp<sup>a</sup>                                 | Thermo Scientific |
| p2NIL                                        | Recombination vector, nonrepliacting in mycobacteria, Kan<sup>b</sup>         | 22          |
| pGOAL17                                      | Source of PacI marker cassette, (sucB, lacZ), Amp<sup>b</sup>                 | 22          |
| pMV306Hyg                                    | Mycobacterial integrating vector, Hyg<sup>a</sup>                            | MedImmune   |
| pMV261                                       | Source of isp60 promoter, Kan<sup>b</sup>                                   | 44          |
| Vectors used for gene replacement            |                                                                            |             |
| pGR2349                                      | p2NIL-based recombination vector carrying the 5’-end of M. marinum          | This study  |
|                                               | MMAR_2349 and its upstream flanking sequence (1434 bp) cloned next to the   |             |
|                                               | 3’-end of the gene and its downstream flanking sequence (907 bp), enriched   |             |
|                                               | with the Pacl cassette from pGOAL17, Kan<sup>b</sup>                          |             |
| pGR2331                                      | p2NIL-based recombination vector carrying the 5’-end of M. marinum          | This study  |
|                                               | MMAR_2331 and its upstream flanking sequence (1034 bp) cloned next to the   |             |
|                                               | 3’-end of the gene and its downstream flanking sequence (1277 bp), enriched  |             |
|                                               | with the Pacl cassette from pGOAL17, Kan<sup>b</sup>                          |             |
| pGR2321                                      | p2NIL-based recombination vector carrying the 5’-end of M. marinum          | This study  |
|                                               | MMAR_2321 and its upstream flanking sequence (996 bp) cloned next to the    |             |
|                                               | 3’-end of the gene and its downstream flanking sequence (1666 bp), enriched  |             |
|                                               | with the Pacl cassette from pGOAL17, Kan<sup>b</sup>                          |             |
| Vectors used for complementation             |                                                                            |             |
| pPown2349-1kb                                | PCR fragment harboring whole MMAR_2349 gene (768 bp) together                | This study  |
|                                              | with 1014-bp sequence upstream from the gene, carrying the gene promoter,   |             |
|                                              | cloned into the KpnI/XbaI site of the pMV306Hyg integrating vector, Hyg<sup>a</sup> |             |
| pPown2331                                    | PCR fragment harboring whole MMAR_2331 gene (801 bp) together                | This study  |
|                                              | with 603-bp sequence upstream from the gene, carrying the gene promoter,    |             |
|                                              | cloned into the XbaI/HindIII site of the pMV306Hyg integrating vector, Hyg<sup>a</sup> |             |
| pPHsp2321                                    | PCR fragment harboring whole MMAR_2321 gene (666 bp) together                | This study  |
|                                              | with 453-bp sequence of the isp60 promoter, cloned into the XbaI/            |             |
|                                               | HindIII site of the pMV306Hyg integrating vector, Hyg<sup>a</sup>           |             |
| Fluorescence-expressing vector for infection |                                                                            |             |
| studies                                      | pMV261 carrying mCherry under the control of the isp60 promoter, Kan<sup>b</sup> | 20          |

Tional mutants harboring a rough morphology were screened on agar plates. The Tn insertion site was determined as described previously (13, 21).

Gene Cloning Strategies—Standard molecular biology protocols were used for all cloning procedures. All PCR products were obtained using thermostable AccuPrime<sup>TM</sup> Pfx DNA polymerase (Invitrogen). They were initially cloned into a pJE1.2/blunt vector (Thermo Scientific) followed by sequencing and digestion with the appropriate restriction enzymes. They were then cloned into the final vectors. To facilitate subcloning, most restriction enzyme recognition sites were incorporated into the primer sequences (Table 2), although in one case (2349GR1PstI-nat primer), a natural restriction site was used.

Construction of MMAR_2349, MMAR_2331, and MMAR_2321 Gene Replacement Vectors—All PCR primers utilized in this study are listed in Table 2. To create an unmarked deletion of the wbbL2, MMAR_2331, and MMAR_2321 genes, three suicidal recombination delivery vectors based on p2NIL were used (22). In each case, the recombination vector carried the region upstream of the gene together with its 5’-end (the GR1-GR2 PCR fragment; 1434 bp for MMAR_2349; 1034 bp for MMAR_2331, and 996 bp for MMAR_2321) cloned next to the 3’-end of the gene and its downstream region (the GR3-GR4 PCR fragment; 907 bp for MMAR_2349, 1277 bp for MMAR_2331, and 1666 bp for MMAR_2321). The GR1-GR2 and GR3-GR4 PCR fragments of each gene were ligated into p2NIL vector so that the resulting plasmid copies of each gene were devoid of an internal sequence (362 bp for MMAR_2349, 415 bp for MMAR_2331, and 302 bp for MMAR_2321). Finally, the Pacl screening cassette from pGOAL17 (22) was inserted into the prepared constructs, yielding the suicide delivery vectors pGR2349, pGR2331, and pGR2321 (Table 1).

Disruption of MMAR_2349, MMAR_2331, and MMAR_2321 by Homologous Recombination—The two-step recombination protocol was used to disrupt the genes of interest at their native loci (22). The plasmid DNA of suicide delivery vectors pGR2349, pGR2331, and pGR2321 was UV-treated (100 mJ), and each of them was electroporated into competent M. marinum strain M where it integrated into the chromosome by homologous recombination. Resulting single crossover (SCO) recombinant mutant colonies were blue, Kan<sup>b</sup>, and sensitive to sucrose (2%). A single MMAR_2349, MMAR_2331, and MMAR_2321 SCO colony was then picked, resuspended in fresh 7H9 medium with OADC, poured onto solid 7H10 medium with OADC without any selective markers, and incubated at 30 °C for 5 days to allow the second crossover to occur. Subsequently, serial dilutions were plated onto medium containing sucrose and X-Gal to select for double crossovers (DCOs). Potential double crossover colonies (white and sucrose-resistant) carrying either wild-type (WT-DCO) or the mutated (ΔMMAR_2349, ΔMMAR_2331, or ΔMMAR_2321) gene (mut-DCO) were screened for kanamycin sensitivity and confirmed by PCR and Southern blotting. The PCR analysis used...
to distinguish among SCO, WT-DCO, and mut-DCO strains was performed on chromosomal template DNA using primers 2349-dco-con-sen and 2349-dco-con-rev for **MMAR** \_2349\_mutants, 2331-dco-con-sen and 2331-dco-con-rev for **MMAR** \_2331\_mutants, or 2321PhspBamHI-sen and 2321PhspBamHI-rev for **MMAR** \_2321\_mutants. The probe for Southern blot hybridization was generated by PCR using the same primers with pGR2349 (\(\Delta**MMAR**\_2349\)), pGR2331 (\(\Delta**MMAR**\_2331\)), or pGR2321 (\(\Delta**MMAR**\_2321\)) as the template. Hybridization was performed on Sall-digested (\(\Delta**MMAR**\_2349\)), PstI-digested (\(\Delta**MMAR**\_2331\)), or BamHI/BgIII-digested (\(\Delta**MMAR**\_2321\)) chromosomal DNA. Probe labeling, hybridization, and signal detection were performed using the AlkPhos Direct labeling and detection system (GE Healthcare) according to the manufacturer’s instructions.

**Construction of Complementation Plasmids**—For complementation of the **MMAR** \_2349\_DCO mutant, we constructed the pPown2349-1kb (Table 1). A PCR fragment (1782 bp) carrying whole **MMAR** \_2349\_gene and a 1014-bp upstream sequence were amplified using primers 2349P1kbKpnI-sen and 2349PownXbal-rev and cloned into the KpnI/XbaI site of the pMV261. The fragment carrying **MMAR** \_2349\_gene together with an additional 603 bp of upstream sequence and cloned it into pMV306Hyg vector (Table 1). **Drug Susceptibility Testing**—Minimal inhibitory concentrations of several antimycobacterial agents were determined against *M. marinum* strains as described previously (20). Briefly, 10-fold serial dilutions of actively growing cultures were spotted on Middlebrook 7H10 agar plates supplemented with OADC and increasing drug concentrations and incubated around 12 days at 30 °C. The minimal inhibitory concentration was defined as the minimum concentration required to inhibiting 99% of mycobacterial growth.

**Polar Lipid Extraction and Analysis**—Bacteria were grown at 30 °C on Sauton’s agar plates containing 10% OADC enrichment and recovered by scraping off the bacterial lawn. Polar lipids were extracted from the bacterial pellets according to established procedures (10) and separated by two-dimensional thin layer chromatography (TLC) on \(10 \times 10\text{-cm}\_2\) plates of aluminum-backed silica Gel 60 (Merck) using chloroform/methanol/water (60:30:6, v/v/v) in the first direction and chloroform/acetate acid/methanol/water (40:25:3:6, v/v/v/v) in the second direction. Glycolipids were visualized by spraying plates with orcinol/sulfuric acid reagent (0.2% (w/v) orcinol in \(\text{H}_2\text{SO}_4\)/water (1:4, v/v)) followed by charring. For subsequent structural analyses, individual LOS species were purified by preparative two-dimensional TLC on \(20 \times 20\text{-cm}\_2\) plates of glass-backed silica Gel 60 (Merck). The lipids were visualized with iodine labeling and recovered by scraping off the spots of interest prior to resuspension in chloroform/methanol/water (1:2:0.8, v/v/v). Solutions were filtered and dried under vacuum prior to structural determination.

**Structural Analyses**—Permethylated LOSs was performed according to the procedure described by Ciucanu and Kerek (23). Briefly, glycolipids were incubated for 2 h in the presence of 200 mg/ml\(^{-1}\) \(\text{NaOH}\) in dry DMSO (300 \(\mu\)l) and iodomethane (200 \(\mu\)l). The methylated LOSs were extracted in

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### TABLE 2

Primers used for PCR amplification

Underlined regions indicate the restriction sites used during the cloning steps.

| Primer Sequence (\(5' \rightarrow 3'\)) | Description |
|-------------------------------------|-------------|
| **Construction of the gene replacement vectors** |
| 2349GR1PstI-sen CGAAGCTTACAACTAGCCCTTGTTCGTCCT | GR1-GR2 PCR fragment of **MMAR** \_2349\_ |
| 2349GR2BamHI-rev GGGATCCGTAGTTAGCTGTCACT | GR3-GR4 PCR fragment of **MMAR** \_2349\_ |
| 2349GR3BamHI-sen CGGATCCACGAAGTGTGCGTCGTC | GR3-GR4 PCR fragment of **MMAR** \_2349\_ |
| 2349GR4PstI-nat GTCTAGAGTTGTCGTCAGATTGAGGAAGCGGCGGG | GR3-GR4 PCR fragment of **MMAR** \_2349\_ |
| 2331GR1HindIII-sen CAAGCTTCCGTTGCCTGCCAATCGTGATC | **MMAR** \_2331\_ PCR fragment, Southern blot probe synthesis |
| 2331GR2BamHI-rev GGGGTACCACGCACAAACGGCCCAACAG | **MMAR** \_2331\_ PCR fragment, Southern blot probe synthesis |
| 2331GR3HindIII-sen CAAGCTTCCGTTGCCTGCCAATCGTGATC | **MMAR** \_2331\_ PCR fragment, Southern blot probe synthesis |
| 2331GR4KpnI-rev GTCTAGAGTTGTCGTCAGATTGAGGAAGCGGCGGG | **MMAR** \_2331\_ PCR fragment, Southern blot probe synthesis |
| 2349GR3BamHI-rev GGGGTACCACGCACAAACGGCCCAACAG | **MMAR** \_2331\_ PCR fragment, Southern blot probe synthesis |
| 2349GR4KpnI-nat GTCTAGAGTTGTCGTCAGATTGAGGAAGCGGCGGG | **MMAR** \_2331\_ PCR fragment, Southern blot probe synthesis |

### SCO/DCO mutant screening

**Southern blot probe synthesis**

| Primer Sequence (\(5' \rightarrow 3'\)) | Description |
|-------------------------------------|-------------|
| 2349-dco-con-sen CGAGATCCATCTCCCTTATCAGATTATTTTCCCG | \(\Delta**MMAR**\_2349\_SCO/DCO mutant screening, Southern blot probe synthesis |
| 2349-dco-con-rev CTTACAGTACTCCCTTATCAGATTATTTTCCCG | \(\Delta**MMAR**\_2349\_SCO/DCO mutant screening, Southern blot probe synthesis |
| 2331-dco-con-sen ACGGCGCATAGCTGGCATAGAC | **MMAR** \_2331\_ PCR fragment, Southern blot probe synthesis |
| 2331-dco-con-rev GGGCTTCCCATCTCCCTTATCAGATTATTTTCCCG | **MMAR** \_2331\_ PCR fragment, Southern blot probe synthesis |

**Construction of the complementation vectors**

| Primer Sequence (\(5' \rightarrow 3'\)) | Description |
|-------------------------------------|-------------|
| 2349PownXbal-rev GTCTAGAGTTGTCGATCGCTGACCT | pPown2349-1kb vector construction |
| 2349P1kbKpnI-sen GTGATACCTTTAAAGGTCCACGGGACACCAC | pPown2331 vector construction |
| 2331PstI-bamHI-rev GGGGTACCACGCACAAACGGCCCAACAG | pPhsp2321 vector construction, \(\Delta**MMAR**\_2321\_SCO/DCO mutant screening, Southern blot probe synthesis |
| 2321PhspBamHI-sen CGGATCCATCTCCCTTATCAGATTATTTTCCCG | pPhsp2321 vector construction, \(\Delta**MMAR**\_2321\_SCO/DCO mutant screening, Southern blot probe synthesis |
| 2321PhspBamHI-rev GGGGTACCACGCACAAACGGCCCAACAG | pPhsp2321 vector construction, \(\Delta**MMAR**\_2321\_SCO/DCO mutant screening, Southern blot probe synthesis |

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chloroform and washed seven times with water. After evaporation of the reagents, the samples were dissolved in methanol prior to mass spectrometry analysis. The molecular masses of native and permethylated compounds were measured by MALDI-TOF on a Voyager Elite reflectron mass spectrometer (PerSeptive Biosystems, Framingham, MA) equipped with a 337-nm UV laser. Samples were prepared by mixing in a tube 5 μl of diluted native or permethylated LOS solutions in methanol and 5 μl of 2,5-dihydroxybenzoic acid matrix solution (10 mg/ml) dissolved in methanol/water (1:1, v/v).

**Macrophage Culture and Phagocytosis Assays**—The macrophage-like cell line J774 was maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal calf serum (FCS) at 37 °C under atmosphere containing 5% CO2. Cells were allowed to adhere in a 24-well plate for 24 h at 37 °C at a density of 5 × 104 cells/well in 0.5 ml. For infections, mycobacteria were grown at 30 °C on Sauton’s agar plates containing 10% OADC enrichment for 5–7 days. Bacterial lawns were scraped off the plates, and bacteria were resuspended in 1 ml of PBS. Bacterial clumps were disrupted by 10 successive passages through a 26-gauge needle. The remaining aggregates were then eliminated with a short spin for 1 min at 1200 rpm in a microcentrifuge. Homogeneity of the bacterial preparations and lack of eventual remaining bacterial clumps were assessed under the microscope following acid-fast staining. Bacterial suspensions were diluted in DMEM supplemented with 10% FCS at a density of 2 × 105 cells/ml and plated onto Middlebrook 7H10 agar supplemented with 10% OADC to determine the exact number of bacteria in the inoculum. The macrophage medium was then replaced by 0.5 ml of the bacterial inoculum to reach a multiplicity of infection (m.o.i.) of 2. Cells were incubated for 3 h at 30 °C under atmosphere containing 5% CO2 to enable phagocytosis. Colonies were formed and detected based on counting cfu. After incubation for 24 h at 30 °C, the medium containing extracellular bacteria was aspirated. The amoeba monolayer was washed three times with fresh PYG medium and treated with 150 μg/ml gentamicin for 1 h to kill the remaining extracellular mycobacteria, allowing study of only internalized microorganisms. The medium was then removed and replaced with fresh PYG.

To assess the cfu, infected A. castellanii or A. polyphaga were treated with Mycopep lysis buffer (BD Biosciences), and mycobacteria were removed by centrifugation at 13,000 rpm for 10 min at room temperature. The resulting bacterial pellet was washed three times with sterile PBS, diluted, and plated on Middlebrook 7H10 agar. cfu were counted after 8–10 days at 30 °C. The percentage of infected Acanthamoeba was determined based on counting >50 amoebae in 10 different microscopic fields. Infected amoebae were stained with Ziehl-Neelsen stain according to the manufacturer’s instructions (BD TB stain kit).

**RESULTS**

**Isolation of M. marinum LOS Mutants with Altered Colony Morphology**—Initial studies dedicated to the genetics of LOS biosynthesis in mycobacteria were based on the isolation of a transposon-insertion mutant of M. marinum 1218R that exhibited altered colony morphology (24). This mutant, carrying a transposon in MMAR_2313/losA and originally thought to be defective in phosphatidylinositol mannoside synthesis, was later proven to be deficient in LOS biosynthesis (10). Similarly, inactivation of MMAR_2333 involved in addition of a caryophylllose moiety in LOSs was also reported to exhibit a rough texture (16). In addition, a recent study aimed to identify novel components involved in the secretion of PE_PGRS proteins led to the identification of several M. marinum mutants with transposon insertions in the LOS biosynthetic gene cluster (14). Most of these mutants also exhibited an altered morphotype (14). Based on these observations and with the aim to provide a large panel of LOS biosynthetic mutants to initiate structure-function relationship activities, we first screened an MmaM::TnS370 transposon library (20) on Middlebrook 7H10 agar plates to select mutants with impaired colony morphology. This led to
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FIGURE 1. Screening for LOS biosynthetic mutants in M. marinum. A, selection of rough morphotypes. Screening the Tn library on Middlebrook 7H10 led to six rough variants carrying Tn insertions in losA (Mmar_2313) (two independent mutants), Mmar_2315, Mmar_2319, Mmar_2333, and Mmar_5103. Positions of the insertions are indicated in parentheses. White arrows indicate the Tn mutants that were further analyzed in this study. An additional papA4/MMAR_Mmar_2343::Tn mutant picked up as a rough mutant and characterized previously (13) is also shown. B, LOS profiles of the M. marinum mutants exhibiting a rough morphotype on 7H10 agar. Following extraction, polar glycolipids were detected with orcinol/sulfuric acid staining and charring. Acetic acid/methanol/water (40:25:3:6, v/v/v/v) in the first direction and chloroform/methanol/water (60:30:6, v/v/v) in the second direction. Glycolipids were detected with orcinol/sulfuric acid staining and charring. PIMs, phosphatidylinositol mannosides.

with insertions in Mmar_2315, encoding a hypothetical methyltransferase, and in Mmar_5103, encoding a protein of unknown function, exhibited a LOS profile comparable with that of the parental strain (Fig. 1B) and therefore were not further investigated.

Targeted Inactivation of LOS Biosynthetic Genes—The above mentioned set of mutants comprises strains that are deficient either in the very early step of LOS biosynthesis (papA4 mutant) or in the final steps involved in the synthesis of LOS-IV (losA or Mmar_2319). However, to provide a wider panel of mutants expressing intermediate structural LOS variants, we opted for targeted gene disruption of three previously uncharacterized genes: Mmar_2349 (wbbL2), Mmar_2321, and Mmar_2331. wbbL2 was selected because it encodes a putative rhamnosyltransferase sharing homology with WbbL1, which uses dTDP-rhamnose to provide the rhamnosyl-containing linker unit responsible for the attachment of the cell wall polymer mycolyl arabinogalactan to the peptidoglycan (25). Methylrhamnose is a monosaccharide found in LOS-I to LOS-IV. Mmar_2331, encoding a hypothetical protein, was chosen because it is located in the close vicinity of Mmar_2333, reported to encode a glycosyltransferase involved in the generation of a lipid-linked caryophylllose donor (16), the caryophylllose consisting of a family of unusual 4-C-branched monosaccharides shared by LOS-II to LOS-IV (11). With respect to the biosynthesis/transfer of the very unusual aglycone moeity of LOS-IV, Mmar_2321 was proposed as a putative candidate for the transfer of the pyrrolidone cycle onto the deoxy-Gal through an amide bond (12). Unmarked deletions of wbbL2, Mmar_2331, and Mmar_2321 were performed by generating the corresponding suicidal recombination delivery vectors, and disruption was achieved by homologous recombination using a two-step recombination protocol as described under “Experimental Procedures.” PCR analyses on chromosomal template DNA were done to distinguish single crossover, wild-type double crossover, and mutant double crossover strains (Fig. 2A). In addition, Southern blot hybridization was performed to confirm the deletion in mutated M. marinum strains (Fig. 2B).

Both the wbbL2 and Mmar_2331 mutants exhibited altered colony morphology with a rough appearance, suggesting potential alterations in the cell wall and presumably in the LOS profile (Fig. 2C). For complementation of the wbbL2 mutant, pPown2349-1kb was constructed (Table 1) in which a DNA segment comprising the whole wbbL2 gene along with an additional 1014-bp fragment sequence upstream of wbbL2 was cloned in the pMV306Hyg integrating vector. For complementation of the Mmar_2331 mutant, pPown2331 was achieved by cloning Mmar_2331 together with its 603-bp upstream sequence cloned into pMV306Hyg (Table 1). In both cases, the colony morphology of the mutants was restored to that of the parental type upon introduction of the corresponding complementing plasmid, indicating that the phenotype observed was due to the loss of WbbL2 and Mmar_2331 activity (Fig. 2C). In contrast, the Mmar_2321 exhibited a colony morphology similar to that of the wild-type strain (data not shown).

LOS Content in the wbbL2, Mmar_2331, and Mmar_2321 Mutants—To determine whether disruptions of wbbL2, Mmar_2331, and Mmar_2321 play a role in LOS biosynthe-
sis, the polar lipid profiles of the corresponding knock-out strains were examined by two-dimensional TLC (Fig. 3). The wbbL2 mutant clearly showed a distinctive LOS pattern with the absence of most major LOS species and a concomitant accumulation of two intermediates, designated spot 1 and spot 2 (Fig. 3). Both spots were purified by preparative TLC and analyzed by MALDI-MS in native and permethylated forms. The MALDI-MS spectrum of the native spot 1 exhibited two distinct 14-mass unit-incremented peak clusters with maximum intensities at \( m/z \) 1221 and 1459, establishing the presence of a di- and tri-acetylated Hex\(_4\) respectively, substituted by complex mixtures of lipids (Fig. 4A). The carbohydrate moiety attribution was confirmed by MS analysis of permethylated glycolipids that showed a sharp signal at \( m/z \) 885 corresponding to a Hex\(_4\) oligosaccharide. In addition, following the per-O-methylation procedure (13), the alkali-labile fatty acyl substituents are incompletely replaced by methyl groups, a fraction of which is retained on the trehalose moiety as demonstrated by clusters around signal at \( m/z \) 1137 and 1389 corresponding to partially deacylated Hex\(_4\) (Fig. 4A). The MS profile is identical to that of LOS-I as reported previously (13) except for a loss of 174 mass units corresponding to a deHex residue. Altogether, these data strongly suggest that spot 1 contains a new biosynthetic intermediate Glc\(_4\) (Ac\(_4\)) designated LOS-0 (Ac\(_4\)) that corresponds to the immediate precursor of LOS-I. The observation of native LOS-0 (Ac\(_4\)) along with LOS-0 (Ac\(_4\)) was
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Firming the involvement of WbbL2 with Rha metabolism. Accumulation of LOS-0 appears to trigger the synthesis of a so far undescribed LOS-0* most likely through the addition of a further glucose residue onto the tetraglucosyl backbone of LOS-0. However, the position of the extra glucose residue could not be fully determined due to low quantities.

The two-dimensional TLC profile of the MMAR_2331 mutant revealed the absence of LOS-II, LOS-III, and LOS-IV paralleled by the accumulation of a glycolipid absent in the parental strain and migrating to an intermediate position between LOS-I and LOS-II (Fig. 3, designated spot 3). However, this TLC profile was highly reminiscent of that reported in an MMAR_2332 mutant (also known as ilvB1_3) (14, 15) or in an MMAR_2333 mutant (16) characterized by the accumulation of a LOS-II* intermediate. To further demonstrate the nature of spot 3, the purified product was subjected to MALDI-MS (Fig. 4C). Analysis of the permethylated spot 3 revealed a major oligosaccharide signal at m/z 1219 attributed to Hex₅deHex₅Pent₁ in agreement with the LOS-II* monosaccharide composition Glc₃Me-Rha₃Xyl₁. Mono-, di-, and triacylated forms of this oligosaccharide substituted by various combinations of lipids were further observed as clusters around signals at m/z 1471, 1423, and 1976. Similarly, LOS-II* (Ac₃) and its partially deacylated LOS-II* (Ac₂) were observed in their native forms as clusters around major signals at m/z 1513 and 1751 (Fig. 4C). Overall, these results clearly establish the identity of spot 3 as LOS-II*, the immediate biosynthetic precursor of LOS-II, which contains δ-Xylp attached to the Glc₃Me-Rha₃ core but lacks the caryophyllose found in LOS-II. These data indicate that the intermediate we isolated from the MMAR_2331 knock-out strain was identical to that produced in the MMAR_2332 (15) and MMAR_2333 (16) mutants, thus connecting MMAR_2331 to either the synthesis of caryophyllose or its transfer on LOS-II* to generate LOS-II.

The MMAR_2321 mutant was characterized by the absence of LOS-IV and the concomitant accumulation of an intermediate glycolipid, designated spot 4 (Fig. 3). MALDI MS analysis (Fig. 4D) of permethylated spot 4 revealed a major signal at m/z 1916.0 tentatively attributed to Car₃Hex₅deHex₅Pent₁ in agreement with the oligosaccharide moiety OS-III identified in LOS-III (Fig. 4D) (11). The presence of two caryophyllose residues at the terminal non-reducing end was confirmed in MS/MS analysis of OS-III by observing two consecutive losses of 278 mass units at m/z 1791.7 and 1513.5 from signal at m/z 2070 (data not shown). Along with OS-III, three clusters of signals around m/z 2168, 2420, and 2644 were identified as partially decylated LOS-III (Ac₃) and LOS-III (Ac₂) and as fully acetylated LOS-III (Ac₄), respectively. The presence of LOS-III (Ac₄) and decacylated LOS-III (Ac₃) was confirmed in the analysis of native spot 4 by the observation of two signals clusters around m/z 2308 (LOS-III (Ac₃)) and 2070 (LOS-III (Ac₄)). The acylation status of LOS-III (Ac₃) and LOS-III (Ac₄) are similar to those of LOS-II* and LOS-0. In addition, two additional clusters of signals were observed around m/z 2196 and 2000. The MS/MS fragmentation pattern of signal at m/z 2196 indicated that this compound is a decylated LOS-III substituted by a set of fatty acids longer by an average of nine carbons than those of LOS-III (Ac₄). These compounds are referred to

![FIGURE 3. LOS content of the various M. marinum mutants.](image)

Overall, these results indicate that this strain corresponds to a high order LOS mutant and that functional inactivation of wbbL2 occurred in the very early biosynthetic steps. The LOS-0 precursor lacks the terminal methyl-Rha residue of LOS-I, contradictingly attributed to a partial decylation of native LOS-0 (Ac₃) that typically occurs during MALDI analysis. In contrast to spot 1, analysis of spot 2 showed a slightly more complicated pattern. In native form, the main signal at m/z 1221 was assigned to LOS-0 (Ac₃), but no LOS-0 (Ac₀) could be observed. Accordingly, the MALDI spectrum of the permethylated sample showed signals consistent with the presence of fully deacetylated oligosaccharide OS-0 as well as partially deacetylated LOS-0 (Ac₂) and LOS-0 (Ac₁) (Fig. 4B). Along with these signals attributed to LOS-0 (Ac₃), another set of signals was found in native forms around m/z 1383 and 1621 that were attributed to Hex₅ (Ac₂) and Hex₅ (Ac₃). As described previously, observation of native Hex₅ (Ac₃) can be interpreted as a partial decylation of native Hex₅ (Ac₄) during MALDI-MS analysis. This assignment was confirmed following permethylation, which led to signals at m/z 1089 and 1341 corresponding to Hex₅ oligosaccharide and Hex₅ (Ac₄) glycolipid, respectively. This established spot 2 as being composed of LOS-0 (Ac₃) and of a so far undescribed Hex₅ (Ac₃) glycolipid designated LOS-0*.

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Importantly, complementation of all the mutants restored a wild-type LOS profile (Fig. 3), indicating that alteration of the LOS pattern could be attributed to the inactivation of the targeted genes and not to an eventual polar effect on the downstream genes.

**Altered LOS Production Facilitates Phagocytosis by Macrophages**—Fig. 5A relates the structures of the major LOS subspecies identified and described in the present study. The availability of a representative set of *M. marinum* mutants with various incomplete LOS variants allowed us to examine their potential biological role in the early interaction with professional phagocytes as would be expected for surface-attached molecules. To this aim, wild-type *M. marinum* strain M (expressing the whole range of LOSs), *papA4* mutant (lacking the entire LOS profile), and other mutants expressing either only highly truncated LOS subspecies (*wbbL2*) or intermediate variants (*MMAR_2331*) or lacking only LOS-IV (*losA*, *MMAR_2319*, and *MMAR_2321*) were used to infect J774 macrophages to examine and compare their ability to enter into the cells. In Sauton’s broth medium, all mutants exhibited growth rates similar to that of the parental strain (data not shown). To examine the entry of bacteria into macrophages, infection experiments were performed at an m.o.i. of 2. Immediately following infection (3 h at 30 °C), cells were treated with gentamycin to kill extracellular mycobacteria and washed three times with PBS. The cfu recovered from macrophages were enumerated by plating on 7H10 agar. Unexpectedly, our results showed an inverse correlation between the size of the LOS and an increased phagocytosis rate, culminating with a reproducibly 7-fold increased rate with the LOS-deficient *papA4* mutant (Fig. 5B). Intermediate phagocytosis rates were observed with the *wbbL2* mutant lacking most LOS subspecies and the *MMAR_2331* mutant characterized by the accumulation of LOS-II* and the defect in more polar LOS subspecies. Conversely, the two LOS-IV-deficient strains (*MMAR_2319* and *losA*) failed to demonstrate increased phagocytosis rates. Importantly, the increased phagocytosis rates of the *wbbL2* and *MMAR_2331* mutants were restored in the complemented mutant strains (Fig. 5C). Analyzing the kinetic of *wbbL2* phagocytosis during the first 3 h of contact with macrophages showed that the mutant entered macrophages very efficiently and rapidly compared with the wild-type strain, resulting in around 30% of *wbbL2*-phagocytized mycobacteria versus 10% of wild type-phagocytized mycobacteria after 3 h of infection (Fig. 5D). That the absence of LOS-IV and/or accumulation of LOS-III failed to affect the uptake of *M. marinum* by macrophages was further supported by infection experiments with the *MMAR_2321* mutant, which exhibited a phagocytosis rate similar to that of the wild-type strain (Fig. 5C). Based on these structure-function relationships, we propose that the different strains should be classified into three groups (Fig. 5A): (i) class I mutants (*papA4*) characterized by the presence of a very early triacetylated LOS precursor, designated LOS-0 (Ac2). A, mass spectrometry analysis of spot 1 from the *wbbL2* mutant revealed the presence of a very early triacetylated LOS precursor, designated LOS-0 (Ac2), whose carbohydrate moiety consists of four Glc residues. B, spot 2 from the *wbbL2* mutant consists of a mixture of LOS-0 (Ac2) and a new pentaglcosylated glycolipid designated LOS-0*. C, analysis of spot 3 from the *MMAR_2331* mutant revealed the presence of a glycolipid whose m/z corresponds to LOS-II*, a LOS intermediate that has been reported to accumulate also in *MMAR_2332* and *MMAR_2333* mutants (15, 16). D, spot 4 from the *MMAR_2321* mutant was identified as LOS-III (Ac2), based on the observation of fully acetylated and partially acetylated signal clusters in permethylated and native forms. The corresponding structures identified are shown in insets. R corresponds either to acylation position or hydrogen.
absence of LOS are very efficiently phagocytosed, (ii) class II mutants (wbbL2 and MMAR2331) producing only the early and less polar LOS subspecies with less pronounced uptakes by macrophages than class I mutants, and (iii) class III mutants (losA, MMAR2319, and MMAR2321) synthesizing all LOS subspecies except the most polar LOS-IV and phagocytosed similarly to the wild-type strain.

To rule out the possibility that the differences in phagocytosis observed may result from mycobacterial aggregation, which would affect the cfu numbers, and to investigate whether the increased phagocytosis rate is associated with an increased number of infected cells, infected macrophages were directly counted under the microscope. For this purpose, several mutants were transformed with pMV261_mCherry and subsequently used to infect macrophages. The number of cells containing red fluorescent mycobacteria was then counted under a fluorescence microscope. As shown in Fig. 5E, the highest proportion of infected macrophages accounted for the papA4 mutant (40% of infected cells), and a proportion of 25% was found for the MMAR2331 and wbbL2 mutants. In contrast, comparable proportions of infected cells were observed with the losA mutant and the wild-type strain (around 10%). Taken together, these data indicate that both class I and class II mutants are more efficiently engulfed by macrophages than class III mutants.

Altered LOS Production Increases Uptake of M. marinum by Amoebae—M. marinum is a waterborne pathogen that causes tuberculosis-like infection in ectotherms, such as frogs and fish (26). Free-living amoebae, such as Acanthamoeba, are professional phagocytes (27), and non-tuberculous mycobacteria, including M. marinum, have been recovered from samples of water colonized by free-living amoebae (28). Distinct mycobacterial species have been shown to resist destruction by amoebae (27, 29–31), which are thought to serve as a reservoir for non-

FIGURE 5. Phagocytosis of LOS mutants by murine macrophages. A, schematic representation of the different subspecies of LOSs and the intermediates identified in this study. NAc-Gal, N-acylated dideoxygalactose. R1, R2, and R3 represent the acyl chains. B, phagocytosis rate of the various LOS mutants by J774 macrophages normalized to 1 for the wild-type strain. Results are expressed as means ± S.E. (error bars) from three to five independent experiments. C, phagocytosis rates of the wbbL2, MMAR2331, and MMAR2321 mutants and complemented (_C) strains. D, kinetics of phagocytosis of the wbbL2 mutant. For C and D, results are representative of two independent experiments and are expressed as means ± S.D. (error bars) of triplicates. E, percentages of mycobacteria-containing macrophages determined by counting infected cells under the microscope. Results are expressed as means ± S.E. (error bars) from three independent experiments.
tuberculous mycobacteria in the environment. Because *M. marinum* and amoeba share the same environmental niche, it is very likely that amoebae represent a natural host for *M. marinum*. A recent study demonstrated that the ESX-1 secretion system is required for *M. marinum* pathogenesis in *A. castellanii* and that *M. marinum* utilizes actin-based mobility in this host (32), indicating that these two virulence pathways used by *M. marinum* in macrophages are conserved during infection of amoebae. Therefore, we addressed whether, like in macrophages, alteration of the LOS profile of *M. marinum* influences the early interaction events with *A. polyphaga* and *A. castellanii*, which are both equally sensitive to gentamycin treatment used to eliminate extracellular mycobacteria. *A. polyphaga* were infected with class II and class III mutants. As shown in Fig. 6A (left panel), the uptake of the *wbbL2* and *MMAR_2331* mutants was significantly increased 3-fold with respect to the parental *M. marinum* strain M and was restored to wild-type levels in the corresponding complemented strains. In contrast, the phagocytosis rate of the *MMAR_2321* mutant by *A. polyphaga* was comparable with that of *M. marinum* strain M (Fig. 6A, left panel).

In an additional set of experiments, the percentage of infected amoebae was established after counting under the microscope. Whereas the percentage of infected *A. polyphaga* was increased by 2–4-fold with the *wbbL2* and the *MMAR_2331* mutants with respect to *M. marinum* strain M, no significant differences were observed with the *MMAR_2321* mutant (Fig. 6B, left panel). The percentage of infected amoebae was restored to wild-type levels in both corresponding complemented strains. Moreover, infection studies performed in another amoeba species, *A. castellanii*, resulted in very similar effects characterized by an increase in phagocytosis of mycobacteria (Fig. 6A, right panel) and infected cells (Fig. 6B, right panel) in the case of *wbbL2* and *MMAR_2331* mutants. As for *A. polyphaga*, the *MMAR_2321* mutant behaved like the parental *M. marinum* M strain in *A. castellanii* (Fig. 6A and B, right panels). Overall, these findings suggest that early interactions and phagocytosis of *Acanthamoeba* are conditioned by the LOS pattern of *M. marinum* and that, as observed in macrophages, truncated LOS variants favor these early infection processes.

**DISCUSSION**

In this study, we identified six distinct LOS mutants among which three had been previously characterized. In agreement with earlier reports (10, 14–16), all mutants except *MMAR_2321* exhibited a drier colony morphology, indicating a change in cell envelope structure. Because changes in the cell wall structure/composition are often correlated to modifications in membrane permeability and susceptibility to drugs (20, 33), we also examined the susceptibility of the *papA4* and *wbbL2* mutant strains to hydrophilic and hydrophobic antibiotics. Both mutants exhibited minimal inhibitory concentrations to isoniazid, streptomycin, rifampicin, and erythromycin that were comparable with those of the parental strain (data not shown), implying that altered LOS composition does not affect drug susceptibility.

Biochemical and structural analysis of the biosynthetic intermediates allowed us to precisely link the participation of the inactivated genes within the LOS pathway, extending our understanding of LOS biosynthesis. In particular, the absence of LOS-I to LOS-IV with the parallel increase of LOS-0, a new biosynthetic intermediate that had not been detected/described previously and corresponding to triacylated Glc\(_\text{p}-(1\rightarrow3)\)-Glc\(_\text{p}-(1\rightarrow4)\)-Glc\(_\text{p}-(1\rightarrow1)\)-Glc\(_\text{p}\) supposes that WbbL2 expresses rhamnosyltransferase activity. Although annotated as a conserved hypothetical protein, BLAST analyses revealed homology between *MMAR_2331* and members of the HpcH/HpaI aldolase/citrate lyase family, suggesting that this enzyme may be
implicated in the synthesis of the 4-C-branched substituent of caryophyllose. Accordingly, disruption of **MMAR_2331** leads to a mutant unable to produce LOS-II to LOS-IV and that accumulates a LOS-II**−** lacking the terminal caryophyllose. **MMAR_2321** presents 44% identity with a putative N-acyltransferase that catalyzes the transfer of a pyrroline derivative onto an α-4-amino-4,6-dideoxy-Galp of the polysaccharide from *Vibrio cholerae* O:5 (34). Therefore, we propose that **MMAR_2321** encodes the N-acyltransferase involved in the linkage of the pyrroline cycle on the α-4-amino-4,6-dideoxy-Galp residue of the LOS-IV (12). Two-dimensional TLC and structural analyses indicated that disruption of **MMAR_2321** led to a mutant lacking LOS-IV and demonstrated that the accumulating product was not a synthetic intermediate between LOS-III and LOS-IV but straight LOS-III. This suggests that the synthesis of a truncated 4-N-acylated-4,6-dideoxy-Galp prevents its block transfer onto LOS-III.

As a result of the generation of a wide panel of isogenic mutants, we were able to initiate structure-function relationship activities, which clearly pointed to an inverse correlation between LOS production and efficient uptake into professional phagocytic cells. This unexpected observation contrasts with a previous study in which LOS mutants demonstrated impaired cell entry efficiency (15). The discrepancies between the two studies may be related to different experimental procedures. First, **Ren et al.** (15) used an m.o.i. of 10 in their study as compared with an m.o.i. of 2 in our study. Second, mycobacterial cultures were performed using different media, which as reported previously affect the cell wall composition, including LOSs (10) and apolar glycolipids (35). In our study, all mutants were grown on Sauton's agar prior to infections as Sauton's agar has been shown to provide higher amounts of LOSs (10). Finally, the different phenotypes observed may also be dependent on the parental *M. marinum* strain used, the 1218R strain in the **Ren et al.** (15) study and the M strain in ours. Based on the assumption that LOSs could mask other cell wall determinants, it is possible that the two strains differ in the nature and composition of these factors, some of them being important for phagocytosis. Nevertheless, our study is based on a large panel of defined mutants that reproducibly shows an increase in macrophage entry efficiency for class I and class II mutants. This view is also further strengthened by very similar phenotypes observed in *A. polyphaga* and *A. castellani*. It is also noteworthy that at later stages of infection all strains exhibited comparable intracellular growth rates (data not shown), suggesting that LOSs do not play a critical role in intramacrophage survival.

Previous infection studies with *M. kansasii* indicated that rough variants devoid of LOSs induce chronic infections, whereas smooth variants producing LOSs are rapidly eliminated (18), leading to the proposal that LOSs may be considered as an avirulence determinant capable of masking other cell wall-associated (glyco)lipids playing a role in virulence, such as lipoarabinomannan and phenolic glycolipids (17). Although the structures of LOSs in *M. kansasii* and *M. marinum* differ significantly, our findings support this “unmasking” hypothesis in *M. marinum* because early interactions with phagocytic cells, which represent the preferred residing niche of *M. marinum*, are amplified in mutants devoid of LOSs or defective in higher order LOS production. The reasons for the enhanced phagocytosis rate of these LOS mutants are currently not known, but it is very likely that the lack of LOSs results in the enhanced exposure of structures at the mycobacterial surface that will be recognized by specific macrophage receptors, eventually leading to a more virulent infection. In this context, it is worth mentioning that an LOS-IV mutant (*wecE*) has been shown to increase the degree of infection and formation of early granulomas in infected zebrafish embryos (14). Thus, these *in vivo* data suggest that LOSs can act as factors that suppress in the early bacterial infections, a view that is supported by our finding that LOS-deficient strains are more avidly phagocytosed by macrophages.

A similar unmasking hypothesis has recently emerged regarding glycopeptidolipids found in *Mycobacterium smegmatis* or *Mycobacterium abscessus* and known as metabolites required for sliding motility and biofilm formation (36). These glycolipids are also suspected to play a role in virulence by inhibiting phagocytosis (37). Importantly, the loss of glycopeptidolipid in *M. abscessus* resulted in the exposure of phosphatidylymyo-inositol (38) and the increased expression and surface localization of lipoproteins (39) responsible for a strong proinflammatory response. This supposes that the outermost portion of the *M. abscessus* cell wall comprising glycopeptidolipids is masking underlying cell wall (glyco)lipids involved in stimulating the innate immune response, thereby affecting subsequent colonization.

Recent investigations demonstrated that LOS biosynthesis defects result in a tighter surface attachment of capsular proteins, including PE_PGRS proteins and EspE (14). Thus, whether the observed increased capacity of the LOS mutants of being phagocytosed is actually caused by the lack of higher order LOS production, the accumulation of lower order LOS structures, or the altered release of surface proteins, such as PE_PGRS, is not clear at this point. Interestingly, using an *in vitro* system, **Stokes et al.** (40) demonstrated that by removing the capsular material of *M. tuberculosis* there was a significant increase in phagocytosis by macrophages compared with the uptake of the wild-type unperturbed bacterium. Thus, by analogy with *M. tuberculosis*, it seems possible that removal of LOSs may perturb the outermost capsular layer of *M. marinum*, resulting in increased uptake by macrophages.

In *M. tuberculosis*, deficiency in diacyltrehalose and pentacetyltrehalose (PAT) affects the surface global composition of the mycobacterial cell envelope, improving the efficiency with which the bacilli bind to and enter phagocytic host cells, implying that PAT production affects early interaction between the bacilli and macrophages (41) similarly to LOSs in *M. marinum*. Because *M. marinum* does not produce diacyltrehalose/PAT and because of the structural relatedness between LOSs and diacyltrehalose/PAT (all are trehalose-based glycolipids), it can be inferred that LOS fulfills a function similar to that of diacyltrehalose/PAT, although the paradigm of the avirulence-promoting role of LOSs awaits further confirmation. LOSs are absent from *M. tuberculosis*, which may be explained by the fact that *M. tuberculosis* possesses fewer genes within the LOS cluster (15). However, LOSs are found in *M. canettii*, also known as
the smooth tubercle bacilli (9), a species restricted to East Africa. With the recent completion of the *M. canetti* genome (42), it now becomes possible to compare its LOS loci with *M. marinum* and to generate isogenic mutants in this species to investigate the role of LOSs in phagocytosis and pathogenicity of *M. canetti*. Whether the presence of LOSs may be related to the decreased persistence and virulence of *M. canetti* compared with *M. tuberculosis* in mice remains an attractive hypothesis that needs to be addressed. Moreover, the recent observation that *M. canetti* can be ingested by amoebal trophozoites (31) implies that *Acanthamoeba* infection also represents an alternative means to dissect and address the role of LOSs in the uptake and virulence of this human pathogen.

As mentioned above, the profile and amount of the different LOS subspecies are dependent on the medium (10), indicating that *M. marinum* has the ability to modulate its LOS content. Therefore, one may speculate that the LOS profile varies under the different growth and environmental conditions encountered and/or during intracellular replication/survival of mycobacteria. That control/regulatory mechanisms of LOS production occur in the *M. marinum*-infected host would directly influence the early interactions with phagocytes and perhaps condition the outcome of the infection. In this context, a *whiB4* (MMAR_5170) mutant showed a highly diminished LOS production, and WhiB4 was proposed to regulate the expression of various LOS biosynthetic genes (14). However, whether WhiB4 plays a role in regulating the LOS pathway during infection remains to be investigated.

There is now an increasing body of evidence suggesting that protozoa are a reservoir for environmental mycobacteria in the environment. In a recent study, Kennedy *et al.* (32) demonstrated that the requirement of both the ESX-1 secretion system and the actin-based motility of *M. marinum* in *A. castellani* is similar to the situation encountered in macrophages (43). Our study shows that the impact of the LOS pattern of *M. marinum* was equivalent in macrophages and in *Acanthamoeba*. These findings not only contribute to the basic molecular understanding of the interactions between *Acanthamoeba* and *M. marinum* but also further support the view that the virulence pathways used by *M. marinum* in macrophage are conserved during *Acanthamoeba* infection.

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