Mycolates of *Mycobacterium tuberculosis* modulate the flow of cholesterol for bacillary proliferation in murine macrophages

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Abstract The differentiation of macrophages into lipidfilled foam cells is a hallmark of the lung granuloma that forms in patients with active tuberculosis (TB). Mycolic acids (MAs), the abundant lipid virulence factors in the cell wall of *Mycobacterium tuberculosis* **(Mtb), can induce this foam phenotype possibly as a way to perturb host cell lipid homeostasis to support the infection. It is not exactly clear how MAs allow differentiation of foam cells during Mtb infection. Here we investigated how chemically synthetic MAs, each with a defined stereochemistry similar to natural Mtbassociated mycolates, influence cell foamy phenotype and mycobacterial proliferation in murine host macrophages. Using light and laser-scanning-confocal microscopy, we assessed the influence of MA structure first on the induction of granuloma cell types, second on intracellular cholesterol accumulation, and finally on mycobacterial growth. While methoxy-MAs (mMAs) effected multi-vacuolar giant cell formation, keto-MAs (kMAs) induced abundant intracellular lipid droplets that were packed with esterified cholesterol. Macrophages from mice treated with kMA were permissive to mycobacterial growth, whereas cells from mMA treatment were not. This suggests a separate yet key involvement of oxygenated MAs in manipulating host cell lipid homeostasis to establish the state of TB.**—Vermeulen, I., M. Baird, J. Al-Dulayymi, M. Smet, J. Verschoor, and J. Grooten. **Mycolates of** *Mycobacterium tuberculosis* **modulate the flow of cholesterol for bacillary proliferation in murine macrophages.** *J. Lipid Res.* **2017.** 58: **709–718.**

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Mycobacterium tuberculosis (Mtb), the etiological agent of tuberculosis (TB), infects approximately two to three

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billion people globally (1). Though only a minority of infected individuals (<15%) will develop pulmonary disease during their lifetime, most remain prolonged asymptomatic carriers that may develop active disease later on (1). In this instance, bacilli are not entirely cleared, but remain in a state of dormancy inside the host. Throughout this period, Mtb resides inside lung granulomas, the main histopathology of TB (2). The granuloma milieu is characterized by a multifaceted host immune response of containment and destruction, yet Mtb bacilli are able to counteract and evade host defenses (3–5). During active TB, granulomas are characterized by large cell aggregates of lymphocytes, neutrophils, dendritic cells, and peripheral fibroblasts (6). The key effector cells of granuloma formation are macrophages, which can differentiate into vacuolar multinucleated giant cells (MGCs) or lipid-laden foam cells (7). Tuberculous bacilli may reside in foam cells or escape into the cell-free caseous center of the granuloma (2, 7). The biochemical signaling pathways involved in foamy phenotype regulation remain understudied. Growing evidence indicates that an advanced metabolic network stands at the center of the unique adaptation of Mtb in its host macrophage (8–10), regulating the manifestation of latent, chronic, or acute TB.

The Mtb cell envelope comprises intricate layers of peptidoglycan, arabinogalactan, glycolipids, mycolic acids (MAs; α -alkyl, β -hydroxy fatty acids) and lipoproteins, with MAs being the dominant constituent (11, 12). The structural and biochemical properties of MAs have been reviewed

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Abbreviations: αΜΑ, α-mycolic acid; BCG, Mycobacterium bovis bacille Calmette-Guérin; BCG-dsRed, *Mycobacterium bovis* bacille Calmette-Guérin expressing the dsRed fluorescent protein; GLM, generalized linear model; kMA, keto-mycolic acid; LD, lipid droplet; Lipo, liposome carrier without synthetic mycolic acid; LXR, liver X receptor; MA, mycolic acid; MGC, multinucleated giant cell; mMA, methoxy-mycolic acid; MOI, multiplicity of infection; Mtb, *Mycobacterium tuberculosis*; PEC, peritoneal exudate cell; TB, tuberculosis; V+, vacuole-positive. ¹

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recently (13). In short, these wax-like hydrophobic lipids comprise a mycolic motif with a long nonfunctionalized alkyl chain and a meromycolate chain with up to two functional groups that can be either oxygenated (distal group) or unoxygenated (distal or proximal group) (13). Three main Mtb MA classes exist: the most abundant unoxygenated α -MA (α MA), the less abundant oxygenated methoxy-MA (mMA), and the least abundant keto-MA (kMA; **Fig. 1**) (14, 15). Orientation of the proximal cyclopropane differs. MA essentially exists in *cis* configuration (though a small amount of *trans* may be present), whereas mMA and kMA contain either *cis*- or *trans*-cyclopropanation with an adjoining methyl branch (11). While the level of α MA is fixed at \sim 53% and oxygenated MA at \sim 47%, the ratio between the two oxygenated MAs varies with methoxy at 32–40% and keto at 7–15% (16), depending on the growth stage of the bacilli (17–19). Mtb envelope-derived lipids potently influence host immunity (20–22), while host cell lipidomes are exploited by intracellular pathogens like Mtb to gain entry and replicate (23). The work by Cole et al. (24) on deciphering the Mtb genome identified two distinct sets of complex enzymatic machinery for successive biosynthesis of fatty acids, meromycolates and long carbon chain MAs. A MA-rich cell wall that can be altered depending on physiological requirements is essential for Mtb virulence (17, 18). We previously showed that in vivo MA treatment induced peritoneal and alveolar macrophages of the foam phenotype in mice (25, 26), similar to that in macrophages from the TB granuloma (27). At the onset of TB infection,

GK324, kMA (17,16,17,23)

Fig. 1. MA structures. The biochemical structures of the synthetic MAs are given for examples of the α MA (JR1080), mMA (JR1046), and kMA (GK324) classes containing *cis*-cyclopropanation. Numbers in brackets represent carbon chain lengths and wiggly line indicates a mixture of stereoisomers at that position.

foam cells form that are characteristically enlarged and filled with multiple lipid droplets (LDs) and vacuoles (28, 29). Cholesterol is abundantly distributed across cell membranes and forms a major constituent of LDs (30, 31). It also plays a unique role in Mtb virulence and pathogenesis. Mtb preferentially catabolizes cholesterol as a nutrient source, while its acquisition, through a unique Mtb import system, is necessary to establish and maintain persistent infection (32, 33). In macrophages, the genetic regulator of cellular cholesterol homeostasis is the liver X receptor (LXR) (34). LXRs are transcription factors that act as cholesterol sensors and that maintain the balance of cholesterol uptake and export through regulation of expression of cholesterol-associated target genes (35).

To better understand the mechanisms responsible for the establishment of active TB-related granulomas, we investigated how in vivo treatment of mice with chemically synthetic MAs, each with a defined stereochemistry representing the separate major classes of Mtb mycolates, influences cell differentiation and support of mycobacteria in peritoneal macrophages. We assessed the influence of MA structure first on the induction of foamy macrophages and MGCs identified by light and laser-scanningconfocal microscopy, second on cholesterol accumulation, and finally on intracellular mycobacterial growth. Our results show that Mtb mycolates differentially steer host macrophages to either an enlarged vacuolar or a lipidladen foamy phenotype. We report that it is kMA that induces mainly cholesterol ester accumulation and intracellular LDs to sustain facilitation of mycobacterial *Mycobacterium bovis* bacille Calmette-Guérin (BCG) proliferation. mMA was found to induce vacuolation with no change in cholesterol ester levels and no improvement in the ability to sustain and facilitate mycobacterial growth. MA treatment had a negligible effect on these parameters. To determine how foam cells brought about by nontuberculous means compare to foam cells induced by the different mycolates, we also investigated cholesterol accumulation and mycobacterial growth of peritoneal macrophages from LXR-deficient mice. In macrophages with a deficiency in LXR activity that is characterized by perturbed cholesterol transport or export, we recorded foam cells with abundant cholesterol ester-containing LDs that showed elevated BCG replication.

MATERIALS AND METHODS

MAs

Natural MA mixture was isolated and single synthetic MAs synthesized as previously described (16, 36–38). The single MAs used for in vivo murine treatment in this study all contained *cis*-cyclopropanation, referring to the orientation of the proximal cyclopropane. MA treatments comprised *cis*-MA, *cis*-mMA, *cis*-kMA consisting of a mixture of both epimers of the distal α -methylketone group with S- and R-stereochemistry, and a natural isolated mixture of all three MA classes (each as a complex mixture of homologs) similar to the natural composition of MAs in the Mtb cell wall (MA mix) (Fig. 1), i.e., $\sim 53\%$ aMA, $\sim 38\%$ mMA, and $\sim9\%$ kMA.

Animals

The mice used were specific pathogen-free C57BL/6 WT females, aged 8–12 weeks (Janvier Labs, France). C57BL/6Bom WT and LXR $\alpha^{-/-} \beta^{-/-}$ mice (39) were bred in the animal facility of Ghent University. Animals were housed individually in a temperature- and light-controlled facility and received mixed ration feed and water ad libitum. Experiments were preapproved by the Ghent University Ethical Committee for Animal Experimentation in accordance with current European laws regarding the welfare and humane use of animals.

Experimental design

Mice were treated intraperitoneally with various control or MA solutions 2 days prior to harvesting of peritoneal exudate cells (PECs). Macrophages from PECs were cultured for 3 days and live cells were fluorescently labeled for foam cell markers and examined by laser-scanning-confocal microscopy on each day (0, 24, and 48 h time points). The mycobacterial model was similar to the foam cell model except that, following overnight adherence, cells were infected for 6 h with BCG expressing the dsRed fluorescent protein (BCG-dsRed; a gift from Prof. Ben Appelmelk from the Department of Medical Microbiology and Infection Control at the Vrije Universiteit University Medical Centre in Amsterdam via the Unit of Medical Biotechnology at the Inflammation Research Center in Ghent). After three washes with endotoxin-free PBS (Lonza), the cells were stained with fluorescent markers for confocal microscopy (0 h) to assess macrophage morphotype or left for up to 5 days to measure mycobacterial growth (48 and 96 h). The LXR model was similar to the mycobacterial model, except that mice were not treated with MAs prior to harvesting PECs.

Mycobacterial culture and infection

BCG-dsRed has been described previously (40). The BCG strain, Copenhagen (Danish 1331), was used here (41), which has no mMA, but similar quantities each of α MA and kMA (42). Bacterial cultures were grown in Middlebrook 7H9 broth (Difco) supplemented with 0.2% glycerol, 0.05% Tween-80, and 10% Middlebrook AODC enrichment containing oleic acid, albumin, dextrose, and catalase (Becton Dickinson). BCG expanded to an $OD_{600 \text{ nm}}$ of 0.8–1.0 was used to infect cells ex vivo at a multiplicity of infection (MOI) of one bacterium per cell for 6 h. Following the 6 h infection, cells were washed three times with warm endotoxin-free PBS and cultured at 37° C and 5% CO₂.

Injectable solutions and macrophage isolation

Liposomes were used as the carrier to deliver the highly hydrophobic MA compounds to target cells. As a first step, $L-\alpha$ phosphatidylcholine (Sigma®) powder was dissolved in chloroform at 100 mg/ml (10% w/v). Phosphatidylcholine and MA dissolved in chloroform were vortexed and heated before undergoing dehydration on a heat block (90°C) and the dried lipids recovered in endotoxin-free PBS. Solutions underwent a series of vortex and sonication steps at 65°C until reaching a homogenous milky consistency. Mice were immediately treated in vivo by intraperitoneal injection (25 μ g MA/100 μ l/mouse). A liposome control [liposome carrier without synthetic MA (Lipo)] was formulated as described for the MA solutions, but without the addition of any synthetic MA. Two days after in vivo treatment, mice were euthanized via cervical dislocation and PECs harvested by peritoneal lavage. Mouse abdomens were decontaminated with 70% ethanol and 10 ml ice-cold endotoxin-free PBS injected intraperitoneally. Following a short abdominal massage, PECs were removed into sterile 15 ml tubes and kept on ice until further processing by centrifugation (180 *g*, 4°C, 10 min) and red blood cell lysis (ACK lysing buffer, 50% v/v; Lonza). PECs were seeded in

250 µl culture medium (5 \times 10⁵ cells) in µ-Slide 8-well microscopy plates (Ibidi®). Culture medium consisted of RPMI 1640 (Gibco®) supplemented with LPS-free and heat-inactivated FCS (10%), sodium pyruvate (2 mM), nonessential amino acids (1%), penicillin/streptomycin antibiotics (0.2%), and β -mercaptoethanol (0.1%). Cultures were enriched for macrophages by overnight adherence. All cells were cultured at 37°C and 5% CO₉.

Light and laser-scanning-confocal microscopy

An aliquot of cell suspension equal to 5×10^4 to 1×10^5 cells was taken for cytospin analysis. The cytospin filter was primed with 100 μ l PBS (20 *g*, 1 min) before addition of 200 μ l cell suspension (20 *g*, 5 min). After an overnight drying step, cells were fixed in methanol for 30 min at -20° C and dried for 2 h. Cells were stained with undiluted May-Grünwald for 5 min (granular stain), washed in PBS, and stained for 20 min in 20× diluted Giemsa (nuclear stain). Cells received a final wash with bi-distilled water and were left to dry overnight. At least 200 cells were counted per treatment with a standard light microscope.

Live cells were stained with fluorescent markers for confocal microscopy at the time points specified in culture medium without amines and serum at 37°C for 30 min. Nuclear DNA was stained with Hoechst $(1 \mu M)$, cellular cytoplasm with CellTrack er^{TM} Red or CellTrackerTMBlue (10 μ M), and neutral LDs with Bodipy[®] 493/503 (8 µg/ml; Molecular Probes). Cells were washed three times in warm endotoxin-free PBS to remove unbound probe and fixed consecutively for 15 min in 2% and then 4% paraformaldehyde. Macrophages were classified as enlarged vacuole-positive (V+) when their cell size was ≥ 24 µm and multiple large vacuoles were present. Fluorescently labeled cells were viewed on a Leica TCS SP5 AOBS inverted confocal microscope with a 63× HCX PL Apo 1.4 oil objective and stacked images taken at 0.42μ m slices with a spectral photomultipliers DFC320 color camera (36-bit, 7 megapixels, nonconfocal). Training and technical support was provided by the Bio Imaging Core facility of the Flemish Institute of Biotechnology (Ghent).

Quantification of intracellular cholesterol content

Macrophage intracellular cholesterol was quantified using the Calbiochem® cholesterol/cholesteryl ester quantitation kit (Merck Millipore; catalog number 428901). Cell pellets $(1 \times 10^6$ cells) were freeze-thawed five times in liquid nitrogen, homogenized with a pestle in $200 \mu l$ chloroform:isopropanol:NP40 $(7:11:0.1, v/v/v)$ and centrifuged for 10 min at 18,000 *g*. The organic lower phase was transferred to a clean microcentrifuge tube and air-dried at 50°C. To remove any residual chloroform, samples were further vacuum-dried for 30 min at 45°C. The dried lipids were dissolved in 200 μ l cholesterol reaction buffer at 40°C for two cycles of heating (10 min) and vortexing with a final vortex of 5 min. All extracted samples and standards were added to a volume of 50 μ l in a black 96-well plate followed by 50 μ l of reaction mix 1 (with cholesterol esterase) or 2 (without cholesterol esterase) for determination of total cholesterol and free cholesterol, respectively. The assay plate was incubated at 37°C for 60 min and then fluorometrically measured on a FLUOstar OMEGA microplate reader (\sim 535/590 nm). Cholesterol standards were plotted against relative fluorescence units and the concentration of total, free, and esterified cholesterol (free subtracted from total) calculated as micrograms per 10^6 cells.

Statistical analyses

The number of mice used for experiments comprised a minimum of five mice per treatment. Data were obtained from at least three independent experiments or 500 cells from no less

than five separate microscopy images. The distribution of all data was determined by a Shapiro-Wilk (*W*) normality test. A nonparametric Mann-Whitney *U* test or Kruskal-Wallis one-way ANOVA (*H*; with Dunn's ranked sum multiple comparisons) was used to assess the proportion of foam cells in PEC cytospins and cellular cholesterol content (dependent variables) for each treatment or cholesterol fraction (categorical variables). Generalized linear model (GLM) analyses (Poisson distribution with sequential Sidak pairwise comparisons) were used to assess counts of the number of cellular vacuoles, LDs, and bacilli (dependent variables) for each treatment or MOI at specified time points (predictors). Image processing and quantification of cellular markers for confocal microscopy were conducted with Volocity 3D image analysis software (PerkinElmer Inc.). All other statistical analyses were performed with IBM SPSS Statistics 23 (IBM, Chicago, IL) and GraphPad Prism 5 (GraphPad Software, Inc.). Results are presented as mean ± SEM and are considered significant at $P \leq 0.05$.

RESULTS

MAs of the methoxy oxygenation class promote the formation of multi-vacuolar foam cells

To assess the influence of MA structure on the induction of multi-vacuolar giant foam cells, mice were intraperitoneally injected with 25 μ g of various MAs (α , methoxy, or keto) (Fig. 1) or control compounds (PBS or liposome carrier without synthetic MA). PECs were harvested 2 days after treatment. Macrophages were examined by cytospin analysis and laser-scanning-confocal microscopy. PECs from mice treated with PBS, Lipo, and α MA-containing liposomes contained <10% vacuolar foam cells (**Fig. 2A, B**). In contrast, PECs from mice treated with liposomes containing kMA or mMA showed \sim 15% and 27–30% vacuolar foam cells, respectively. GLM analysis showed a clear difference among the treatments in their ability to induce vacuoles. mMA clearly effected vacuole formation with significantly more enlarged V+ cells (25–35%) as compared with kMA $(5-10\%)$ and other treatments (5%) (Fig. 2C, D). In addition, the proportion of enlarged V+ cells from the mMA cell population remained significantly elevated over time when kept in culture. At the 48 h time point, the macrophages from mMA treatment remained enlarged, while all other treatments resulted in macrophages returning to normal size (<5% enlarged V+ cells) (Fig. 2C). MGCs were seen mainly among the enlarged V+ cells induced by mMA (Fig. 2E).

MAs of the keto oxygenation class induce foam cells rich in cholesterol-laden LDs

A clear difference was observed among treatments in their ability to induce LD accumulation (**Fig. 3**). kMA significantly induced LDs, up to 3-fold higher in comparison to all other treatments (Fig. 3A). Clearly, kMA was the strongest inducer of intracellular lipids in peritoneal mouse macrophages at all time points assayed. In order to normalize for LDs involved in regular macrophage metabolic activity, the LD number in PBS-treated cells was

Fig. 2. MAs of the methoxy oxygenation class promote the formation of multi-vacuolar foam cells. The proportion of vacuole-rich macrophages was determined by light microscope images of May-Grünwald-Giemsa-stained cells on cytospins and by laser-scanningconfocal microscopy (mean ± SEM). A: Percentage of foam cells in PEC fraction from the control (PBS and Lipo), α MA, mMA, and kMA treatments. (Shapiro-Wilk: *W* = 0.801, ***P* < 0.01; Kruskal-Wallis: $H = 12.100, *P < 0.05, df = 4; n = 3$ independent experiments; ns, not significant). B: Light microscope images of cytospins showing vacuolar foam cells in PEC fraction. Images were taken at 100× oil magnification. Arrows indicate MGCs. Scale bar: $10 \mu m$. C: Induction of enlarged V+ cells is shown for control (PBS and Lipo) and the various MA-treated mouse peritoneal macrophages over time, as measured by laser-scanning-confocal microscopy (GLM: Wald chi-square = 753.924, ****P* < 0.001, $df = 14$; n = 5 per time point; ns, not significant). D, E: Laser scanning confocal microscopy images showing enlarged V+ cells for PBS, α MA, and mMA treatments. Arrows indicate MGCs. Stacked images, 63× oil objective. Scale bar: 20 m. E: Zoomed images from the mMA treatment in (D).

selected as the baseline value against which relative increases or decreases of LDs in other treatments were compared. From this relative analysis, kMA again emerged as the MA class that significantly and prominently induced an increment in LDs in the macrophages (Fig. 3B, C). To further substantiate and quantify the accumulation of LDs in kMA-treated macrophages, the levels of intracellular active cholesterol and stored cholesteryl ester were determined in peritoneal macrophages isolated 2 days after oxygenated mMA or kMA treatment. Macrophages from the kMA treatment contained substantial intracellular esterified cholesterol, which also accounted for 77% of the total cholesterol content (Fig. 3D). Thus, the ratio of esterified-to-free cholesterol was distinctly elevated in the

Fig. 3. MAs of the keto oxygenation class induce foam cells rich in cholesterol-laden LDs. The induction of LDs is shown for peritoneal macrophages from control (PBS and Lipo) and α MA-, mMA-, and kMAtreated mice (mean ± SEM). A: LD accumulation in murine macrophages harvested at specified time points following the control and various MA treatments (GLM: Wald chi-square = 353.662, *P* < 0.001, $df = 14$; n = 5 per time point). B: Relative induction of LDs over time as compared with PBS (broken line). C: Laser-scanning-confocal microscopy images depicting variation in LD induction for PBS, α MA, and kMA treatments. Arrows indicate LD-filled cells. D: Cellular cholesterol content of variously treated cells. Bars represent total cholesterol, subdivided in the amount of free and esterified cholesterol for each of the treatments (micrograms per 10^6 cells). Upper panel, fraction percentages of free and esterified cholesterol are shown in the bars ($n = 12$ mice; Shapiro-Wilk: $W =$ 0.813, *P* < 0.05; Kruskal-Wallis: *H* = 24.726, *P* < 0.001, *df* = 3). Lower panel, the ratio of esterified-to-free cholesterol is given for the various treatments (Kruskal-Wallis: *H* = 26.554, *P* < 0.001, *df* = 3). E: kMA-induced LDs in peritoneal macrophages as identified by the neutral lipid probe, Bodipy® 493/503. C, E: Stacked images, $63 \times$ oil objective. Scale bar: $20 \mu m$. Significant *P* values were ranked as **P* < 0.05, ***P* < 0.01, and *** $P < 0.001$; ns, not significant.

LD-inducing kMA treatment and not in vacuole-inducing mMA treated cells (Fig. 3D-E).

MA-induced LD accumulation in macrophages promotes BCG proliferation

In order to determine to what extent LD induction by kMA or vacuole induction by mMA may affect mycobacterial growth, peritoneal macrophages from mice treated with kMA or mMA were infected with BCG-dsRed and cultured for 5 days (**Fig. 4A**). Control groups again consisted of peritoneal macrophages isolated from mice injected with PBS (placebo) or the unoxygenated α MA. Infection of the macrophages with BCG did not abrogate the accumulation of LDs in kMA-treated macrophages and did not induce, as such, an accumulation of LDs in placebo- or mMA-treated macrophages (Fig. 4B, upper panel). Strikingly, BCG replication was strongly enhanced in the kMA-treated macrophages (Fig. 4C), resulting in a near 3-fold increment in BCG numbers compared with the placebo-treated macrophages or macrophages treated with mMA (Fig. 4B, lower panel; Fig. 4D). Macrophages from mice treated with a natural mixture of MA made up of α MA (\sim 53%), kMA (\sim 9%), and mMA $(\sim 38\%)$ showed an intermediate induction of LDs as well as BCG replication (Fig. 4B), indicating that the presence of either mMA and/or α MA does not inhibit the LD-inducing and BCG-proliferative biological functions of kMA.

Assessment of mycobacterial growth in LD-accumulating macrophages from LXR-deficient mice

In order to determine whether foamy macrophages brought about by nontuberculous means would also display elevated intracellular mycobacterial growth similar to kMA-treated macrophages, we investigated this parameter in peritoneal macrophages from LXR-deficient KO mice. Macrophages deficient in LXR contained abundant LDs sustained over time (**Fig. 5A**). This was accompanied by intracellular accumulation of esterified-to-free cholesterol at a ratio of \sim 8:1 in KO mice (Fig. 5B). Strikingly, these LDaccumulating macrophages, elicited by a deficiency in LXR activity, also showed significantly increased mycobacterial growth as compared with macrophages from WT mice (Fig. 5C).

DISCUSSION

The hallmark response to Mtb infection is the formation of granulomas that, in spite of being crucial to host protection, are exploited by mycobacteria for survival and persistence (43). Granulomas contain distinct macrophage populations of which foam cells loaded with neutral lipids and giant cells containing abundant vacuoles are characteristic (44). It was shown previously that natural purified Mtb MA

Fig. 4. MA-induced LD accumulation in macrophages promotes BCG proliferation. Following in vivo treatment with MA, peritoneal macrophages were infected ex vivo with BCG-dsRed (MOI 1, broken line) and cultured until timed interval measurements were taken by laser-scanning-confocal microscopy (mean ± SEM). A: Laser-scanning-confocal microscopy images (zoomed) showing macrophages from mock or BCG infections. The presence of a BCG-dsRed bacillus can be clearly distinguished. Scale bar: $2.5 \mu m$. B: Upper panel, LD accumulation in peritoneal macrophages over time (GLM: Wald chi-square = 636.496, *P* < 0.001, *df* = 17; n = 5 per time point). Lower panel, proliferation of BCG bacilli over time (GLM: Wald chi-square = 250.303, *P* < 0.001, *df* = 17; n = 5 per time point). On the x-axis, "Mix" denotes a natural purified MA extract consisting of \sim 53% αMA, \sim 38% mMA, and \sim 9% kMA. C, D: Laser-scanning-confocal microscopy images depicting a clear difference in the presence of BCG-dsRed bacilli in peritoneal macrophages from mice treated with either kMA or mMA. Stacked images, $63 \times$ oil objective. Scale bar: $20 \mu m$. Arrows indicate BCG-dsRed bacilli. Significant *P* values were ranked as **P* < 0.05, ***P* < 0.01, and ****P* < 0.001; ns, not significant.

induces cellular features similar to those observed in granuloma cell populations (25). However, as MAs are heterogeneous, in this study, we determined to what extent examples of individual MA classes sustain the foamy and/or vacuolar giant cellular traits in macrophages. The clear difference reported here between mMA and kMA treatment in murine macrophages, a distinct induction by mMA of vacuoles without LDs and an upregulation of LDs without vacuoles by kMA, indicates the occurrence of independent triggers for giant versus foamy cell phenotypes during Mtb infection. The pathophysiological significance of these independent macrophage responses possibly reflects a strategy by Mtb to persist inside the key immune effector cell responsible for its elimination (45–47). In nature, the inflammatory neutral MA class makes up approximately 50% of all MAs, whereas the ratio of pro-inflammatory mMA to kMA classes in the Mtb bacillus cell wall is variable and may be tuned to complement growth and persistence (13, 17, 18). Korf et al. (25) previously showed that macrophages from mice treated with the natural MA mixture (similar to the MA mix used in this study) containing approximately 32–40% methoxy mycolates, accumulated large intracellular vacuoles like those reported by us here after treatment with pure (100%) mMA. Our results furthermore showed that the macrophages harvested from mice after natural MA mix treatment, which contains 7–15% keto mycolates, significantly accumulated intracellular LDs and allowed for mycobacterial growth, comparable to the macrophages from the pure (100%) kMA treatment group, but reduced by about half. One could therefore anticipate that the changes brought about by regulation of the MA class composition by Mtb will not be extreme.

LDs are widespread cell organelles dedicated to the storage of neutral lipids like cholesteryl esters (48). Numerous

studies have reported on the close association of this distinctive lipid organelle with Mtb pathogenesis (28, 46, 49–52). The dynamic interaction of the LD with the Mtbcontaining vacuole was recorded in an in vitro granuloma model by Peyron et al. (28). Electron microscopy analyses of foam cells showed that phagosomes containing bacilli migrated toward LDs and eventually engulfed them. D'Avila et al. (53) observed a dose-response in LD formation over time in a murine model of TB using the vaccine strain, BCG. Close associations between LDs and phagosomes were recorded in BCG-infected treatments, with significant LD induction observed after 24 h that remained elevated for about 2 weeks after infection. Cytoskeleton arrangement and phosphatidylinositol-3-kinase, an enzyme involved in cellular regulation of proliferation, trafficking, and metabolism (54), were identified to play key roles in the LD recruitment to bacilli-enclosed phagosomes (55). It is thus clear that pathogenic mycobacteria regulate biogenesis or accumulation of host cell lipids into LD-confined organelles, which may be further manipulated to intimately associate with the phagosomes enclosing the bacilli. An important question is the pathophysiological relevance of this association. Research suggests that the LD-phagosome interaction serves a dual purpose: at one end to benefit the microorganism and at the other to aid the host cell in microbe elimination (28, 32, 56). LDs may serve as a nutrient source to the bacillus through direct assimilation of host cell lipids (28) or as a nutrient reservoir during latency (32). In contrast, host cells employ LDs to disseminate important anti-bacterial factors, as was recorded in an in vitro study of latex bead phagosomes and pathogenic mycobacteria (56). Phagosomes infected with Mtb and exposed to distinct lipids (i.e., arachidonate, ceramide, and sphingomyelin) induced actin assembly in macrophages, a process

Fig. 5. Mycobacterial growth in LD-accumulating macrophages from LXR-deficient mice. A: Macrophages from WT and LXR-deficient mice were cultured for 5 days and the intracellular LD prevalence analyzed by laser-scanning-confocal microscopy. KO-cells contained \sim 10-fold more cellular LDs in comparison to WTcells, though no significant change in the number of LDs per 100 cells was observed over time (GLM: Wald chi-square = 99.093, *P* < 0.001, *df* = 5; n = 4 per time point; sequential Sidak pairwise comparisons *P* > 0.05). Laser-scanning-confocal microscopy images depicting clear morphological differences in LD prevalence between WT and KO macrophages. Scale bar: 20 μ m. B: Cholesteryl esters accounted for 88% of total cholesterol in LXR-deficient cells (left panel) that also contained significantly elevated ratios of intracellular esterified-to-free cholesterol (right panel; Mann-Whitney: *U* = 144.000, *P* < 0.001, *df* = 1; n = 12 mice). C: Macrophages from WT and KO mice were infected ex vivo for 6 h with BCG-dsRed (MOI 1, broken line) and cultured for 5 days. Mycobacterial growth was measured by laser-scanning-confocal microscopy at specified time points, and was significantly increased in LXR-deficient cells over time (GLM: Wald chi-square = 576.689, *P* < 0.001, *df* = 17; n = 6 per time point; sequential Sidak pairwise comparisons *P* < 0.05). Data represent mean ± SEM. Significant *P* values were ranked as **P* < 0.05 and ****P* < 0.001; ns, not significant.

involved in microtubule organization and LD trafficking (55). Lipid exposure also effected phagolysosomal fusion and decreased phagosome pH, events indicative of phagosome maturation and engagement of bactericidal machinery (56). It was noted that the kMA-induced LD accumulation in macrophages tended to rapidly return to normal after removing the kMA stimulation, while the mMA-induced vacuolation of the macrophage was maintained after removal of mMA. Enlarged intracellular vacuoles, like those induced after treatment with mMA, represent structures formed by invagination of the external cell membrane phospholipid bilayer into the cytosol (57). These double-layered vacuoles form stable structures that essentially provide a compartment for upholding the bacterium. LDs are phospholipid monolayer organelles critically involved in cellular lipid metabolism (58). As dynamic structures mediating lipid exchange or consumption, removal of the LD-inducing stimulus can be expected to coincide with rapid turnover and/or catabolism of the LDs by cellular machinery.

As indicated above, we confirmed that α MA is functionally neutral and showed that the oxygenated MAs effected pronounced changes with remarkable segregation of foam cell features: mMA induced enlarged multi-vacuolar cells, whereas kMA stimulated cells to accumulate LDs with most of its cholesterol converted to the cholesteryl ester storage form. We next verified the relevance of these responses to intracellular growth of mycobacteria. Using kMA-induced foamy cells and mMA-induced giant cells, we infected these macrophages ex vivo with BCG-dsRed and measured replication of the bacilli over time. Our results showed that intracellular growth was promoted in kMA-induced foamy macrophages, but not in mMA-induced giant cells. Macrophages classically convert into foam cells through a dysregulation in the balance between influx and efflux of cholesterol. Cholesterol is transported to peripheral macrophages in LDL that are taken up by the cell through the LDL receptor, or in the oxidized form, through the scavenger receptors, SRA and CD36. Cholesterol efflux is mediated by the transporters ABCA1 and ABCG1 that load the cholesterol on HDL for transfer to the liver, a mechanism known as reverse cholesterol transport. These processes are regulated by the lipid sensing nuclear receptors, LXR and PPAR γ . A direct role for LXR in mycobacterial foam cell formation has not been reported. However, mice deficient in LXR show spontaneous foam cell formation (59). LXR-deficient macrophages may be considered a cell model of the late stage of Mtb infection, where the macrophage has been modified to shut off its cholesterol export without limiting cholesterol import, aiming at providing a rich sterol nutrition source for mycobacterial growth and replication. We therefore investigated whether foam cells generated through a noninfectious condition, namely, a deficiency in LXR activity, would also facilitate growth and proliferation of BCG bacilli. Similar to macrophages from mice treated with kMA, LXR-deficient macrophages accumulated cholesteryl ester-rich LDs and were facilitative toward mycobacterial growth. LXR-deficient macrophages contained more than double the amount of cholesterol, compared with what could be induced with kMA, and facilitated proliferation to almost double the number of BCG mycobacteria per 100 cells after 96 h. These results suggest that LXR may indirectly function as a negative regulator of mycobacterial foam cell formation and mycobacterial growth. Although speculative, this proposition is in line with a previous report showing that LXRactivity suppresses the outgrowth of Mtb bacilli in a mouse model of pulmonary Mtb infection (60).

Our study addressed the individual contribution that examples of each of the three main classes of MAs from Mtb make toward the induction of foam cells and multi-vacuolar giant cells, as well as toward the facilitation of intracellular mycobacterial proliferation. All three features are important elements of the manifestation of TB in human lungs and were elicited to varying degrees by the individual MA classes studied. Thus, we showed here that it is kMA that induces mainly cholesterol ester accumulation within intracellular LDs and facilitates mycobacterial BCG proliferation. In contrast, mMA was found to induce vacuole formation, characteristic of giant cells found in Mtb lung granulomas, with no change in cholesteryl ester content or an improved ability to sustain and facilitate mycobacterial growth. Finally, MA treatment exerted a negligible effect on all three parameters. These findings suggest separate important roles for keto and methoxy oxygenated MA classes in manipulating the host macrophage response during the establishment of TB. This new insight may assist in deciphering and targeting the Achilles' heel of the tubercle bacillus according to an approach recently reviewed by Nataraj et al. (61), relying on the most recent understanding of how MA is synthesized and differentiated into classes by Mtb. Cyclopropanation provides the minimum functionalization associated with MA virulence (17). Dubnau et al. (18) provided evidence that cyclopropanation or functionalization of the distal group of the meromycolic chain represents the first step of a common pathway responsible for generating methoxy and keto mycolates. Differential regulation of this single mechanism can thus allow mycobacteria to manipulate MA class composition of α , methoxy, and keto mycolates during growth progression. Evidence for this was provided by Yuan et al. (19), who showed that keto mycolates are more abundant during the early stage of mycobacterial infection, but give way to methoxy mycolates that become more abundant in the later phase of infection. Our results imply that much may be accomplished by attempting to control the switch between methoxy and keto mycolates as a principle of drug treatment to combat TB. One argument for this was provided by Slama et al. (62), who showed that mMA was much more abundant in the virulent Mtb H37Rv strain than in the attenuated Mtb H37Ra strain. This has to be understood in the wider context that other differences between the H37Rv and H37Ra strains were also identified, but the researchers then achieved better resolution of the evidence (62): A recombinant Mtb H37Rv was generated with a single

gene KO of a protein involved in the late stage of MA biosynthesis, namely HadC. HadC is a dehydratase that is not catalytically active, but holds the longer form of the elongating merochain of MA in a complex with HadB, which catalyzes dehydration of a β -hydroxyacyl in the merochain to an enoyl. The *hadC* KO Mtb H37Rv showed a reduction in the abundance of mMA similar to that found in Mtb H37Ra (62). Accordingly, it also had lower virulence, approximating that of Mtb H37Ra, lost its cording and biofilming ability, and its envelope integrity, thus making it more susceptible to the hydrophobic drug, rifampicin. It should be noted that these biological effects could not only be ascribed to the lowering of the abundance of mMA, because the *hadC* KO Mtb H37Rv also had a larger degree of unsaturation in the merochain and could not produce the small amount of extra-long chain MAs of all three classes that are present in the WT Mtb H37Rv.

Whereas Slama et al. (62) demonstrated the advantage of being able to reduce the abundance of mMA in Mtb, our results could imply that even more may be achieved by interfering with kMA production in Mtb. However, Dubnau et al. (18) managed to create a *hma*-KO mutant Mtb H37Rv that produced no oxygenated MA at all. The mutant Mtb also showed severe loss of virulence, but was still able to grow and multiply in a monocytic cell line or in the lungs and spleen of infected mice. Thus, kMA seems not to be essential for survival of Mtb and is, hence, not a prime target for the conceptualization of a new anti-TB drug; although we have demonstrated here how kMA allows the mycobacterium to thrive, but not survive, in the target host cell.

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