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In Brief
First quantitative phosphoproteome analysis of TDM-activated macrophages reveals new insights in biological processes of macrophages stimulated with the mycobacterial cord factor. Surprisingly, the bioinformatic results revealed Mincle-dependent and -independent phosphorylation, which appear to affect different biological processes. Whereas PI3K/AKT signaling, dependent on Mincle, is involved in TDM-induced cytokine regulation, Mincle-independent phosphorylation and transcriptomic changes were linked to cell cycle regulation. Collectively, the observed reprogramming of macrophages by TDM might be relevant in the mycobacteria-macrophage interaction.

Highlights
- quantitative phosphoproteome analysis of TDM-activated macrophages.
- distinct Mincle-dependent and independent phosphorylation and gene regulations.
- Mincle-dependent activation of PI3K/AKT signaling by TDM.
- Mincle-independent macrophage response is linked to cell cycle regulation.

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Macrophage Phosphoproteome Analysis Reveals MINCLE-dependent and -independent Mycobacterial Cord Factor Signaling*

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Immune sensing of *Mycobacterium tuberculosis* relies on recognition by macrophages. Mycobacterial cord factor, trehalose-6,6'-dimycolate (TDM), is the most abundant cell wall glycolipid and binds to the C-type lectin receptor (CLR) MINCLE. To explore the kinase signaling linking the TDM-MINCLE interaction to gene expression, we employed quantitative phosphoproteome analysis. TDM caused up-regulation of 6.7% and suppressed 3.8% of the 14,000 phospho-sites identified on 3727 proteins. MINCLE-dependent phosphorylation was observed for canonical players of CLR signaling (e.g. PLCγ, PKCδ), and was enriched for PKCδ and GSK3 kinase motifs. MINCLE-dependent activation of the PI3K-AKT-GSK3 pathway contributed to inflammatory gene expression and required the PI3K regulatory subunit p85α. Unexpectedly, a substantial fraction of TDM-induced phosphorylation was MINCLE-independent, a finding paralleled by transcriptome data. Bioinformatics analysis of both data sets concurred in the requirement for MINCLE for innate immune response pathways and processes. In contrast, MINCLE-independent phosphorylation and transcriptome responses were linked to cell cycle regulation. Collectively, our global analyses show substantial reprogramming of macrophages by TDM and reveal a dichotomy of MINCLE-dependent and -independent signaling linked to distinct biological responses. *Molecular & Cellular Proteomics* 18: 669–685, 2019. DOI: 10.1074/mcp.RA118.000929.

Macrophages play a dual role in tuberculosis, because they harbor the intracellular *Mycobacterium (M.) tuberculosis* in the phagosome, but also initiate the successful immune response by production of chemokines and cytokines. Recognition of mycobacteria by macrophages depends on the presence of pattern recognition receptors of the Toll-like receptor (TLR) and C-type lectin receptor (CLR) families, and their interaction with mycobacterial ligands derived from the cell wall or mycobacterial nucleic acids. TLR2 recognizes the 19 kDa lipopeptide and Lipoarabinomannan (LAM) (1), whereas TLR9 is activated by mycobacterial DNA containing unmethylated CpG motifs (2, 3). Several SYK-coupled activating CLR contribute to the macrophage response to mycobacteria, including Dectin-1 (4, 5), Dectin-2 binding manLAM (6), DCAR as a receptor for phosphorytidyl-inositol mannosides (7), and MINCLE and MCL, which are receptors for the mycobacterial cord factor (8–11). The cord factor, trehalose-6,6'-dimycolate (TDM), is the major cell wall glycolipid and is enough to elicit several hallmarks of the mycobacteria-host interaction, i.e. inflammatory gene expression in macrophages, adjuvant activity toward Th1/Th17 responses, and granuloma formation in vivo (8, 12–14). TDM, and its synthetic glycolipid analog trehalose-6,6'-dibehenate (TDB), directly bind to MINCLE, and with lesser affinity to MCL (9). MINCLE-deficiency in macrophages and mice abrogates TDM-induced cytokine production, adjuvanticity and granuloma formation (8, 10, 15).

The transcriptional responses induced by ligation of TLR or CLR family members are overlapping, as may be expected based on their similarity in function as pattern recognition receptors, and include expression of pro-inflamma-
pitory chemokines and cytokines. However, there are also distinct response patterns linked to specific biological responses (16). For example, the TLR9 ligand CpG DNA induces high level IL-12 production and thereby provides a strong signal for Th1 differentiation. In contrast, the CLR ligands Curdlan (via DECTIN-1) and TDB (via Mincle) induce robust Th17-type CD4+ T cell responses through production of IL-1β and IL-23 (10, 14, 15, 17). These distinct gene expression patterns of macrophages stimulated with ligands for either TLR or CLR can be attributed to the differences in signal transduction pathways between both receptor families. TLR signaling is dependent on the adapter protein MYD88 (engaged by all TLR except TLR3) and/or TRIF (engaged by TLR3 and TLR4), the kinases IRAK4 and TBK1, and the E3 ligase TRAF6, to activate the transcription factors NF-κB and Interferon regulatory factors (IRF) (reviewed in (18)). TLR signaling has been intensely investigated in the last 15 years, including several proteomic and phosphoproteomic studies, revealing new kinases and signaling modules relevant for the transcriptional response to TLR ligands (19–21).

Activating CLR share signaling through an ITAM motif and the kinase SYK (22). Dectin-1 directly recruits SYK via a hemi-ITAM motif (23), whereas MCL, Mincle and DECTIN-2 associate with the ITAM-containing FcRγ chain to activate SYK (16). Downstream of SYK, activation of PLCγ2, phosphorylation of PKCδ (24), and assembly of the CARD9-BCL10-MALT1 complex is required for activation of NF-κB (25). PLCγ2 also triggers activation of the transcription factor NFAT and induces expression of EGR family members (26). However, in contrast to the TLR family, the signaling cascades downstream of SYK-coupled CLR have not been systematically studied using unbiased global proteomic and transcriptomic approaches.

Rapid signal transduction from pattern recognition receptors located at the cell membrane to the nucleus relies on multiple forms of post-translational modification (PTM) of proteins, altering the localization, interaction with other proteins, enzymatic activity or stability of the substrate proteins. Together with ubiquitination and acetylation, kinase-mediated phosphorylation is one of the best characterized molecular switches in cellular signaling (27). In CLR-induced macrophage activation, reversible phosphorylation regulated by kinases and phosphatases likely plays an essential role, as evidenced by the essential function of the kinases SYK and PKCδ, or the activation of the MAPK module, which lead to activation of transcription factors and thereby induce inflammatory gene expression. A comprehensive analysis of the changes in protein phosphorylation in response of macrophages to stimulation through a SYK-coupled CLR has not been reported yet. It is therefore unknown which kinases, transcription factors and other signaling proteins in addition to the above-mentioned players are regulated by phosphorylation in response to CLR stimulation in macrophages.

Here, we investigated global phosphorylation of proteins in macrophages of wild type and Mincle-deficient mice in response to stimulation with TDM to obtain new insight into the signaling and biological processes regulated through phosphorylation. To determine to which extent TDM-triggered protein phosphorylation is initiated through its receptor Mincle, we compared TDM-stimulated wild type and Mincle-deficient macrophages. Although the phosphorylation of many canonical kinases in the SYK-CARD9 pathway was induced in a Mincle-dependent manner, we observed a surprisingly large extent of Mincle-independent protein phosphorylation. To investigate whether gene expression induced by TDM depends entirely on its receptor Mincle, RNAseq analysis was performed. Consistent with the phosphoproteomic data, RNAseq analysis showed that a substantial fraction of the transcriptional response was at least partially independent of Mincle. These Mincle-independent changes in phosphorylation and gene expression were associated with cell cycle regulation and metabolic changes, whereas Mincle was required for signaling and transcription programs linked to cytokine production and innate immune response. Because activation of the PI3K-AKT-GSK3 axis was Mincle-dependent, we employed macrophages deficient in the PI3K regulatory subunit p85α and observed a significant contribution of this pathway to the production of cytokines by cord factor-activated macrophages.

**EXPERIMENTAL PROCEDURES**

**Mice and Macrophage Differentiation—** C57BL/6 and Mincle−/− mice (originally provided by the Consortium of Functional Glycomics) were bred at the animal facility of the Medical Faculty in Erlangen. PI3K p85α-deficient mice (28) were provided by Dr. Gernot Schabauer. Msr1−/− mice were provided by Dr. Matthias Trost, Fyb−/− mice by Dr. Annegret Reinhold, CASaR1−/− mice by Dr. Markus Boman, and p110γ,6−/− mice (29) by Dr. Sandra Beer-Hammer. All mice were bred on or backcrossed to a C57BL/6 background. Bone marrow cells were isolated and after erythrocyte lysis cultured in CD34+ (26) conditioned medium (LCCM) for 7 days. After overnight depletion of adherent cells, 5 to 6 x 10^6 of the nonadherent cells were seeded per 10 cm Petri dish. On day 3, 5 ml cDMEM + 10% LCCM were added, and differentiated bone marrow macrophages (BMM) were harvested on day 7.

**Stimulation, Cell Lysis and Phosphopeptide Preparation—** The glycolipid compounds TDM (Bioclot, Aidenbach, Germany) and TDB (Avanti Polar Lipids, Alabaster) were used in a plate-bound form. For coating onto tissue culture plates, TDM/TDB were dissolved in isopropanol (ISO) by incubation at 60 °C for several minutes and added to cell culture plates. ISO was evaporated by incubation at room temperature in a biological safety cabinet for several hours. TDM and TDB were used at 2 μg/ml or 5 μg/ml. Trehalose monoester (TMX) and diester (TDX) glycolipids containing C12 and C16 acyl chains were synthesized and purified as previously described (30) and coated for stimulation of macrophages at a concentration of 2 μg/ml as

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1 The abbreviations used are: TLR, toll-like receptor; CLR, C-type lectin receptor; LAM, lipooligosaccharide; IRF, interferon regulatory factor; PTM, post-translational modification.
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described above for TDM and TDB. Control ligands 0.5 μM CpG ODN 1826 (TIB Molbiol, Berlin, Germany) and 20 ng/ml LPS (Escherichia coli 055:BS; Sigma Aldrich, Schnelldorf, Germany) were diluted in cDMEM. overnight pre-stimulation of macrophages with 10 ng/ml LPS was performed in Petri dishes. The pharmacological AKT inhibitors Triciribine (5 μM) and AKT-Inhibitor VIII (10 μM; Merck Millipore, Darmstadt, Germany) as well as the PI3K p110 isoform-specific inhibitors PIK-75 (1 μM; Selleckchem) and TXG-221 (1 μM; Selleckchem, Houston, TX) and the PI3K inhibitor LY294002 (20 μM, EMD Millipore) were simultaneously added to the stimulation.

For phosphoproteome analysis 50 × 10⁶ cells per sample were stimulated for 45 min. After washing cells with cold PBS, macrophages were harvested in PBS containing protease and phosphatase inhibitors (1:50 Roche Complete (50×); 1:50 NaF (0.5 mM); 1:100 β-Glycerophosphat (1 mM); 1:200 Na₂VO₄ (200 μM)). The samples were centrifuged for 5 min, 1400 rpm at 4 °C and the pellets were lysed with 100 to 200 μl lysis buffer (1% Rapiigest, 50 mM Tris pH8.0, 1 mM TCEP). Afterward, 1 μl Benzonase was added.

pY Immunoprecipitation—After stimulation, cells were lysed with RIPA buffer containing protease and phosphatase inhibitors (Roche complete, 0.5 μM sodium fluoride, 1 μM β-glycerophosphate, 200 mM sodium orthovanadate). Cell lysates were sonicated and protein concentrations were assessed. Afterwards 500 μg protein were incubated with Pure Proteome Protein G Magnetic Beads (MERCK Millipore) decorated with 4G10 anti-phospho Tyrosine antibody overnight. After removing the supernatant and washing the beads, the phosphoproteins on the beads were reduced with Dithiotreitol and alkylated with iodoacetamide. The proteins were digested with Trypsin ( Pierce). Supernatants were collected, centrifuged (20.000 × g, 20 min, 4 °C), dried in the Speedvac and analyzed by LC-MS.

Experimental Design and Statistical Rationale—For the phosphoproteome analysis, we used LPS-primed macrophages from WT and Mincle-deficient mice and stimulated them with plate-coated TDM or ISO solvent control for 45 min. Three completely independent experiments were performed for macrophage differentiation and stimulation, thus a sample size of three biological replicates was used. After cell lysis the proteins were reduced, alkylated and digested with trypsin. Then the peptide samples from experimental conditions to be directly compared (WT-TDM versus WT ISO; and WT-TDM versus Mincle −/−) were labeled with light or heavy formaldehyde and dimethyl labeled (32) as follows: WT-ISO and WT-TDM (light), and subjected to fractionation by hydrophilic interaction liquid chromatography (HILIC) (33). Peptides were collected in 13 fractions and subjected to phosphopeptide enrichment using TiO² spin columns before LC-MS/MS analysis (34–36).

Dimethyl-labeled samples were separated on an Ultimate 3000 Rapid Separation LC Systems chromatography (Thermo-Fisher Scientific, Waltham, MA) with a C18 PepMap, serving as a trapping column (2 cm × 100 μm ID, PepMap C18, 5 μm particles, 100 Å pore size) followed by a 50 cm EASY-Spray column (50 cm × 75 μm ID, PepMap C18, 2 μm particles, 100 Å pore size)(Thermo-Fisher Scientific) with a linear gradient consisting of 2.4–28% (ACN, 0.1% formic acid (FA)) over 120 min at 300 nl/min. Mass spectrometric identification was performed on an Orbitrap Fusion Tribrid mass spectrometer (Thermo-Fisher Scientific) operated in data-dependent, positive ion mode similar to previously published (35, 37). FullScan spectra were acquired in a range from 400 m/z to 1500 m/z, at a resolution of 120,000, with an automated gain control (AGC) of 300,000 ions and a maximum injection time of 50 ms. HCD fragmentation was performed at 33% collision energy for all included precursor ions and MS/MS fragments were detected in the linear ion trap mass analyzer in rapid mode.

Protein identification and quantification were performed using MaxQuant Version 1.5.3.17 (38) with the following parameters: stable modification carbamidomethyl (C); variable modifications phosphorylation (STY), oxidation (M), acetylation (protein N terminus), deamini- dation (NQ), hydroxyproline (P), quantification labels Dimethyl and Dimethyl:2H₄ on N-terminal and/or lysine, and trypsin as enzyme with 2 missed cleavages. Search was conducted using the Uniprot-Trembl Mouse database (55,505 entries, downloaded 2015, including common contaminants. Mass accuracy was set to 4.5 ppm for precursor ions and 0.5 Da for ion trap MS/MS data. Identifications were filtered at a 1% false-discovery rate (FDR) at the protein level, accepting a minimum peptide length of 5 amino acids. Quantification of identified proteins referred to razor and unique peptides and required a minimum ratio count of 2. Dimethyl-based relative ratios were extracted for each protein/conditions and were used for downstream analyses.

Next Generation Sequencing—For RNAseq analysis 0.5 × 10⁶ BMM were stimulated with 2 μg/ml plate-coated TDM or evaporated isopropanol as solvent control for 24 h. Total RNA was then isolated using the PEGold RNA Micro Kit (Peqlab Biotechnology GmbH, Erlangen, Germany) according to the manufacturer’s guidelines. RNAs were stored at −80 °C and sent to the Next Generation Sequencing Core Unit of the University Hospital Erlangen for RNA sequencing. Quality and integrity of the isolated RNAs was confirmed by the Agilent 2100 Bioanalyzer (Agilent Technologies, Waldbrunn, Germany) with all RNAs having RIN > 8.5. RNAseq library preparation was done with pooled technical replicates using the TrueSeq stranded mRNA Library Prep Kit (Illumina, Inc., San Diego, CA) and sequencing was performed on the Illumina HiSeq 2500 platform (100 bp single-end) (Illumina, Inc.). Trimmed sequencing reads were aligned to the Mus musculus reference genome GRCm38 using the RNA-seq aligner STAR (version 2.5.3a) (39). For gene level quantification the software package Salmon (40) was used. Data normalization (TMM, edgeR, Bioconductor R-package) and statistical analysis for identification of differentially expressed genes were performed using the limma Bioconductor R-package (41). Because of consistent sequencing depth
across all RNA samples, limma-trend was used for differential expression analysis. Based on a classical interaction model, differentially expressed genes were determined according to the following criteria: adjusted (Benjamini-Hochberg) p value < 0.05, log2 fold-change > 1. Gene ontology (GO) analyses were performed using Cytoscape BiNGO (Cytoscape version 3.5.1) (42) (Benjamini-Hochberg false discovery rate (FDR) correction p value <0.05 and hypergeonomic distribution). For pathway enrichment analysis the InnateDB (http://www.innate.db.com) analysis platform was used (Benjamini-Hochberg (FDR) correction p value <0.05 and hypergeonomic distribution).

**Biological Formative Analysis**—Hierarchical clustering and visualization of regulated phosphopeptides in WT and Mincle+/− macrophages by TDM was done in a heat map and histogram using GProX Version 1.1.13, (43). To define sets of MINCLE-dependently and -independently regulated phosphorylation sites, the following parameters were used: MINCLE-dependent phosphorylation sites (TDM versus WT: fold change ≥ 1.5; p ≤ 0.05 and WT versus Mincle−/− fold change ≥ 1.5) and MINCLE-independent phosphorylation sites (TDM versus WT: fold change ≥ 1.5; p ≤ 0.05 and WT versus Mincle+/− fold change ≤ 1.25).

**GO and Pathway Analysis**—Using innateDB (www.innate.db.com Version 5.4) the over-represented GO terms and pathways over the genomic background for MINCLE-dependent or -independent phosphorylation sites were analyzed (hypergeometric distribution and Benjamini-Hochberg correction).

**Kinase Motifs**—Phosphorylation sites were analyzed for over-represented kinase motifs using motif-x (motif-x.med.harvard.edu, Version 1.2 10.05.06). MINCLE-dependent or independent phosphorylation sites were analyzed versus our own reference list containing all phosphorylation sites detected in at least 2 replicates (7436 peptides). The following parameters were used: width = 13; occurrences = 10; significance 0.002.

**Western Blotting**—For Western blot analysis, cellular lysates were prepared in RIPA buffer containing proteinase and phosphatase inhibitors (Roche complete, 0.5 μg sodium fluoride, 1 μg β-glycerophosphate, 200 μM sodium orthovanadate). Western blotting was performed by 12%-SDS-PAGE and wet-blotting. The following Abs were used: anti-pSYK (Tyr525/526), anti-SYK, anti-pERK (Thr202/Tyr204), anti-pGSK-3 (Ser9), anti-phospho-p38 (Thr180/Tyr182) and anti-p38 (Cell Signaling, Frankfurt, Germany), anti-Mincle (clone 4A9, MBL, Woburn, MA), anti-phospho-Tyrosine 4G10 (MERCK Millipore), anti-α-Tubulin (Sigma Aldrich) and anti-GRB2 (BD Biosciences, Heidelberg, Germany) as loading control and HPR-conjugated secondary Abs (Jackson Immuno Research Laboratories, Ely, UK).

**qRT-PCR**—Cellular lysates for quantitative RT-PCR were prepared with Trifast (Peqlab, Erlangen, Germany) to perform phenol/chloroform isolation according to manufacturer’s protocol. cDNA synthesis was done using the High Capacity cDNA Reverse Transcription Kit (Applied Biosystems). For qRT-PCR primers and probes were selected from the Universal Probe library (Roche) and purchased from Metabion. Hprt was used as housekeeping control and fold changes were calculated by the ∆∆CT method.

**Griess-assay and ELISA**—Nitrite and cytokine concentrations were analyzed in the supernatant of 200,000 cells in 96-well plates (triplicates) after 48 h stimulation unless otherwise stated. Cytokine concentrations were determined by DuoSet Sandwich ELISA (R&D Systems, Wiesbaden, Germany) according to the manufacturer’s protocol. NO production was assessed by measuring nitrite levels with the Griess assay. 10 ng/ml IFNγ (Peprotech, Hamburg, Germany) was used as costimulatory ligand for the Griess-assay.

**Protein Concentration**—To quantitate the protein content, 200,000 cells per well in 96-well plates (triplicates) were lysed with RIPA buffer containing proteinase and phosphatase inhibitors (see WB) and protein concentrations were assessed by BCA Protein Assay Kit (Thermo Fisher) according to the manufacturer’s protocol.

**MTT Conversion Assay**—To quantitate macrophage content in the wells, a colorimetric assay measuring the activity of cellular NAD(P)H oxidoreductases as conversion of the tetrazolium dye MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) was performed. MTT (20 μl of a 5 mg/ml stock solution) was added to the cultures at the indicated time points. After 3 h of incubation at 37 °C, the formazan crystals formed were dissolved by addition of 150 μl of a 10% SDS solution in HCl and incubation overnight. The OD570 nm was then measured in an ELISA reader.

**Statistical Analysis**—Statistical analyses were performed using Prism5 (GraphPad Software). Significance was determined by unpaired Mann-Whitney test for non-Gaussian distribution; *p ≤ 0.5, ** p ≤ 0.01.

**RESULTS**

**Kinetics and MINCLE-dependence of TDM/TDB-induced SYK/MAPK Activation and Cytokine Expression**—MINCLE expression is low in resting macrophages but induced by TLR and CLR stimuli. To achieve robust MINCLE expression, enabling a synchronized signaling response to MINCLE triggering, we primed bone marrow-derived macrophages (BMM) over night with a relatively low dose of LPS (10 ng/ml). Stimulation with the cord factor trehalose-6-6’-dimycolate (TDM) or the synthetic glycolipid adjuvant trehalose-6,6’-dibehenate (TDB), both used in plate-coated form, triggered robust phosphorylation of SYK and of the MAPK ERK1/2 and p38, peaking between 40’ and 120’ after stimulation (Fig. 1A, 1B). Production of the cytokines G-CSF and TNF in response to TDM/TDB stimulation was detectable first after 120’ (G-CSF) and 60’ (TNF) (Fig. 1C, 1D). Confirming previous results (10, 45), kinase activation and cytokine production were almost completely MINCLE-dependent (Fig. 1). Activation of SYK and MAPK preceded expression and secretion of cytokines; between 40’ to 60’ after stimulation, we observed strong protein phosphorylation but no cytokine secretion. Thus, we decided to use this time window for unbiased phosphoproteome analysis, when changes in phosphorylation are likely directly induced by interaction of TDM with its receptor and not influenced by secondary effects of secreted cytokines.

**Unbiased Detection of TDM-induced Protein Phosphorylation**—Ligand binding by MINCLE leads to tyrosine phosphorylation of the kinases SYK and PKCδ (24, 45, 46). We therefore first used the antibody 4G10 to detect changes in tyrosine phosphorylation of proteins after macrophage stimulation by Western blotting, which revealed several differential bands in response to the MINCLE ligands TDB/TDM and the TLR9 ligand CpG ODN, respectively (supplemental Fig. S1A). Mass spectrometry analysis of 4G10-enriched immunopre-
cipitation samples obtained 20’ and 60’ after stimulation with TDB was then used to identify tyrosine-phosphorylated proteins (supplemental Fig. S1). A total of 42 tyrosine-phosphorylated proteins were detected in two independent experiments (supplemental Table S1), among them the established MINCLE-signaling protein PKCδ, the PI3K regulatory subunit 4 (Pik3r4, aka VPS15), and GSK-3, consistent with activation of the PI3K-AKT-GSK3 signaling pathway; the small GTPase RAB22a, shown to be recruited to phagosomes; and the cell cycle-associated proteins PRP4 and RBM14 (supplemental Fig. S1).

Tyrosine phosphorylation constitutes only a small fraction of cellular protein phosphorylation, whereas phosphorylation of serine and threonine residues accounts for most protein kinase substrates. To globally detect protein phosphorylation on serine, threonine and tyrosine residues in a quantitative manner, we employed a protocol combining dimethyl labeling of cell lysates, enrichment of phosphopeptides by HILIC and titanium dioxide beads, and LC-MS/MS (Fig. 2A). In three independent experiments, we identified a total of 14173 phosphopeptides from 3727 phosphoproteins (Fig. 2B; supplemental Table S2). In accordance with previous work in LPS-stimulated BMM (21), most phosphorylated amino acids were serine (83%) and threonine (14%) residues, and only 3.4% of all sites were phospho-tyrosines (Fig. 2C). Treatment of WT macrophages with TDM significantly (1.5-fold, \( p < 0.05 \)) up-regulated 846 (6.7%) and suppressed 484 (3.8%) of all phosphopeptides (Fig. 2D, left panel), whereas the comparison of WT and MINCLE-deficient TDM-stimulated macrophages showed 641 (5.5%) induced and 388 (3.4%) reduced phosphopeptides (Fig. 2D, right panel). Given the similar extent of regulation in these comparisons, we next determined whether TDM-induced regulation is MINCLE-dependent.

MINCLE-dependence of TDM-induced Protein Phosphorylation—The 1330 phosphopeptides significantly regulated by TDM in WT macrophages were analyzed by hierarchical clustering and visualized using a heatmap representation to determine whether up- or downregulation depended on MINCLE (Fig. 3A). For a large fraction of these phosphopeptides, regulation was similar in both comparisons, indicating that TDM-induced regulation was MINCLE-dependent (indicated by subgroups C1 and C5 in Fig. 3A). However, a substantial portion of TDM-regulated phosphopeptides did not differ between WT and MINCLE-deficient macrophages or were even regulated more strongly in the absence of MINCLE (subgroups C4 and C6 in Fig. 3A). Focusing on phosphopeptides induced by TDM-stimulation in WT macrophages, we used histogram plots to overlay the log2-fold change distribution of WT versus MINCLE-deficient TDM-stimulated macrophages, which revealed that indeed around half of all phosphopeptides upregulated by TDM were not significantly more abundant in WT than in MINCLE-deficient macrophages (Fig. 3B).

To analyze whether MINCLE-dependent and -independent kinase pathways may be linked to distinct cellular components and biological responses to TDM stimulation, we performed bioinformatic analyses using the InnateDB Gene Ontology and Pathway enrichment analysis tools (Fig. 3C, 3D; supplemental Table S3). Indeed, MINCLE-dependent phosphoproteins were associated with the GO terms “late endosome/lysosomal membrane,” “endoplasmic reticulum,” and
“cytoskeleton,” “innate immune response,” and with signaling pathways controlled by the adapter protein FcRγ and by PLC, two canonical players of MINCLE-signaling through SYK. In contrast, Mincle-independent phosphoproteins were enriched for the cellular components “early endosomal membrane” and “nuclear membrane,” for the biological processes or pathway terms “RNA splicing,” “chromatin organization/remodeling” and “cell cycle.”

Next, we employed the algorithm motif-X to search for footprints of MINCLE-dependent and -independent kinase activation in the phosphopeptide data set. The sets of significantly induced phosphopeptides were analyzed for enrichment compared with a reference set of 7436 peptides detected in at least 2 experimental replicates (supplemental Fig. S2). Mincle-dependent phosphopeptides showed the strongest enrichment for a novel LxSP motif, which has not been associated with a putative mammalian kinase yet, but was recently reported as a candidate substrate motif for the CMV-encoded kinase pUL97 (47). The substrate motifs of PKA and of PKCδ, an established kinase in the MINCLE pathway linking SYK to CARD9 phosphorylation (24), were enriched in the MINCLE-dependent set of phosphopeptides. In addition, a substrate motif for the kinase GSK-3, which was tyrosine-phosphorylated in response to TDB (supplemental Fig. S1), was also detected in the MINCLE-dependent phosphopeptide set. On the other hand, motif-X analysis of MINCLE-independently induced phosphopeptides yielded exclusively a motif for the DNA-damage kinases ATM/ATR, which was previously also found to be enriched in the phosphoproteome of macrophages activated through TLR4 by LPS (21).

A graphical overview of the phosphoproteome data set is provided in Fig. 4. First, we confirmed MINCLE-dependent phosphorylation of several established C-type lectin receptor signaling proteins, including the phospholipase PLCγ2 and PKCδ, which are upstream of the CARD9-BCL10-MALT1 complex; activation of the MAPK signaling module with phosphorylation of MEKK1 (MAP3K1), and ERK1 (MAPK3) and ERK2 (MAPK1). Next, we identified the Src-family kinases LYN and FYN, as well as the tyrosine kinase FES, as Mincle-dependent TDM-activated phosphoproteins. The serine/thre-
 kinase PRKD2, a downstream kinase mediating effects of PKC, has not been described in the Mincle-pathway yet.

Several members of the NF-κB signaling module were detected as phosphoproteins, including IKBKB (IKKβ), IKBKG (NEMO), and the transcription factors NFKB2 (p100), NFKBIE (IκBβ) and NFKBIZ (IκBζ). Although regulation of the phosphosite in these proteins did not pass our thresholds, a closer look at the data in supplemental Table S2 indicated regulation of NF-κB signaling by TDM: phosphorylation of the kinase IKKβ at Ser-672 was reduced 1.43-fold (p < 0.03) and phosphorylation of the transcription factor NFKB2 at Ser-858 was increased 1.52-fold (p = 0.09). Consistent with the GO term enrichment of endosomal proteins, several transporters for chloride (CLCN3, CLCN7) and protons (ATP6V0A2, TCIRG1 (48)), the metallocarboxylase STEAP3 (49), and the endosomal GTPase RAB8A (50), were MINCLE-dependently phosphorylated in response to TDM, pointing to regulation of endosomal maturation and trafficking processes. The phosphorylation of 9 solute carrier proteins (SLC) suggests that MINCLE signaling may have a larger impact on the exchange of amino acids, sugars and other nutrients into and out of the cell, or between different macrophage compartments.

With regard to Mincle-independent phosphorylation, the adapter proteins DAB2 and FYB, the tyrosine kinase HCK and the focal adhesion kinase PTK2 may be involved in the phosphorylation of several cytoskeletal (MYOF, MARCKS), ribosomal (EIF4G1, EIF4H1) and nuclear proteins, such as LMNA and PCM1, the transcription factors ARNT and FOXN3, chromatin remodelers (MECP2, REST) and constituents of PML nuclear bodies (ATRX, SP100, SUMO1).

Interestingly, several cell surface receptors were phosphorylated in response to TDM stimulation (including the scavenger receptor MSR1, the complement receptor C5ar1, the chemokine receptor CXCR4, and the cation channel TRPV2, raising the possibility of parallel signaling pathways triggered by TDM stimulation, which may account for at least some of the MINCLE-independent phosphorylation events. Macrophages from several knockout mouse lines (Msr1−/−, Fyb−/−, C5ar1−/−) were used to test whether these cell surface receptors and adapter proteins phosphorylated in response to TDM play an essential role for the signaling and cytokine production. No differences were observed in the extent of phosphorylation of SYK, AKT or ERK1/2, nor in the release of G-CSF, IL-6 or CCL2, in the absence of MSR1 (supplemental

Fig. 3. Mincle-dependence of TDM-induced protein phosphorylation. A, Heatmap of regulated phosphopeptides WT and Mincle−/− macrophages (significantly regulated in TDM versus ISO; p < 0.05). B, Histogram of TDM-upregulated phosphorylation sites in WT and Mincle−/− macrophages (TDM versus ISO fc ≥ 1.5 and p < 0.05). Red bars show distribution of fold changes for the TDM versus ISO comparison in WT macrophages; blue bars for the comparison WT versus MINCLE-deficient macrophages after stimulation with TDM. C, D, Bioinformatic analysis of MINCLE-dependent and -independent phosphoproteins using InnateDB. The MINCLE-dependent group was defined as TDM versus ISO fc ≥ 1.5 p < 0.05 and WT versus Mincle−/− fc ≥ 1.5 (300 phosphorylation sites/226 phosphoproteins). The MINCLE-independent group was defined as TDM versus ISO fc ≥ 1.5 p < 0.05 and WT versus Mincle−/− fc ≤ 1.25 (396 phosphorylation sites/311 phosphoproteins). C, GO term enrichment. D, Pathway enrichment analysis.
MINCLE-dependent Activation of PI3K-AKT Signaling by TDM—Several phosphoproteins associated with PI3K-AKT signaling were detected in the phospho-tyrosine and phosphoproteome data sets, including the regulatory PI3K subunit p85α (PIK3R1), AKT1/2, and its substrates AKT1s, TSC2 and GSK-3 (Fig. 4; supplemental Fig. S1). Although AKT activation in response to TDB was recently reported (51), the canonical AKT phosphosites Thr308 and Ser473 were not detected in the proteomic data sets. We therefore analyzed AKT phosphorylation by immunoblot and observed that stimulation of LPS-primed macrophages with TDM and TDB, but not with the TLR9 ligand CpG, induced robust phosphorylation of AKT at Thr308 and Ser473 (Fig. 5A), which correlates with AKT kinase activity. Phosphorylation of AKT started 40 sec after stimulation and was almost completely dependent on MINCLE (Fig. 5B).

We next tested the functional importance of AKT activity by employing the inhibitors Triciribine and AKT-Inhibitor VIII, which both suppressed the production of G-CSF, whereas release of IL-6 and NO were not affected (Fig. 5C). Phosphorylation of AKT in response to TDM was completely abrogated by treatment with the PI3K inhibitor LY294002, which also inhibited G-CSF production (supplemental Fig. S6A–S6C).

There are four isoforms of the class I PI3K catalytic subunit, of which p110α and p110δ are mainly expressed in hematopoietic cells.
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Fig. 5. Activation of PI3K-AKT signaling by TDM. A, Phosphorylation of AKT on T308 and S473 by TDM and TDB. Stimulation of LPS-primed BMM for 1h with TDM, TDB or CpG. B, Phosphorylation of AKT is MINCLE-dependent. Stimulation of LPS-primed BMM with TDM for indicated time points in C57BL/6 or Mincle−/− mice. C, Cytokine production and NO release were analyzed from TDM-stimulated or non-stimulated BMM which were simultaneously treated with AKT inhibitor Triciribine or AKT Inhibitor VIII (n = 3). D, Reduced TDM-induced AKT phosphorylation in PI3K p85α−/− BMM compared with WT. Analysis of AKT phosphorylation after 1 h stimulation with TDM or CpG in LPS-primed and after 4 h stimulation in non-LPS-primed WT and p85α−/− macrophages. E, Reduced TDM-induced cytokine (48 h, +IFNγ) and NO (48 h, +LPS) levels in p85α−/− BMM compared with WT (2 independent experiments with 2 mice each per genotype). F Gene expression analyzed by RT-qPCR of LPS-primed BMM stimulated with TDM or CpG for 24h. n = 5 mice pooled from 3 independent experiments. n.s. = not significant.

edic cells (52). Macrophages derived from p110γ/δ double-deficient mice responded with normal AKT phosphorylation and unaltered G-CSF production to TDM; in contrast, combined treatment with inhibitors specific for p110α and p110β (PIK-75 and TGX-221, resp.) prevented AKT phosphorylation and G-CSF release in WT and p110γ/δ-deficient macrophages (supplemental Fig. S6D, S6E), suggesting that p110α and/or p110β are the catalytic isoforms responsible for phosphorylation of AKT and expression of G-CSF. The catalytic subunits p110α, β, δ all associate with the regulatory subunit PIK3R1 (aka p85α) (supplemental Fig. S6F), which stabilizes them and contains an SH2 domain for interaction with tyrosine phosphorylated proteins (52). Therefore, we employed p85α−/− macrophages as a genetic tool to probe the functional role of PI3K activation. In the absence of p85α, phosphorylation of AKT on Ser473 and on Thr308 was strongly reduced after TDM-stimulation of LPS-primed and of resting macrophages (Fig. 5D). Phosphorylation of the AKT substrate GSK-3 at Ser9 was also reduced in non-primed p85α−/− macrophages (Fig. 5D). These signaling defects were selective, because phosphorylation of SYK and ERK1/2 was unaltered (not shown). Production of NO, G-CSF and IL-6 were all significantly reduced in p85α−/− BMM compared with WT, although the levels of IL-10 detected in the supernatant were not altered (Fig. 5E), p85-deficient macrophages showed significantly reduced expression of IL-10 mRNA (Fig. 5F).

Transcriptome Analysis of Macrophage Activation by TDM Reveals MINCLE-dependent and -Independent Gene Expression—The unexpectedly prominent MINCLE-independent phosphorylation events prompted us to re-assess the notion that transcriptional activation of macrophages by TDM is entirely dependent on MINCLE-signaling via FcRγ-SYK-CARD9 (10, 45). WT and MINCLE-deficient macrophages stimulated for 24 h with TDM were analyzed by RNAseq using Illumina HiSeq. Hierarchical clustering of differentially expressed genes showed that resting WT and MINCLE-deficient
Macrophages clustered closely together. Stimulation with TDM caused substantial up- and downregulation of gene expression in WT macrophages, with subclusters of apparently MINCLE-dependent and -independent gene expression (Fig. 6A). This impression was confirmed by applying fold change filtering, with 211 clearly MINCLE-dependent and 339 -independent TDM-induced genes (Fig. 6B). K-means clustering of TDM-regulated genes (supplemental Table S4) was then used (Fig. 6C) to define gene sets with a characteristic pattern of induction and MINCLE-dependence for further bioinformatics analysis (Fig. 6D,6E). Clusters 1 and 8 contained the largely or partially MINCLE-dependently induced TDM target genes, which were enriched for pathway annotations of Interferon signaling, TLR- and NF-κB-signaling, and innate immune response. In contrast, MINCLE-independent gene expression (clusters 4 and 6) were associated with metabolic alterations (cholesterol biosynthesis, glycolysis, metabolism of lipids) and cell cycle regulation, respectively (Fig. 6C, 6D). Cluster 6 was also associated with “ATR activation,” consistent with the enrichment of the ATM/ATR kinase motif found in the set of phosphopeptides upregulated in a MINCLE-independent fashion by TDM (supplemental Fig. S2). Together, the RNAseq data set revealed a similar degree of MINCLE-independent responses as observed in the phosphoproteome data set, supporting the notion that not all TDM-triggered signaling is caused by binding to MINCLE. Moreover, although the transcriptomic and phosphoproteome data set cannot be compared directly, bioinformatics analysis yielded a concordant enrichment of terms associated with innate immune response for the MINCLE-dependent changes at the level of transcription and phosphoproteins. In contrast, the MINCLE-independent responses in both data sets were enriched for annotations linked to cell cycle regulation.

The MINCLE-independent TDM-induced Expression Program Is Also Upregulated by Trehalose Ester Glycolipids Not Binding to Mincle—To validate the findings from the RNAseq data set, qRT-PCR analysis of samples from independent experiments was performed. As expected, induction of mRNA expression for G-CSF and iNOS (belonging to the MINCLE-dependent TDM-induced expression program) was confirmed. Interestingly, MINCLE-independent upregulation of these genes was also observed. This suggests that Trehalose Ester Glycolipids, which do not bind to MINCLE, can activate the same cholesterol-dependent signaling pathways as TDM.

**Fig. 6. RNAseq of TDM-induced macrophage activation.** A, Hierarchical clustering of differentially expressed genes. B, Numbers of differentially expressed genes: TDM-induced genes (Log2FC (WT_TDM/WT_mock) > 1.58, adj. p value < 0.05), TDM-repressed genes (Log2FC (WT_TDM/WT_mock) < -1.58, adj. p value < 0.05); MINCLE-dependent: Log2 FC (WT_TDM/Mincle ~ TDM) > 1.5, and MINCLE-independent: Log2FC (WT_TDM/Mincle ~ TDM) < 1/~1.5. C, K-means clustering showing separation in MINCLE-dependent and -independent TDM target genes. D, selected GO and pathway terms enriched in different clusters.
dependent clusters 1 and 8) was largely dependent on MINCLE (Fig. 7A). In contrast, the expression of the chemokine CCL22 and the chemokine receptor CCR1 (both contained in cluster 4) was partially (Fig. 7B), and of CCL2 and the NAPDH oxidase NOX1 (two cluster 6 genes) was largely MINCLE-independent (Fig. 7C, 7D). To determine whether the MINCLE-independent response to TDM is indeed specific for the cord factor or can be caused more broadly by (glyco-)lipids, we tested a set of trehalose monoester (TMX) and diesters (TDX) with acyl chains of various length (C12 and C16). In previous work, we had observed that the trehalose diester form of C16 bound to MINCLE and triggered cytokine production form macrophages (53). Confirming our previous results, the trehalose diester of C16 and TDM diesters (TDX) with C12 and C16 acyl chains, and by the fatty acid (FA) behenate (C22). E, Csf3, F, Dusp5, G, Nox1, H, Ccl2. Mean and S.E., n = 5–6 mice, pooled from 3 independent experiments, * p < 0.05 WT versus Mincle−/−; # p < 0.05 treatment versus mock in WT BMM. All other responses to identical stimuli were not significantly different between WT and Mincle−/− BMM.

The MINCLE-independent Response to TDM and Related Glycolipids Is Linked to Macrophage Survival Or Proliferation—Because GO and pathway enrichment analysis revealed that cluster 6 was enriched for “cell cycle” and “DNA synthesis,” we validated the expression of the cluster 6 genes Myc, Ccne1, Ccne2, and Chek1 by qRT-PCR and confirmed their MINCLE-independent induction after TDM stimulation (Fig. 7F–7H). Upregulation of Dusp5, Nox1 and Ccl2 was largely independent of MINCLE, with a partial reduction after TDM stimulation.
Fig. 8. **Cell cycle-associated gene expression is MINCLE-independent.** A, WT and Mincle^-/- BMM were stimulated for 24 h as indicated with TDM, TDB, the trehalose monoesters (TMX) and diesters (TDX) with C12 and C16 acyl chains, and the fatty acid (FA) behenate (C22). Gene expression of Ccne1, Ccne2, Chek1 and Myc was then analyzed by qRT-PCR. Mean and S.E., n = 5–6 mice, pooled from 3 independent experiments. *p < 0.05 WT versus Mincle^-/-; # p < 0.05 treatment versus mock in WT BMM. All other responses to identical stimuli were not significantly different between WT and Mincle^-/- BMM. B, BCA assay for analysis of cellular protein content in BMM cultures stimulated for the indicated times with TDM, TDB or CpG ODN 1826. Pooled from four experiments. C, MTT conversion assay from BMM stimulated for 48h. Mean and S.E., n = 5–6 mice, pooled from 3 independent experiments. *p < 0.05 WT versus Mincle^-/-; # p < 0.05 treatment versus mock in WT BMM.

8A). As observed for DUSP5, CCL2 and NOX1 (Fig. 7E–7G), expression of these cell cycle-associated genes was not only upregulated in a MINCLE-independent manner by the cord factor, but to a similar extent also by synthetic trehalose diesters and monoesters of C12 and C16 fatty acids, and more weakly by behenic acid alone (Fig. 8A). Ccne1 and Ccne2 encode the cyclins E1 and E2, whose expression peaks in mid-G1 to early S-phase of the cell cycle (54). The proto-oncogene Myc encodes the E-box transcription factor MYC which is associated with proliferation and self-renewal of macrophages (55, 56). CHEK1 is activated by the DNA-damage response kinase ATR and contributes to p53-dependent responses (57). We observed that macrophage cultures treated with TDM appeared much denser than control cells cultured in medium alone (supplemental Fig. S7A). This effect was also observed with the trehalose monoester and diester glycolipids, and appeared to be partially MINCLE-independent (supplemental Fig. S7A). This impression was quantitatively confirmed by measuring the protein content of the macrophage cultures (Fig. 8B). Of note, when analyzed...
over time, the protein content of control cultures diminished, but stayed constant in the TDM-treated samples, suggesting that TDM prevented a loss of macrophages during starvation from growth factors, which was partially MINCLE-dependent after 48 h (Fig. 8B). Interestingly, the MINCLE-ligand glycolipids (TDM, TDB and the trehalose diester of C16) lead to a stronger enhancement of cellular protein content in the wells than the monoesters, and this difference was MINCLE-dependent (supplemental Fig. S7B). As an additional method to quantify effects of TDM on macrophage survival and proliferation, we employed the MTT conversion assay. First, the effect of TDM and TDB was comparable to the activity of the macrophage growth factor M-CSF (supplemental Fig. S7C). MTT conversion after 48 h was significantly increased by all glycolipids, but not by behenic acid, with the strongest effect observed for the MINCLE ligands (TDM, TDB and the trehalose diester of C16) (Fig. 8C). Similar to the changes in total protein content, the MTT conversion rates of the untreated macrophages decreased over time (see 24 h time point in supplemental Fig. S7D), suggesting that the glycolipids caused increased survival of macrophages through the MINCLE-independent upregulation of cell cycle-associated genes after 24 h and additional MINCLE-dependent effects at the later 48 h time point (Fig. 8B, 8C).

DISCUSSION

Recognition of the mycobacterial cord factor TDM by the CLR Mincle and the key role of SYK-CARD9-BCL10-MALT1 signaling has been firmly established by genetic deletion in mice. Compared with the well-studied TLR signaling pathways, an unbiased analysis of the signaling and transcriptome response to activation of MINCLE, or of related SYK-coupled CLR, is missing. This manuscript provides the first global assessment of macrophage activation by the mycobacterial cord factor TDM at the phosphoproteome level, combined with a corresponding RNAseq-based transcriptomic data set. Our results show substantial regulation of phosphopeptides in response to TDM, which was comparable in breadth with similar studies investigating the phosphoproteome of TLR-activated macrophages (21). We consider most of the changes in phosphoprotein abundance as a direct consequence of TDM-induced signaling, because at the early time point of 45 min after stimulation no secreted cytokines could be detected in the supernatant. Importantly, the comparison of WT and MINCLE-deficient macrophages in phosphoproteome and transcriptomic analysis allowed us to confirm the importance of the TDM-Mincle interaction, yet it also revealed an unexpectedly large fraction of MINCLE-independent changes, which are functionally separate from the Mincle-dependent responses.

MINCLE-dependent phosphorylation included several canonical players of this pathway, such as PKCΔ and PLCγ2, which was consistent with the enrichment of PKCΔ kinase motifs among the phosphopeptides induced by TDM in a MINCLE-dependent fashion. Because PKCΔ is required for formation of the CBM complex and subsequent activation of NF-kB (24), the strong enrichment of the pathway term “NF-kB activation” and the GO category “inflammatory response” among the cluster of MINCLE-dependent TDM target genes in our RNAseq data fits very well.

Several Src-family kinases were activated by TDM-MINCLE, including FYN and LYN. Of interest in this context, LYN was very recently reported to directly interact with MINCLE after TDM stimulation and to attenuate its signaling by recruitment of SIRP1α and SHP1 (58). The signaling via the Src kinase LYN was triggered by MINCLE-dependent activation of the integrin CD11b (58); thus, the detection of induced LYN phosphorylation in our data set may indicate integrin activation as a regulatory pathway of the response to TDM.

Triggering of PI3K-AKT signaling by mycobacterial cord factor or its synthetic analog has recently been described in human DC (51) and in human neutrophil–like HL-60 cells (59). Although we detected phosphorylation of several PI3K-AKT-associated proteins, including AKT1/2 and the regulatory PI3K subunit p85, the canonical phosphopeptides indicating activated AKT (Thr308 and Ser473) were not found in the phosphoproteome data set, but readily detected as upregulated by TDB/TDM using immunoblot analysis. Our further data on the role of the PI3K/AKT pathway in TDM-induced macrophage activation indicate that the PI3K catalytic subunit isoforms p110α and p110δ are redundant in macrophages, whereas pharmacological inhibition of p110α and p110δ was enough to completely block AKT phosphorylation and TDM-induced G-CSF and IL-6 production. Although the lack of a phenotype in p110α/δ-deficient macrophages was at first surprising, given reports that p110δ is the main isoform activated by tyrosine-kinase linked receptors in T cells and innate immune cells (60), these findings are concordant with previous reports showing redundancy between different p110 isoforms (61) and demonstrating a function for p110β in macrophages (62). The regulatory PI3K subunit p85α binds p110α, p110β and p110δ and therefore appears to play a broader role in PI3K activation (52). Indeed, our data obtained with p85α−/− macrophages showed a strong reduction of TDM-induced phosphorylation of AKT and of several cytokines (Fig. 5). Strongly attenuated phosphorylation of AKT in the absence of p85α was found previously in murine peritoneal macrophages stimulated with LPS (63) but has not been demonstrated after stimulation with a CLR ligand before. In contrast to earlier reports showing a selective reduction of the anti-inflammatory IL-10 and reciprocal overproduction of IL-12 by p85α−/− DC after stimulation with LPS (28), in our system both pro- and anti-inflammatory responses were similarly affected by p85α deficiency (and by pharmacological inhibition of PI3K catalytic subunits p110α and p110δ). Of note, Luyendyk et al. observed increased IL-6 and TNF production by p85α-deficient peritoneal macrophages stimulated with LPS (63). At present we do not know whether this discrepancy reflects a differen-
tial role of p85-PI3K signaling in macrophages versus DC, or rather points to differences in the regulation of TLR- and CLR-driven responses by PI3K.

The functional annotation of Mincle-dependently regulated phosphoproteins showed strong enrichment for endo-/lysosomal and transport proteins (Figs. 3, 4), an effect we had not observed in our analysis of LPS-induced phosphoproteins (21). The strong phosphorylation observed for many soluble carrier proteins may reflect ongoing signaling at the plasma membrane when macrophages are stimulated with plate-bound TDM as used here, whereas soluble ligands such as LPS could result in more rapid internalization and diversion of signalosomes to intracellular compartments. On the other hand, the strong Mincle-dependent phosphorylation of phagolysosomal proteins is of interest regarding the intracellular mycobacterial life-style. Several of these phosphoproteins control phagosomal maturation or trafficking processes, suggesting that TDM-Mincle signaling modulates the fate of the mycobacterial niche in macrophages. In fact, the proton pump TCRIRG1 (alias ATP6V0A3) is important for acidification of the mycobacterial phagolysosome (48, 64). The metallo-ductase STEAP3 controls iron homeostasis in macrophages (49) but has not been implicated in the mycobacteria-macrophage interaction yet. The endosomal GTPase RAB8A recruits PI3K to regulate the signaling from endosomal TLRs (50), which is of special interest regarding the activation of PI3K-AKT signaling by TDM discussed above.

The robust Mincle-independent kinase activation and transcriptional changes was a surprising result of our unbiased approach to macrophage reprogramming by the cord factor, given the strong Mincle-dependence of TDM-induced inflammatory responses and adjuvant activity reported by us and others before (8, 10, 65). The availability of synthetic mono- and diesters of trehalose with fatty acids of various length (C12, C16) allowed us to test whether the Mincle-independent responses to TDM may not be caused by a specific propensity of the cord factor itself, but rather by more generic features of glycolipids. The results demonstrated a clear separation in the capacity of the glycolipids to induce inflammatory genes (e.g., Csf3) associated with Mincle-dependent clusters 1 and 8 (Fig. 6), which required the diester form of C16, and the Mincle-independent induction of genes associated with cluster 4 and 6 (Fig. 6), which was triggered also by the trehalose monoesters (which do not bind to Mincle). This finding raises the question which alternative signaling mechanism(s) may be triggered by the cord factor and related glycolipids in macrophages. An obvious candidate is the close Mincle relative MCL, which was identified as second cord factor receptor following the detection of residual transcriptional responses in Mincle−/− mice (9) and can form heterodimers with Mincle (66, 67). Although we have not tested whether MCL is responsible for kinase activation and gene expression in the absence of Mincle in our system, we consider this unlikely for two reasons: first, Mincle-deficiency abrogates completely the activation of several canonical CLR pathway proteins (e.g., SYK, PKCζ; ERK1/2) which would be shared by MCL signaling; second, the Mincle-independent response appears to be functionally distinct based on the bioinformatic analysis of pathway and GO association of phosphoproteome and transcriptome changes. To confirm or reject this notion, Mincle-independent phosphoprotein regulation and TDM target gene expression needs to be tested in macrophages treated with SYK-inhibitors and in FcγR-deficient macrophages in future experiments. TLR/MYD88-dependent activation by the cord factor has been reported previously (12, 68), which could provide an alternative explanation for the Mincle-independent signaling. Although we have not tested this possibility by using Myd88−/− macrophages for analysis of Mincle-independent target gene expression, the finding that pathway enrichment associated MYD88 signaling with the Mincle-dependent response argues against this explanation. The scavenger receptors MARCO and MSR1 can also bind TDM (68); therefore, the phosphorylation of MSR1 prompted us to test its functional role in TDM-induced signaling and gene expression, but we did not observe differences in Msr1−/− macrophages. Finally, our attempts to assign a role in Mincle-independent target gene expression to two other candidate phosphoproteins, the complement receptor C5aR1 and the adapter protein FYB, were not successful. Activation of integrins by TDM may be another mechanism of Mincle-independent signaling and is supported by phosphorylation of the kinases HCK and PTK2, both involved in integrin-signaling (69, 70). Importantly, receptor-independent mechanisms such as membrane alterations induced by TDM and related glycolipids encountered by macrophages should also be considered as potential cause of the Mincle-independent phosphorylation events and transcriptional changes. Such direct interactions with membrane lipids and alterations in signaling molecule assembly have been described for urate acid crystals and aluminum hydroxide (71, 72).

Regardless of whether Mincle-independent phosphorylation and transcriptional regulation are induced by receptor-independent membrane alterations or triggered by a specific receptor of yet unknown identity, the type of biological processes and pathway terms associated with it were remarkably distinct. Especially prominent was the association of cell cycle-related genes and phosphoproteins that included regulators of the DNA-damage response like the kinase ATR. In turn, the only significantly enriched kinase motif identified in the Mincle-independent phosphosites was the ATM/ATR motif. The Mincle-independent upregulation of several cyclins and the pro-proliferative transcription factors MYC and of the kinase CHEK1, by TDM correlated with increased cell density in macrophage cultures. Our observation that macrophage survival/growth is induced Mincle-independently by several TDM-related glycolipids after 24h, but Mincle-dependently further enhanced by the cord factor after 48h, suggests that...
both pathways contribute to TDM-induced macrophage growth and survival. Phosphorylation of laminas, such as of LMNA observed here, contributes to regulation of the cell cycle (73). Thus, these data suggest that TDM promotes the survival and/or proliferation of macrophages and may thereby contribute to the dynamics of the granuloma response during mycobacterial infection. The concurrent activation of a DNA-damage response signature is at present unexplained but may be caused by the action of genotoxic substances like ROS and NO generated after stimulation with TDM, or could be a consequence of ATR activation because of replication stress. Recently, MYC-dependent DNA-damage response signaling was shown to be triggered in macrophages infected with M. bovis BCG, leading to the formation of multi-nucleated, polyploid macrophages through modified cell division (74). This pathway was MYD88-dependent and could be triggered by chronic stimulation with a bacterial lipopeptide TLR2 ligand (74). We have not determined whether the cell cycle/DNA-damage response observed here after TDM stimulation also leads to generation of multi-nucleated macrophages, but the similarity in the Myc-induction and ATM/ATR activity clearly suggests this possibility.

Taken together, we presented in this manuscript the first combined global analysis of the macrophage response to a C-type lectin receptor ligand at the phosphoproteome and transcriptome level. Our unbiased phosphoproteomic investigation of macrophage activation by the mycobacterial cord factor revealed many novel regulated phosphoproteins as components of MINCLE-dependent and -independent signaling, which appear to affect different biological processes and cellular components. The targeting of phagolysosomal and transport phosphoproteins by TDM-MINCLE signaling points to a role in the mycobacteria-macrophage interaction that merits further investigation. Likewise, the identification of robust AKT activation by TDM-MINCLE signaling and the essential role of p85α-PI3K in this process and its downstream effects indicate an important role of this pathway in macrophage reprogramming during mycobacterial infection. The strong MINCLE-independent component of the global phosphorylation and transcriptional response was consistently characterized by alterations in cell cycle regulators and DNA-damage response genes. Although it remains to be clarified by which receptor and mechanism this MINCLE-independent TDM effect is brought about, it appears to constitute a significant aspect of macrophage reprogramming by the cord factor and resembles recently reported features of mycobacteria-induced macrophage phenotypical changes.

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**DATA AVAILABILITY**

Mass spectrometric raw data and the MaxQuant outputs are available through the PRIDE repository ([https://www.ebi.ac.uk/pride/archive/](https://www.ebi.ac.uk/pride/archive/)) (44) and have been assigned the identifier PXD009513. Individual MS/MS spectra of phosphopeptides can be found here: [http://msviewer.ucsf.edu/prospector/cgi-bin/mssearch.cgi?report_title=MS-Viewer&search_key=dtfndgcpu&search_name=msviewer](http://msviewer.ucsf.edu/prospector/cgi-bin/mssearch.cgi?report_title=MS-Viewer&search_key=dtfndgcpu&search_name=msviewer). RNAseq data sets have been submitted to Gene Expression Omnibus and will be available under accession number GSE115322.

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Additional information is available under accession number GSE115322. RNAseq data sets have been submitted to Gene Expression Omnibus and will be available under accession number GSE115322.

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