- 1 G9a and Sirtuin6 epigenetically modulate host cholesterol accumulation to
- 2 facilitate mycobacterial survival
- 3 Praveen Prakhar<sup>a,1</sup>, Bharat Bhatt<sup>a,1</sup>, Tanushree Mukherjee<sup>a,1</sup>, Gaurav Kumar Lohia<sup>a</sup>,
- 4 Ullas Kolthur-Seetharam<sup>b</sup>, Nagalingam Ravi Sundaresan<sup>a</sup>, R.S. Rajmani<sup>c</sup>,
- 5 Kithiganahalli Narayanaswamy Balaji<sup>a,c,2</sup>
- <sup>a</sup> Department of Microbiology and Cell Biology, Indian Institute of Science, Bangalore
- 7 560012, Karnataka, India
- 8 b Department of Biological Sciences, Tata Institute of Fundamental Research,
- 9 Mumbai 400005, Maharashtra, India
- 10 <sup>c</sup> Centre for Infectious Disease Research, Indian Institute of Science, Bangalore -
- 11 560012, Karnataka, India
- 12 P.P., B.B. and T.M. contributed equally to this work.
- 13 <sup>2</sup> To whom correspondence may be addressed: Kithiganahalli Narayanaswamy
- 14 Balaji, Department of Microbiology and Cell Biology, Indian Institute of Science,
- 15 Bangalore 560012, Karnataka, India; Ph: +91-80-22933223; Email:
- 16 balaji@iisc.ac.in

24

25

- 18 **Keywords:** *Mycobacterium tuberculosis*, epigenetic modifications, G9a, SIRT6,
- 19 cholesterol accumulation
- 20 **Author contributions**: P.P. and K.N.B. conceived and designed the experiments.
- 21 P.P., T.M., B.B., G.K.L., and R.S.R. performed experiments. P.P., T.M., B.B., G.K.L.
- 22 and K.N.B. analyzed the data. U.K. and N.R.S. contributed reagents and materials.
- P.P., T.M., B.B., and K.N.B. wrote the manuscript and K.N.B. supervised the study.

Abstract:

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50

Cholesterol derived from the host milieu forms a critical factor for mycobacterial pathogenesis. However, the molecular circuitry co-opted by Mycobacterium tuberculosis (Mtb) to accumulate cholesterol in host cells remains obscure. Here, we report that a functional amalgamation of WNT-responsive histone modifiers G9a (H3K9 methyltransferase) and Sirt6 (H3K9 deacetylase) orchestrate cholesterol build-up in *in-vitro* and *in-vivo* models of Mtb infection. Mechanistically, G9a, along with SREBP2, drives the expression of cholesterol biosynthesis and uptake genes; while Sirt6 represses the genes involved in cholesterol efflux. The accumulated cholesterol promotes the expression of antioxidant genes leading to reduced oxidative stress, thereby supporting Mtb survival. In corroboration, loss-of-function of G9a in vitro and in vivo by pharmacological inhibition; or utilization of BMDMs derived from Sirt6 KO mice or in vivo infection in Sirt6 heterozygous mice; hampers host cholesterol accumulation and restricts Mtb burden. These findings shed light on the novel roles of G9a and Sirt6 during Mtb infection and highlight the previously unknown contribution of host cholesterol in potentiating anti-oxidative responses for aiding Mtb survival.

### Introduction:

*Mycobacterium tuberculosis* (Mtb) rewires host cellular machinery to subvert protective immune responses and achieve a secure and nutrient-rich niche. Emerging evidences highlight the implication of epigenetic factors in Mtb-driven tuning of gene expression to effectuate such immune evasion<sup>1–3</sup>. Reports suggest that the histone methyltransferase (HMT) EZH2 epigenetically down-modulates

52

53

54

55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

MHC-II presentation<sup>4</sup>; SET8 HMT governs immune processes such as apoptosis, oxidative stress and cytokine secretion<sup>5</sup>; while certain mycobacterial proteins themselves gain access to host chromatin and modulate a plenitude of immune genes<sup>6</sup>. One of the classical features of tuberculosis (TB) infection involves the accumulation of neutral lipids, cholesterol and cholesteryl esters to generate foamy macrophage (FM) phenotype<sup>7,8</sup>. Although, certain studies report that lipid droplets do not serve as an important source of nutrient for Mtb, hence, do not affect Mtb growth<sup>9</sup> and inhibition of fatty acid oxidation restricts intracellular growth of Mtb via ROS production<sup>10</sup>; there are evidences to support that the formation of FMs positively correlates with mycobacterial virulence and the loss of lipids from these cells compromises mycobacterial survival by not only limiting nutrients but also by curbing the requisite cues for altering hosts' ER stress, survival pathways and autophagy levels<sup>11–19</sup>. In this context, cholesterol serves essential functions for Mtb in the acquisition of dormancy and resistance to antibiotics in both, in vitro and in vivo systems<sup>14,20</sup>. To our interest, it has been recently reported that host cells such as macrophages form a major source of cholesterol for intracellular Mtb <sup>21</sup> and also possibly for extracellular Mtb released into the caseated or cavitated TB granuloma lesions<sup>14</sup>. However, information regarding the mechanisms regulating cholesterol accumulation in hosts during Mtb infection requires extensive investigation. Existing literature suggests that genes of cholesterol biosynthesis and homeostasis are epigenetically governed by miRNAs (miR-33a, miR-185), histone deacetylases (HDAC3, Sirt2, Sirt6) and HMTs (G9a) under distinct conditions<sup>22,23</sup>. Amongst these, the study on Sirt2 does not correlate its in vivo activity with chronic Mtb infection<sup>24</sup>. Contrastingly, activation of the nuclear Sirtuin, Sirt1, has recently been ascribed with restriction of Mtb growth by augmenting autophagy<sup>25</sup>. Sirt1 has also been shown to

77

78

79

80

81

82

83

84

85

86

87

88

89

90

91

92

93

94

95

96

97

98

99

100

be involved in lipid metabolism, stress response, anti-inflammatory response and cellular senescence in diverse contexts<sup>26–30</sup>. However, the contribution of the other nuclear Sirtuin, i.e. Sirt6, during infections, specifically mycobacterial infection, has not been addressed so far; although it has been enlisted as an upregulated gene in Mtb infection-related transcriptomic dataset<sup>31</sup>. Sirt6 majorly confers H3K9- and H3K56- deacetylation and is known to be associated with life span, genome stability and tumorigenesis<sup>32–35</sup>. Strikingly, Sirt6 has been identified as a potential regulator of SREBP1/2 functions, a major transcription factor for cholesterol metabolism<sup>36</sup>. We were piqued to unravel the epigenetic contribution of Sirt6 in accumulating cholesterol during TB infection. Additionally, evidences from the literature suggest that apart from acetylation, methylation of H3K9 (mono- and di-), conferred by G9a, imparts crucial epigenetic signatures for shaping immunological fates during various pathophysiological conditions, such as T cell differentiation, immunological memory, viral latency and endotoxin tolerance<sup>37-42</sup>. With this premise, we focused on elucidating the interplay of H3K9 methylation and acetylation by G9a and Sirt6, respectively, in defining cholesterol accumulation during TB infection. We find that Mtb induces the production of G9a and Sirt6, which contribute to epigenetically driven differential expression of cholesterol biosynthesis, uptake and efflux genes. thereby allowing cholesterol accumulation durina Mechanistically, WNT signaling was found to govern the levels of G9a and Sirt6 upon Mtb infection. WNT signaling has earlier been implicated in cell proliferation, migration, immunological processes and also in shaping immune responses during Mtb infection 43,44. Further, the accumulated cholesterol was found to aid mycobacterial survival by promoting anti-oxidative factors. Loss-of-function of G9a using a pharmacological inhibitor and that of Sirt6 using Sirt6 heterozygous mice in an *in vivo* mouse TB model was found to hamper host cholesterol accumulation and restrict Mtb burden. This was also corroborated by lung histopathology, which indicated a reduced severity of TB in mice lacking G9a and Sirt6 functions. Together, we show for the first time that G9a and Sirt6 are upregulated during Mtb infection; and in conjunction mediate TB pathogenesis by epigenetically reprogramming cholesterol accumulation. Besides, this study underscores the relevance of specific G9a and Sirt6 inhibitors as plausible anti-TB adjuvants.

#### **Results:**

## Interception of G9a and Sirt6 leads to restricted mycobacterial burden

Infection of host macrophages with pathogenic Mtb H37Rv was found to induce the expression of the HMT G9a (encoded by *Ehmt2*) at transcript as well as protein level; without any significant change in the global H3K9 monomethylation pattern (Fig. 1A; S1A, B). Alongside, we report that Mtb H37Rv augments the expression of the HDAC Sirt6. However, the corresponding global H3K9 acetylation mark also remains unaltered (Fig. 1A; S1A, B). Importantly, we corroborated our findings in human primary macrophages (Fig. 1B; S1C) as well as in a mouse model of pulmonary TB infection (Fig. 1C, D; S1D), wherein, we observed an enhanced expression of the concerned epigenetic factors at the transcript and protein level during Mtb H37Rv infection. Further, the induction of G9a and Sirt6 was found to be specific to virulent species of mycobacterium as infection of mouse peritoneal macrophages with the non-pathogenic *Mycobacterium smegmatis* showed only a weak expression of G9a and Sirt6 (Fig. S1E). To evaluate the possible contribution of these epigenetic factors to Mtb infection, *in vitro* CFU assays were performed. Inhibition of G9a using a specific pharmacological inhibitor (BIX-01294) was found to

compromise Mtb H37Rv burden after 48 h of *in vitro* infection (**Fig. 1E**). Next, mycobacterial CFU was assessed in BMDMs derived from WT (littermate control) and *Sirt6* knockout (KO) mice as the premature ageing and death of Sirt6 KO mice by 4 weeks post-natally<sup>45</sup> hinders the isolation of thioglycolate-elicited peritoneal macrophages and long-term *in vivo* experiments. Sirt6 expression was assessed in the lung homogenate of WT, Sirt6 heterozygous and Sirt6 KO mice (**Fig S1F**). We found the Mtb H37Rv burden to be restricted in *Sirt6* KO BMDMs (**Fig. 1F**). Further, *in vitro* silencing of *Ehmt2* or *Sirt6* individually lead to a compromised mycobacterial CFU, that was further diminished in mouse peritoneal macrophages knocked down for both *Ehmt2* and *Sirt6* (**Fig. 1G**; **Fig. S1G**: knockdown validation; **Fig. S1H**: cell viability assessment). These results suggest a critical role for the epigenetic modifiers G9a and Sirt6 in the successful pathogenesis of mycobacterium.

# G9a and Sirt6 effectuate cholesterol accumulation during mycobacterial pathogenesis

As described earlier, among various factors, host-derived cholesterol forms an integral part of mycobacterial pathogenesis *in vitro* and *in vivo*. In this context, virulent Mtb H37Rv infection was found to trigger cholesterol accumulation in host macrophages, unlike *M. smegmatis* infection, as assessed by Filipin staining (**Fig. S2A, B**). The same was mirrored in the lungs of mice infected with Mtb H37Rv, wherein staining lung cryosections with Filipin showed a significant increase in cholesterol accumulation specifically in macrophages (**Fig. S2C, D**). With the premise that both G9a and Sirt6 have been reported to epigenetically regulate cholesterol levels<sup>22</sup>, we sought to explore their contribution to cholesterol accumulation in the context of Mtb infection. It was observed that *in vitro* silencing of

Ehmt2 and Sirt6 via specific siRNAs led to a marked decline in the ability of Mtb H37Rv to furnish cholesterol accretion (**Fig. 2A, B**). Further, significantly less cholesterol was detected by Filipin staining in BMDMs derived from Sirt6 KO mice, even in the presence of Mtb H37Rv infection (**Fig. 2C, D**). Substantiating the same, macrophage-specific accumulation of cholesterol was reduced in the lungs of G9a inhibitor-treated mice (**Fig. 2E, F**) and Sirt6 heterozygous mice (**Fig. 2G, H**). These evidences strongly indicate the ability of Mtb to utilize G9a and Sirt6 for mediating the essential process of cholesterol accumulation in host cells.

## Cholesterol biosynthesis and transport genes are differentially regulated by

## G9a and Sirt6

The accumulation of cholesterol in a given cell or tissue results from the coordinated interplay of genes involved in its biosynthesis and uptake on one hand, and its efflux on the other. To this end, the status of the pertinent markers (23 genes) of each function was assessed for their transcript level expression during infection with Mtb H37Rv *in vitro* and *in vivo* (**Fig. S2E, F**). We observed that genes involved in cholesterol uptake (*Lrp2*) and biosynthesis (*Aacs, Hmgcs1, Mvd, Dhcr24*) were significantly upregulated during infection; while those implicated in efflux (*Abca1, Abcg1*) showed a marked downregulation. We further assessed the transcript levels of the altered genes in human PBMCs infected with Mtb H37Rv. Corroborating our *in vitro* and *in vivo* mice data, we observed a significant increase in the transcript levels of genes involved in cholesterol uptake (*Lrp2*) and biosynthesis (*Aacs, Hmgcs1, Mvd, Dhcr24*) with a downregulation in the expression of efflux genes (*Abca1, Abcg1*) in H37Rv-infected human PBMCs (**Fig. S2G**). Interestingly, this differential gene expression was found to be finely tuned by the combined activities of G9a and

Sirt6. While depletion of G9a function by siRNA-mediated knock-down compromised the expression of the biosynthesis and uptake genes (*Lrp2, Aacs, Hmgcs, Mvd, Dhcr24*) at the transcript level (**Fig. 3A**); that of Sirt6 rescued the Mtb-dependent downregulation of cholesterol efflux genes (*Abca1, Abcg1*) (**Fig. 3B**). Inline, overexpressing Sirt6 led to a marked decrease in *Abca1* and *Abcg1* expression (**Fig. S3A**). The transcript level profiling performed for the concerned genes in the lungs of mice treated with G9a inhibitor *in vivo* or *Sirt6* heterozygous mice also yielded a similar pattern (**Fig. 3C, D**). These findings were validated at the protein level, where G9a inhibitor limited the surface expression of LRP2 during infection, both *in vitro* and *in vivo* (**Fig. 3E, F**); and ABCA1 protein expression was found to be elevated in the lungs of Mtb H37Rv-infected *Sirt6* heterozygous mice (**Fig. 3G**), compared to that in the infected wild type controls. The protein level of ABCA1 was also rescued in macrophages knocked down for Sirt6 (**Fig. S3B**). The current set of observations prompted us to delineate the G9a- and Sirt6-driven mechanism of differential regulation of cholesterol biosynthesis, uptake and efflux genes.

G9a-SREBP2 and Sirt6 fine-tune cholesterol accumulation during Mtb infection
The transcription factor SREBP2 (encoded by *Srebf2*) is a well-established master
regulator of cholesterol biosynthesis genes. However, it functions in tight association
with accessory transcription activators and regulators<sup>46</sup>. We hypothesized the
possibility of SREBP2 and G9a to interact and together bring about the augmented
expression of cholesterol biosynthesis and uptake genes. In this regard, first we
show that Mtb H37Rv induces SREBP2 expression and immunopulldown analysis
indicates that SREBP2 interacts with G9a during Mtb H37Rv infection (Fig. 4A).
Loss-of-function of SREBP2 using specific siRNA (Fig. S3C, knockdown validation)

202

203

204

205

206

207

208

209

210

211

212

213

214

215

216

217

218

219

220

221

222

223

224

225

compromised the expression of positive factors of cholesterol accumulation (Lrp2, Aacs, Hmgcs, Mvd, Dhcr24) (Fig. 4B). In line with this, Mtb H37Rv burden was significantly restricted in macrophages knocked down for Srebf2 in vitro (Fig. 4C). Next, we verified the role of G9a HMT at the chromatin level by assessing the recruitment of G9a to SREBP2 binding sites at the promoters of cholesterol biosynthesis and uptake genes. Both, G9a occupancy and associated H3K9me1 marks were found to be elevated during Mtb H37Rv infection (Fig. 4D, E). Further, sequential ChIP analysis showed an enhanced co-occupancy of the concerned promoters with G9a and SREBP2 (Fig. 4F), thus confirming the concerted action of SREBP2 and G9a in driving the expression of cholesterol biosynthesis and uptake genes (Lrp2, Aacs, Hmgcs, Mvd, Dhcr24). Contrary to this, Sirt6 was found to occupy the promoters of Abca1 and Abcg1 during Mtb H37Rv infection, leading to concomitantly decreased acetylation marks; and supporting the initially observed downregulation of the cholesterol efflux genes during Mtb infection (Fig. 4G, H). Further, since H3K9me2 conferred by G9a renders a closed chromatin state and transcriptional downregulation, the contribution of G9a in the reduced expression of ABCA1 was analyzed. It was found that mycobacteria lost the ability to downregulate ABCA1 in G9a knocked down macrophages (Fig. S3D, E); indicating the partial dependence of cholesterol efflux genes on the repressive function of G9a. To understand this interplay of G9a and Sirt6, a time kinetics ChIP assay was performed. Interestingly, it was observed that the temporal recruitment of Sirt6 and G9a allows the repression of cholesterol efflux genes by both Sirt6 and G9a at early time points (12 h) (Fig. S4A, B), that is then sustained only by G9a at later times of infection (24 h) (Fig. S4C). However, unlike for the biosynthesis/uptake genes, SREBP2 was not found to be involved in the regulation of the efflux genes (Fig.

**S4D**). Also, the cholesterol biosynthesis and uptake genes (*Lrp2, Aacs, Hmgcs, Mvd, Dhcr24*) were only regulated by G9a and were found to be independent of Sirt6

(Fig. S4E-G).

## Cholesterol accumulation mitigates oxidative stress during mycobacterial

infection

226

227

228

229

230

231

232

233

234

235

236

237

238

239

240

241

242

243

244

245

246

247

248

249

250

The orchestrated accumulation of cholesterol by Mtb-induced G9a and Sirt6 provides insights into the crucial functions that cholesterol might effectuate to favor Mtb survival. As discussed, the contribution of cholesterol as a source of nutrition for mycobacteria is widely accepted. However, evidences from the literature suggest numerous alternate implications of cholesterol in cellular homeostasis and responses to stimuli<sup>47,48</sup>. Moreover, supplementation of exogenous cholesterol helped in mitigating the toxic effects of bile acid and lipids in Nonalcoholic steatohepatitis by enhancing the expression of NRF2 and HIF- $1\alpha^{49}$ . Similarly, Cholesterol crystals present in the atherosclerotic plagues also act as a stimulus to regulate Nrf-2<sup>50</sup>. We report that mycobacteria-induced cholesterol accumulation renders the expression of a principal transcription activator of antioxidant genes, NRF2 (encoded by Nfe2l2) that then leads to the expression of its target genes (Fig. S5A-D). In this line, perturbation of G9a or Sirt6 using specific siRNAs compromised the expression of NRF2-target genes Trxrd1, Ngo1, Hmox1, Gsr, Gpx1, Sod1 at the transcript level (Fig. S5E) and protein level (Fig. 5A). We next verified that the observed loss of antioxidant gene expression indeed resulted from hampered accumulation of cholesterol in G9a- and Sirt6-knocked down cells. To this end, first we utilized siRNAs against the five G9a-dependent genes found to be essential for cholesterol biosynthesis and uptake (Lrp2, Aacs, Hmgcs, Mvd, Dhcr24); and verified the

252

253

254

255

256

257

258

259

260

261

262

263

264

265

266

267

268

269

270

271

272

273

274

275

abrogation of cholesterol accumulation during Mtb H37Rv infection to result from the specific downregulation of the five concerned targets (Fig. S6A-C), while negating the implication of any other Mtb-unresponsive cholesterol biosynthesis genes in the process (Fig. S6D). In these cholesterol deficient cells, we found a significant reduction in the expression of antioxidant genes at the transcript and protein level (Fig. S5E, 5B); implicating cholesterol in driving antioxidant gene expression. Consequently, increased oxidative stress was observed in G9a and Sirt6 knocked down cells (Fig 5C-D) as well as in cells deficient of G9a-dependent cholesterol biosynthesis and uptake gene expression (Fig. 5E-F). Corollary to this, we also observed that in vitro depletion of cholesterol accumulation genes (Fig. 5G) or Nfe2l2 (Fig. 5H) in macrophages significantly compromised Mtb H37Rv burden (Fig. **5G**, **H**). Furthermore, to identify the importance of each of the G9a-dependent cholesterol biosynthesis and uptake genes, each gene was individually knocked down and assessed for their effect on the expression of antioxidant genes (Fig. S7) as well as Mtb burden (Fig. 51). Our observations suggest a dominant role for the genes *Hmgcs1* and *Aacs*, catalyzing the pioneer steps of cholesterol biosynthesis, in impacting oxidative stress and mycobacterial burden. These evidences highlight the critical functions of cholesterol accumulation in mycobacteria-infected hosts. However, further investigation into the exact implication of *Hmgcs1* and *Aacs* in regulating antioxidant gene expression is warranted. In the perspective of the above-mentioned observations, we explored the role for WNT signaling pathway during Mtb-driven expression of G9a and Sirt6. WNT pathway is known to modulate various cellular events like autophagy during Mtb

infection<sup>51</sup>. Importantly, it has been associated with antioxidants such as NRF2 for

defining neuronal developmental pathways<sup>52</sup>. Activated WNT signaling, driven by

277

278

279

280

281

282

283

284

285

286

287

288

289

290

291

292

293

294

295

296

297

298

299

300

WNT3A, has also been shown to enhance NRF2-mediated anti-oxidant gene expression by preventing the GSK3β-dependent phosphorylation and subsequent proteasomal degradation of NRF2 in hepatocytes<sup>53</sup>. Further, its contribution in regulating lipid accumulation by endocytosis of LDL-derived cholesterol <sup>54</sup> indicated its possible role in yet another aspect of Mtb infection, i.e. cholesterol accumulation. G9a and Sirt6 expression was found to be dependent on Mtb H37Rv-activated WNT pathway (Fig. S8A-C; S8A: hallmarks of Mtb-activated WNT signaling: increased pGSK3\beta and reduced p\beta-CATENIN); as inhibition of the pathway with pharmacological inhibitors (IWP2 and β-CATENIN inhibitor) (**Fig. S8C. right panel**) or knockdown of *Ctnbb1* (**Fig. S8C**, **middle panel**; **Fig. S8B**: knockdown validation) compromised the levels of G9a and Sirt6. Conversely, β-CATENIN over-expression alone induced the expression of the concerned histone modifiers (Fig. S8C, left panel). Further, β-CATENIN was found to be recruited to the promoters of *Ehmt2* and Sirt6, (Fig. S8D). siRNA-mediated knockdown of Ctnbb1 compromised the ability of Mtb to differentially regulate cholesterol metabolism genes (Fig. S8G); subsequent cholesterol accumulation (Fig. S8E, F) and hence Mtb H37Rv survival (Fig. S8H). These findings indicate that Mtb infection leads to the WNT signaling pathway-dependent expression of G9a/Sirt6 as well as accumulation of cholesterol, which drives a secure niche for the pathogen to survive.

## G9a and Sirt6 contribute to mycobacterial pathogenesis

The observed G9a/Sirt6-dependent accumulation of cholesterol and the related abatement of mycobacterial burden upon their functional loss incited us to determine the impact of G9a and Sirt6 in defining *in vivo* Mtb burden and associated lung tissue pathology during Mtb H37Rv infection. We found that therapeutic treatment of Mtb

H37Rv-infected mice with G9a inhibitor not only compromised cholesterol accumulation but also reduced mycobacterial CFU and led to a decreased level of Mtb infection-specific granulomatous lesions. Lung histopathological examination by Hematoxylin and Eosin (H and E) staining also revealed a marked reduction in the percentage of lung area covered with the characteristic TB granulomatous lesions, with an overall decline in total granuloma score compared to the untreated counterparts (Fig. 6A-E). Further, we observed limited Mtb H37Rv CFU in the lungs and spleen in *Sirt6* heterozygous mice and up to 50% restriction in the ability of *Sirt6* heterozygous mice to effectively develop TB granulomatous lesion (Fig. 6F-J). Therefore, the partial normalization of total lung architecture in mice lacking G9a or Sirt6 functions strongly indicates the relevance of the histone modifications conferred by G9a and Sirt6 in the pathogenesis of TB. We believe that thwarted cholesterol accumulation, leading to enhanced oxidative stress, jeopardizes mycobacterial survival strategies, thereby restricting overall TB progression in mice with abrogated G9a/Sirt6 functions.

## **Discussion**

The formation of FMs has been described as an integral part of TB pathogenesis and the constituents of the lipid droplets (LDs) contained in the FMs associate with diverse functions. Specifically, cholesterol uptake by Mtb and utilization to achieve survival advantages has been vividly elucidated<sup>55–58</sup>. We uncover the Mtb-driven host molecular players that lead to the accumulation of this essential factor in host cells during infection. Despite the presence of compelling evidences for the implication of cholesterol in the pathogenesis of Mtb, the epidemiological surveys depict a nonlinear and complex relationship between high cholesterol and TB

327

328

329

330

331

332

333

334

335

336

337

338

339

340

341

342

343

344

345

346

347

348

349

350

progression<sup>59</sup>. Similarly, in mouse models of pulmonary TB, a cholesterol-rich diet (high serum cholesterol levels) has been related to distinct disease outcomes. For instance, in ApoE KO mice, high serum cholesterol impairs host defense against Mtb<sup>60</sup>; while that in *Ldlr* KO mice does not alter the capacity of the host to restrict mycobacterial replication<sup>61</sup>. These uncertainties may be explained by the differences in cholesterol availability that arise from its esterification or association with lipoproteins to form VLDLs, LDLs and HDLs. Therefore, a clear picture defining the role of cholesterol still warrants investigation. The accumulation of cholesterol imparts regulatory effects on several aspects of host immunity by altering processes ranging from plasma membrane dynamics to maintaining serum cholesterol levels and epigenetic deregulations. Cholesterol is important for the adaptive immune system for its contribution to the formation of plasma membrane lipid rafts, which facilitate immune functions such as T-cell and Bcell signaling, their activation and proliferation 62,63. Further, high serum cholesterol leads to autoimmune and inflammatory manifestations via aberrant immune activation<sup>64</sup>. Alongside these important roles, cholesterol accumulation also shapes the innate immune arm by modulating functions such as TLR signaling, monocyte proliferation, macrophage polarization, apoptosis as well as dendritic cell maturation and activation under distinct conditions<sup>65-69</sup>, including infections. For instance, cholesterol has been shown to play a crucial role in regulating Salmonella-induced autophagy 70 and lowering free cholesterol by their conversion to oxysterols has been implicated in providing immunity against *Listeria monocytogenes* infection 1. During mycobacterial infection, in particular, suppression of intracellular cholesterol accumulation via oxysterols (natural LXR activators) or by inhibition of SREBP2 has been shown to enhance the production of anti-microbial peptides and restrict Mtb

352

353

354

355

356

357

358

359

360

361

362

363

364

365

366

367

368

369

370

371

372

373

374

375

burden<sup>72</sup>. Inline, loss of function of LXR $\alpha$  and LXR $\beta$  (leading to reduced expression of Abca1) has been reported to render mice more susceptible to Mtb infection due to defective recruitment of innate effector cells and innate immune functions as well as severely compromised Th1/Th17 functions<sup>73</sup>. In the current study, we present a novel mechanism by which free cholesterol accumulated within host cells can aid in Mtb pathogenesis. We show that G9a-SREBP2 and SIRT6 independently regulates the cholesterol homeostasis wherein G9a-SREBP2 regulates cholesterol biosynthesis genes, whereas, SIRT6 regulates the expression of cholesterol efflux genes but has no effect on cholesterol biosynthesis genes. We find that cholesterol accumulation modulates the innate immune arm by driving the expression of antioxidative genes that would favor Mtb survival by circumventing oxidative stress responses and mediators. This aligns with the observation that cells with high cholesterol upregulate anti-oxidants such as NRF2 and HO-1 to mitigate oxidative stress<sup>74</sup>. A recent report from our lab proposes that mycobacterial clearance pathways such as apoptosis and pro-inflammatory cytokine production are hampered by classical anti-oxidative molecules TRXR1 and NQO1<sup>5</sup>. Therefore, cholesterol-dependent antioxidant production and subsequent innate and adaptive immune alterations not reported as yet, can potentially help in strengthening the understanding of the survival strategies employed by Mtb. In the light of host-directed therapeutics, our finding is in congruence with a previous study where statins, that decrease cholesterol levels by inhibiting HMGCoA reductase (a rate-limiting step of cholesterol biosynthesis), had been reported to inhibit mycobacterial growth<sup>75</sup>. With the individual knockdown of G9a-dependent cholesterol biosynthesis genes, we tease out the specific contribution of *Hmgcs1* and Aacs in regulating cholesterol-driven mitigation of oxidative stress and subsequently,

mycobacterial burden. Therefore, this study provides an avenue for testing alternate targets for effective combinatorial therapy against TB and for dedicated studies on metabolic homeostasis and mycobacterial pathogenesis in *Hmgcs1* or *Aacs* knockout conditions. Recently, mammalian sirtuins have been proposed as a potential target for host-directed therapy against TB. For example, SIRT1 activators ameliorates lung pathology, SIRT3 promotes antimycobacterial responses whereas SIRT2 inhibition has been shown to reduce Mtb burden <sup>25,76,77</sup>. In the current study, we find that SIRT6 benefits the Mtb survival and impacts lung pathology, thereby establishing the class of sirtuins as potential targets for TB therapeutics. Together, we report that epigenetic modifiers G9a and Sirt6 are induced by Mtb, and the two enzymes differentially occupy the promoters of distinct arms of cholesterol biosynthesis, uptake and efflux genes, in order to build up cholesterol within host cells. Interception of G9a and Sirt6 restricts mycobacterial burden and limits TB pathology, plausibly by compromising free cholesterol accumulation and thereby increasing oxidative stress in host cells (Fig. 7). We believe that an organ-specific and carefully titrated delivery of therapeutics against these epigenetic factors would provide rational and clinically relevant adjuvants for TB treatment.

#### **Materials and Methods**

### **Mice and Cells**

376

377

378

379

380

381

382

383

384

385

386

387

388

389

390

391

392

393

394

395

396

397

398

399

400

Male and female mice of the following strains were utilized in the study: BALB/c (stock number 000651, The Jackson Laboratory, USA), *Sirt6* KO (kind gift from Dr. Ullas Kolthur-Seetharam, TIFR, India and Dr. Nagalingam Ravi Sundaresan, IISc, India; primary source: The Jackson Laboratory, USA, stock number 006050) and *Sirt6* heterozygous (kind gift from Dr. Ullas Kolthur-Seetharam,

TIFR, India and Dr. Nagalingam Ravi Sundaresan, IISc, India; primarily generated by crossing *Sirt6* KO mice with WT 129S6 mice). Mouse primary macrophages were isolated from peritoneal exudates using ice-cold PBS four days post intraperitoneal injection of 1.5ml of brewer thioglycollate (8%). RAW 264.7 mouse macrophages cell line was obtained from National Center for Cell Sciences, Pune, India; and used for transient transfection experiments using plasmids as they are better suited for transfection as compared to the peritoneal macrophages that are known to be highly sensitive to external DNA<sup>78,79</sup>. Primary macrophage and RAW 264.7 macrophage cell line was cultured in Dulbecco's Minimal Eagle Medium (Gibco, Thermo Fisher Scientific) supplemented with 10% heat-inactivated Fetal Bovine Serum (Gibco, Thermo Fisher Scientific) and maintained at 37°C in 5% CO<sub>2</sub> incubator. All strains of mice were obtained from The Jackson Laboratory and maintained in the Central Animal Facility (CAF), Indian Institute of Science (IISc) under 12 h light and dark cycle.

#### **Ethics Statement**

Experiments involving mice and virulent mycobacteria (Mtb H37Rv) were carried out after the approval from Institutional Ethics Committee for animal experimentation and Institutional Biosafety Committee. The animal care and use protocol adhered were approved by national guidelines of the Committee for the Purpose of Control and Supervision of Experiments on Animals (CPCSEA), Government of India.

#### **Bacteria**

Mtb H37Rv was a kind research gift from Prof. Kanury Venkata Subba Rao,

THSTI, India. tdTomato Mtb H37Rv was a kind research gift from Dr. Amit Singh, IISc, India. Mycobacterial cultures were grown to mid-log phase in Middlebrook 7H9 medium (Difco, USA) supplemented with 10% OADC (oleic acid, albumin, dextrose, catalase) and hygromycin for tdTomato Mtb H37Rv. Single-cell suspensions of mycobacteria were obtained by passing mid-log phase culture through 23, 28 and 30 gauge needle 10 times each and used at a multiplicity of infection 10 unless mentioned otherwise. The studies involving virulent mycobacterial strains were carried out at the biosafety level 3 (BSL-3) facility at CIDR, IISc.

## Reagents and antibodies

All general laboratory chemicals were obtained from Sigma Aldrich/Merck Millipore, Thermo Fisher Scientific, HiMedia or Promega. Tissue culture plasticware was purchased from Jet Biofil or Tarsons Products Pvt Ltd. Further details are provided in the supplementary file.

## **Transient transfection studies**

RAW 264.7 macrophages were transiently transfected with 5 $\mu$ g of overexpression constructs of  $\beta$ -CATENIN and SIRT6; or peritoneal macrophages were transfected with 100 nM each of siGLO Lamin A/C, non-targeting siRNA or specific siRNAs against *Ehmt2*, *Sirt6*, *Ctnnb1*, *Lrp2*, *Aacs*, *Hmgcs1*, *Mvd*, *Dhcr24*, *Srebf2*, *Nfe2I2* (purchased from Dharmacon as siGENOMETM SMARTpool reagents) with polyethyleneimine. 70-80% transfection efficiency was observed by counting the number of siGLO Lamin A/C positive cells in a microscopic field using fluorescence microscopy. 36 h post-transfection (for experiments with RAW 264.7 cells) or 24h post-transfection (for experiments with peritoneal macrophages), the cells were treated or infected as indicated and processed for analyses.

#### *In vivo* mouse model and inhibitor treatment

451

452

453

454

455

456

457

458

459

460

461

462

463

464

465

466

467

468

469

470

471

472

473

474

475

BALB/c mice (n=40) were infected with mid-log phase Mtb H37Rv, using a Madison chamber aerosol generation instrument calibrated to 200 CFU/animal. Aerosolized animals were maintained in a securely commissioned BSL3 facility. Post 28 days of established infection, mice were administered a daily dose of G9a inhibitor BIX-01294 (40mg/kg) 80 intra-peritoneally for 28 days. Alternately, wild type (littermate control) mice or Sirt6 heterozygous mice were infected as described above. In each case, on the 56<sup>th</sup> day, mice were sacrificed, spleen and left lung lobe and spleen were homogenized in sterile PBS, serially diluted and plated on 7H11 agar containing OADC to quantify CFU. Upper right lung lobes were fixed in formalin, embedded in paraffin and stained with hematoxylin and eosin and immunofluorescence analysis. For Granuloma scoring, different scores were assigned based on the characteristic granulomatous features that is granuloma with necrosis (Score=5), without necrosis (Score = 2.5) and with fibrosis (Score = 1)81. For total granuloma scoring, the number of granulomas in each lung lobe was multiplied with the characterized feature score. The granulomatous area of lung sections stained with H&E was measured using Image J software (granulomatous area/ total area \*100).

## Statistical analysis

Levels of significance for comparison between samples were determined by the Student's t-test and one-way ANOVA followed by Tukey's multiple-comparisons. The data in the graphs are expressed as the mean ± SEM for the values from at least 3 or more independent experiments and P values < 0.05 were defined as

477

478

479

480

481

482

483

484

485

486

487

488

489

490

491

492

493

494

495

496

497

498

499

significant. GraphPad Prism 6.0 software (GraphPad Software, USA) was used for all the statistical analyses. All the details concerning the pharmacological reagents, antibodies, in vitro experiments and procedures have been provided as supplementary information. Acknowledgments: We thank CAF, IISc for providing mice for experimentation. β-CATENIN cDNA was gifted by Dr. Roel Nusse, Stanford University School of Medicine, USA. We acknowledge the BSL-3 facility for allowing the experiments on Mtb H37Rv to be carried out. **Funding**: This work was supported by funds from the Department of Biotechnology (DBT No. BT/PR27352/BRB/10/1639/2017, DT.30/8/2018 and BT/PR13522/COE/34/27/2015, DT.22/8/2017 to K.N.B) and the Department of Science and Technology (DST, EMR/2014/000875, DT.4/12/15 to K.N.B.), New Delhi, India. K.N.B. thanks Science and Engineering Research Board (SERB), DST, for the award of J. C. Bose National Fellowship (No.SB/S2/JCB-025/2016, DT.25/7/15) and for the funding (SP/DSTO-19-0176, DT.06/02/2020). The authors thank DST-FIST, UGC Centre for Advanced Study and DBT-IISc Partnership Program (Phase-II at IISc BT/PR27952/INF/22/212/2018) for the funding and infrastructure support. Fellowships were received from IISc (P.P., T.M., and G.K.L.) and UGC (B.B.). **Competing interests**: No competing financial interests.

Abbreviations: HMT, histone methyl transferase; TB, tuberculosis; FM, foamy macrophage; ER, endoplasmic reticulum; HDAC, histone deacetylase; CFU, colony forming unit; BMDM, bone marrow derived macrophage; KO, knock out; LRP, Low density lipoprotein receptor-related protein 2; SREBP, Sterol response element binding protein; ABC, ATP-binding cassette; LDL, low density lipoprotein; LD, lipid droplet; HDL, high density lipoprotein; NRF2, nuclear factor erythroid 2 (NFE2) related factor 2; VLDL, very low density lipoprotein; PBMC, polymorphic blood mononuclear cells.

#### References:

500

501

502

503

504

505

506

507

508

509

510

- 511 1. Cole, J., Morris, P., Dickman, M. J. & Dockrell, D. H. The therapeutic potential of epigenetic manipulation during infectious diseases. *Pharmacol. Ther.* **167**, 85–99 (2016).
- 514 2. Esterhuyse, M. M., Linhart, H. G. & Kaufmann, S. H. E. Can the battle against tuberculosis gain from epigenetic research? *Trends Microbiol.* **20**, 220–226 (2012).
- 517 3. Kathirvel, M. & Mahadevan, S. The role of epigenetics in tuberculosis infection. 518 Epigenomics vol. 8 537–549 (2016).
- 519 4. Ghorpade, D. S., Holla, S., Sinha, A. Y., Alagesan, S. K. & Balaji, K. N. Nitric 520 oxide and KLF4 protein epigenetically modify class II transactivator to repress 521 major histocompatibility complex II expression during Mycobacterium bovis 522 bacillus Calmette-Guérin infection. *J. Biol. Chem.* **288**, 20592–20606 (2013).
- 523 5. Singh, V. *et al.* Histone methyltransferase SET8 epigenetically reprograms host immune responses to assist mycobacterial survival. *J. Infect. Dis.* **216**, 477–488 (2017).
- 526 6. Yaseen, I., Kaur, P., Nandicoori, V. K. & Khosla, S. Mycobacteria modulate 527 host epigenetic machinery by Rv1988 methylation of a non-tail arginine of 528 histone H3. *Nat. Commun.* **6**, 1–13 (2015).
- 7. Holla, S. *et al.* MUSASHI-Mediated Expression of JMJD3, a H3K27me3 Demethylase, Is Involved in Foamy Macrophage Generation during Mycobacterial Infection. *PLOS Pathog.* **12**, e1005814 (2016).
- 8. Russell, D. G., Cardona, P. J., Kim, M. J., Allain, S. & Altare, F. Foamy macrophages and the progression of the human tuberculosis granuloma. *Nature Immunology* vol. 10 943–948 (2009).

- 535 9. Knight, M., Braverman, J., Asfaha, K., Gronert, K. & Stanley, S. Lipid droplet formation in Mycobacterium tuberculosis infected macrophages requires IFN-537 y/HIF-1α signaling and supports host defense. *PLoS Pathog.* **14**, (2018).
- 538 10. Chandra, P. *et al.* Inhibition of fatty acid oxidation promotes macrophage control of mycobacterium tuberculosis. *MBio* **11**, 1–15 (2020).
- 540 11. Daniel, J., Maamar, H., Deb, C., Sirakova, T. D. & Kolattukudy, P. E.
  541 Mycobacterium tuberculosis uses host triacylglycerol to accumulate lipid
  542 droplets and acquires a dormancy-like phenotype in lipid-loaded macrophages.
  543 *PLoS Pathog.* **7**, (2011).
- 544 12. D'Avila, H. *et al.* Mycobacterium bovis Bacillus Calmette-Guérin Induces 545 TLR2-Mediated Formation of Lipid Bodies: Intracellular Domains for 546 Eicosanoid Synthesis In Vivo . *J. Immunol.* **176**, 3087–3097 (2006).
- Dodd, C. E., Pyle, C. J., Glowinski, R., Rajaram, M. V. S. & Schlesinger, L. S.
   CD36-Mediated Uptake of Surfactant Lipids by Human Macrophages
   Promotes Intracellular Growth of Mycobacterium tuberculosis . *J. Immunol.* 197, 4727–4735 (2016).
- 551 14. Kim, M. J. *et al.* Caseation of human tuberculosis granulomas correlates with elevated host lipid metabolism. *EMBO Mol. Med.* **2**, 258–274 (2010).
- 553 15. Peyron, P. *et al.* Foamy macrophages from tuberculous patients' granulomas constitute a nutrient-rich reservoir for M. tuberculosis persistence. *PLoS Pathog.* **4**, (2008).
- 556 16. Singh, V. *et al.* Mycobacterium tuberculosis-driven targeted recalibration of macrophage lipid homeostasis promotes the foamy phenotype. *Cell Host Microbe* **12**, 669–681 (2012).
- 559 17. Brzostek, A., Pawelczyk, J., Rumijowska-Galewicz, A., Dziadek, B. & Dziadek, 560 J. Mycobacterium tuberculosis is able to accumulate and utilize cholesterol. *J. Bacteriol.* **191**, 6584–6591 (2009).
- 562 18. Ouimet, M. *et al.* Mycobacterium tuberculosis induces the MIR-33 locus to 563 reprogram autophagy and host lipid metabolism. *Nat. Immunol.* **17**, 677–686 564 (2016).
- 565 19. Kim, Y. S. *et al.* PPAR-alphamActivation Mediates Innate Host Defense 566 through Induction of TFEB and Lipid Catabolism. *J. Immunol.* **198**, 3283–3295 567 (2017).
- 568 20. Gatfield, J. & Pieters, J. Essential role for cholesterol in entry of mycobacteria into macrophages. *Science (80-. ).* **288**, 1647–1650 (2000).
- 570 21. Lee, W., VanderVen, B. C., Fahey, R. J. & Russell, D. G. Intracellular 571 Mycobacterium tuberculosis exploits host-derived fatty acids to limit metabolic 572 stress. *J. Biol. Chem.* **288**, 6788–6800 (2013).
- 573 22. Meaney, S. Epigenetic regulation of cholesterol homeostasis. *Front. Genet.* **5**, 311 (2014).
- 575 23. Rayner, K. J. *et al.* Antagonism of miR-33 in mice promotes reverse 576 cholesterol transport and regression of atherosclerosis. *J. Clin. Invest.* **121**, 577 2921–2931 (2011).
- 578 24. Cardoso, F. et al. Myeloid Sirtuin 2 Expression Does Not Impact Long-Term

- Mycobacterium tuberculosis Control. *PLoS One* **10**, e0131904 (2015).
- 580 25. Cheng, C. Y. *et al.* Host sirtuin 1 regulates mycobacterial immunopathogenesis and represents a therapeutic target against tuberculosis. *Sci. Immunol.* **2**, 582 (2017).
- 583 26. Ghosh, H. S., Reizis, B. & Robbins, P. D. SIRT1 associates with eIF2-alpha and regulates the cellular stress response. *Sci. Rep.* **1**, 1–9 (2011).
- 585 27. Hayakawa, T. *et al.* SIRT1 suppresses the senescence-associated secretory phenotype through epigenetic gene regulation. *PLoS One* **10**, (2015).
- 587 28. Hou, X. *et al.* SIRT1 regulates hepatocyte lipid metabolism through activating AMP-activated protein kinase. *J. Biol. Chem.* **283**, 20015–20026 (2008).
- 589 29. Liu, T. F. & McCall, C. E. Deacetylation by SIRT1 Reprograms Inflammation and Cancer. *Genes and Cancer* **4**, 135–147 (2013).
- 591 30. Yuan, H. *et al.* Activation of stress response gene SIRT1 by BCR-ABL promotes leukemogenesis. *Blood* **119**, 1904–1914 (2012).
- 593 31. Shi, L., Jiang, Q., Bushkin, Y., Subbian, S. & Tyagi, S. Biphasic dynamics of macrophage immunometabolism during Mycobacterium tuberculosis infection. *MBio* **10**, (2019).
- 596 32. Dang, W. The controversial world of sirtuins. *Drug Discovery Today: Technologies* vol. 12 e9 (2014).
- 598 33. Kanfi, Y. *et al.* The sirtuin SIRT6 regulates lifespan in male mice. *Nature* **483**, 599 218–221 (2012).
- Sebastián, C. *et al.* The histone deacetylase SIRT6 Is a tumor suppressor that controls cancer metabolism. *Cell* **151**, 1185–1199 (2012).
- 602 35. Wu, X., Cao, N., Fenech, M. & Wang, X. Role of Sirtuins in Maintenance of Genomic Stability: Relevance to Cancer and Healthy Aging. *DNA and Cell Biology* vol. 35 542–575 (2016).
- 605 36. Elhanati, S. *et al.* Multiple regulatory layers of SREBP1/2 by SIRT6. *Cell Rep.* **4**, 905–912 (2013).
- 607 37. Gazzar, M. E. E. El *et al.* Chromatin-Specific Remodeling by HMGB1 and Linker Histone H1 Silences Proinflammatory Genes during Endotoxin Tolerance. *Mol. Cell. Biol.* **29**, 1959–1971 (2009).
- 610 38. El Gazzar, M. *et al.* G9a and HP1 couple histone and DNA methylation to TNF\$\alpha{\\$}transcription silencing during endotoxin tolerance. *J. Biol. Chem.* **283**, 32198–32208 (2008).
- 613 39. Imai, K., Togami, H. & Okamoto, T. Involvement of histone H3 lysine 9 (H3K9)
  614 methyltransferase G9a in the maintenance of HIV-1 latency and its reactivation
  615 by BIX01294. *J. Biol. Chem.* **285**, 16538–16545 (2010).
- 616 40. Merkling, S. H. *et al.* The Epigenetic Regulator G9a Mediates Tolerance to RNA Virus Infection in Drosophila. *PLoS Pathog.* **11**, (2015).
- 618 41. Scheer, S. & Zaph, C. The lysine methyltransferase G9a in immune cell differentiation and function. *Frontiers in Immunology* vol. 8 429 (2017).
- 620 42. Tachibana, M. et al. G9a histone methyltransferase plays a dominant role in

- euchromatic histone H3 lysine 9 methylation and is essential for early embryogenesis. *Genes Dev.* **16**, 1779–1791 (2002).
- 43. Mukherjee, T. & Balaji, K. N. The WNT framework in shaping immune cell
   responses during bacterial infections. *Frontiers in Immunology* (2019)
   doi:10.3389/fimmu.2019.01985.
- 44. Villaseñor, T. *et al.* Activation of the Wnt pathway by Mycobacterium
   tuberculosis: A Wnt-Wnt Situation. *Frontiers in Immunology* vol. 8 50 (2017).
- 628 45. Mostoslavsky, R. *et al.* Genomic instability and aging-like phenotype in the absence of mammalian SIRT6. *Cell* **124**, 315–329 (2006).
- 630 46. Espenshade, P. J. SREBPs: Sterol-regulated transcription factors. *J. Cell Sci.* 631 **119**, 973–976 (2006).
- 632 47. DeVries-Seimon, T. *et al.* Cholesterol-induced macrophage apoptosis requires 633 ER stress pathways and engagement of the type A scavenger receptor. *J. Cell Biol.* **171**, 61–73 (2005).
- 48. Jin, X. *et al.* HO-1/EBP interaction alleviates cholesterol-induced hypoxia through the activation of the AKT and Nrf2/mTOR pathways and inhibition of carbohydrate metabolism in cardiomyocytes. *Int. J. Mol. Med.* **39**, 1409–1420 (2017).
- Kaminsky-Kolesnikov, Y. *et al.* Cholesterol Induces Nrf-2-and HIF-1 α Dependent Hepatocyte Proliferation and Liver Regeneration to Ameliorate Bile
   Acid Toxicity in Mouse Models of NASH and Fibrosis. *Oxid. Med. Cell. Longev.* **2020**, (2020).
- 50. Freigang, S. *et al.* Nrf2 is essential for cholesterol crystal-induced inflammasome activation and exacerbation of atherosclerosis. *Eur. J. Immunol.* 41, 2040–2051 (2011).
- Holla, S., Kurowska-Stolarska, M., Bayry, J. & Balaji, K. N. Selective inhibition of IFNG-induced autophagy by Mir155- and Mir31-responsive WNT5A and SHH signaling. *Autophagy* 10, 311–330 (2014).
- 649 52. Bell, K. F. S. *et al.* Neuronal development is promoted by weakened intrinsic 650 antioxidant defences due to epigenetic repression of Nrf2. *Nat. Commun.* **6**, 1– 651 15 (2015).
- 652 53. Rada, P. *et al.* WNT-3A regulates an Axin1/NRF2 complex that regulates antioxidant metabolism in hepatocytes. *Antioxidants Redox Signal.* **22**, 555–654 571 (2015).
- 54. Scott, C. C. *et al.* Wnt directs the endosomal flux of LDL -derived cholesterol and lipid droplet homeostasis . *EMBO Rep.* **16**, 741–752 (2015).
- 657 55. Mohn, W. W. *et al.* The actinobacterial mce4 locus encodes a steroid transporter. *J. Biol. Chem.* **283**, 35368–35374 (2008).
- 659 56. Nazarova, E. V. *et al.* Rv3723/LucA coordinates fatty acid and cholesterol uptake in Mycobacterium tuberculosis. *Elife* **6**, (2017).
- 661 57. Pandey, A. K. & Sassetti, C. M. Mycobacterial persistence requires the utilization of host cholesterol. *Proc. Natl. Acad. Sci. U. S. A.* **105**, 4376–4380 (2008).

- de Chastellier, C. & Thilo, L. Cholesterol depletion in Mycobacterium avium infected macrophages overcomes the block in phagosome maturation and
   leads to the reversible sequestration of viable mycobacteria in
- phagolysosome-derived autophagic vacuoles. *Cell. Microbiol.* **8**, 242–256 (2006).
- 59. Lin, H. H. *et al.* Association of obesity, diabetes, and risk of tuberculosis: Two population-based cohorts. *Clin. Infect. Dis.* **66**, 699–705 (2018).
- 671 60. Martens, G. W. *et al.* Hypercholesterolemia impairs immunity to tuberculosis. 672 *Infect. Immun.* **76**, 3464–3472 (2008).
- 61. Martens, G. W., Vallerskog, T. & Kornfeld, H. Hypercholesterolemic LDL receptor-deficient mice mount a neutrophilic response to tuberculosis despite the timely expression of protective immunity. *J. Leukoc. Biol.* **91**, 849–857 (2012).
- 677 62. Shimabukuro-Vornhagen, A. *et al.* Inhibition of Protein Geranylgeranylation 678 Specifically Interferes with CD40-Dependent B Cell Activation, Resulting in a 679 Reduced Capacity To Induce T Cell Immunity. *J. Immunol.* **193**, 5294–5305 680 (2014).
- 63. Yang, W. *et al.* Potentiating the antitumour response of CD8+ T cells by modulating cholesterol metabolism. *Nature* **531**, 651–655 (2016).
- 683 64. Ito, A. *et al.* Cholesterol Accumulation in CD11c+ Immune Cells Is a Causal and Targetable Factor in Autoimmune Disease. *Immunity* **45**, 1311–1326 (2016).
- 686 65. Kim, K. D. *et al.* Apolipoprotein A-I induces IL-10 and PGE2 production in human monocytes and inhibits dendritic cell differentiation and maturation.

  888 *Biochem. Biophys. Res. Commun.* **338**, 1126–1136 (2005).
- 689 66. Swirski, F. K. *et al.* Ly-6Chi monocytes dominate hypercholesterolemia-690 associated monocytosis and give rise to macrophages in atheromata. *J. Clin.* 691 *Invest.* **117**, 195–205 (2007).
- 692 67. Yvan-Charvet, L. *et al.* ABCA1 and ABCG1 protect against oxidative stress-693 induced macrophage apoptosis during efferocytosis. *Circ. Res.* **106**, 1861– 694 1869 (2010).
- 695 68. Zhu, X. *et al.* Increased cellular free cholesterol in macrophage-specific Abca1 knock-out mice enhances pro-inflammatory response of macrophages. *J. Biol. Chem.* **283**, 22930–22941 (2008).
- 698 69. Zhu, X. *et al.* Macrophage ABCA1 reduces MyD88-dependent toll-like receptor 699 trafficking to lipid rafts by reduction of lipid raft cholesterol. *J. Lipid Res.* **51**, 700 3196–3206 (2010).
- 701 70. Huang, F. C. The critical role of membrane cholesterol in Salmonella-induced autophagy in intestinal epithelial cells. *Int. J. Mol. Sci.* **15**, 12558–12572 (2014).
- 71. Abrams, M. E. *et al.* Oxysterols provide innate immunity to bacterial infection by mobilizing cell surface accessible cholesterol. *Nat. Microbiol.* **5**, 929–942 (2020).
- 707 72. Ahsan, F., Maertzdorf, J., Guhlich-Bornhof, U., Kaufmann, S. H. E. & Moura-

- Alves, P. IL-36/LXR axis modulates cholesterol metabolism and immune defense to Mycobacterium tuberculosis. *Sci. Rep.* **8**, 1520 (2018).
- 73. Korf, H. *et al.* Liver X receptors contribute to the protective immune response against Mycobacterium tuberculosis in mice. *J. Clin. Invest.* **119**, 1626–1637 (2009).
- 713 74. Jin, X. *et al.* HO-1 alleviates cholesterol-induced oxidative stress through activation of Nrf2/ERK and inhibition of PI3K/AKT pathways in endothelial cells. *Mol. Med. Rep.* **16**, 3519–3527 (2017).
- 75. Parihar, S. P. *et al.* Statin therapy reduces the mycobacterium tuberculosis burden in human macrophages and in mice by enhancing autophagy and phagosome maturation. *J. Infect. Dis.* **209**, 754–763 (2014).
- 719 76. Kim, T. S. *et al.* SIRT3 promotes antimycobacterial defenses by coordinating mitochondrial and autophagic functions. *Autophagy* **15**, 1356–1375 (2019).
- 721 77. Bhaskar, A. *et al.* Host sirtuin 2 as an immunotherapeutic target against tuberculosis. *Elife* (2020) doi:10.7554/eLife.55415.
- 78. Hamers, A. A. J. *et al.* Nur77-deficiency in bone marrow-derived macrophages modulates inflammatory responses, extracellular matrix homeostasis, phagocytosis and tolerance. *BMC Genomics* **17**, 162 (2016).
- 726 79. Zhang, X., Edwards, J. P. & Mosser, D. M. The expression of exogenous genes in macrophages: Obstacles and opportunities. *Methods Mol. Biol.* **531**, 123–143 (2009).
- 729 80. Malmquist, N. A., Moss, T. A., Mecheri, S., Scherf, A. & Fuchter, M. J. Small-730 molecule histone methyltransferase inhibitors display rapid antimalarial activity 731 against all blood stage forms in Plasmodium falciparum. *Proc. Natl. Acad. Sci.* 732 *U. S. A.* **109**, 16708–16713 (2012).
- 733 81. Singh, R. *et al.* Polyphosphate deficiency in Mycobacterium tuberculosis is associated with enhanced drug susceptibility and impaired growth in guinea pigs. *J. Bacteriol.* **195**, 2839–2851 (2013).

# 738 Figure legends

736 737

- 739 Figure 1. Interception of G9a and Sirt6 leads to restricted mycobacterial
- 540 **burden. (A)** BALB/c peritoneal macrophages were infected with H37Rv for 12 h,
- 741 protein level of G9a, SIRT6 and their respective histone modification marks,
- H3K9me1 and H3K9Ac, were assessed by immunoblotting. (B) Immunoblot for G9a
- and SIRT6 in human PBMCs infected with H37Rv for 12 h. (C, D) *In vivo* expression
- of G9a (C) and SIRT6 (D) was analyzed in lung cryosections of uninfected mice and
- mice infected with H37Rv for 56 days by immunofluorescence. (E-G) In vitro CFU

was assessed 48 h post H37Rv infection under the following conditions: **(E)** in BALB/c mouse peritoneal macrophages treated with G9a inhibitor (5 μΜ) or (**F)** in BMDMs derived from WT (littermate control) or *Sirt6* KO mice or **(G)** in BALB/c mouse peritoneal macrophages transiently transfected with siRNAs against G9a or Sirt6 or both. The MOI of infection is 1:10 (macrophage:mycobacteria) for all the *in vitro* experiments. All data represents the mean ± SEM from 3 independent experiments; \*, P < 0.05, \*\*, P < 0.01; ns, not significant (Student's t- test for E-F and One-way ANOVA for G) and the blots are representative of 3 independent experiments. Med, medium (uninfected/untreated cells maintained in DMEM supplemented with 10% heat inactivated FBS for the entire duration of the experiment); WT, wild type; KO, knock out; inh., inhibitor; NT, non-targeting; BMDM, bone marrow derived macrophages; PBMC, peripheral blood mononuclear cells. Scale bar, 10μm.

**Figure 2. G9a and Sirt6 induce cholesterol accumulation during mycobacterial pathogenesis. (A, B)** BALB/c mouse peritoneal macrophages transfected with NT or *Ehmt2* or *Sirt6* siRNA were assessed for free cholesterol level upon infection with tdTomato-expressing H37Rv for 48 h by immunofluorescence. **(A)** Representative images of Filipin stained macrophages and **(B)** its quantification (n=200-300). **(C, D)** BMDMs from WT (littermate control) and *Sirt6* KO mice were utilized to assess free cholesterol by Filipin staining upon tdTomato-expressing H37Rv infection for 48 h. **(C)** Representative images and **(D)** its quantification (n=200-300). **(E, F)** Lung cryosections from uninfected or 56 days H37Rv-infected/ G9a inhibitor (40mg/kg) treated BALB/c mice were assessed for free cholesterol by Filipin staining in macrophages stained with F4/80: **(E)** representative images and **(F)** quantification of

Filipin staining in F4/80 positive cells (i.e. macrophages). **(G, H)** Lung cryosections of uninfected and infected WT (littermate control) and *Sirt6* het mice were assessed for free cholesterol levels by Filipin staining in macrophages stained by F4/80: **(G)** representative images and **(H)** quantification of Filipin staining in F4/80 positive cells (i.e. macrophages). *In vivo* data represents the mean ± SEM from 2-3 mice. The MOI of infection is 1:10 (macrophage: mycobacteria) for all the *in vitro* experiments. All data represents the mean ± SEM from 3 independent experiments, \*, P < 0.05; \*\*, P < 0.01; \*\*\*, P < 0.001 (One-way ANOVA for B, D, F and H). Med, medium; WT, wild type; KO, knock out; het, heterozygous; inh., inhibitor; MFI, mean fluorescence intensity; CTCF, corrected total cell fluorescence; F4/80, macrophage marker; BMDM, bone marrow derived macrophages. Scale bar, 10µm.

Figure 3. Cholesterol biosynthesis and transport genes are selectively regulated by G9a and Sirt6. (A, B) BALB/c mouse peritoneal macrophages were transfected with NT or *Ehmt2* or *Sirt6* siRNA. Transfected cells were infected with H37Rv for 12 h and the expression of the indicated genes was assessed by qRT-PCR. (C, D) RNA was isolated from the lung homogenates from the indicated groups of mice after 56 days of total infection and the transcript levels of the indicated cholesterol metabolism genes was analyzed by qRT-PCR. (E) Surface expression of LRP2 was analyzed by immunofluorescence in BALB/c peritoneal macrophages pretreated with G9a specific inhibitor (5μM) for 1 h followed by infection with tdTomato-expressing H37Rv for 12 h. (F) Lung cryosections from uninfected or 56 days H37Rv-infected/ G9a inhibitor (40mg/kg) treated BALB/c mice were assessed for surface expression of LRP2 by immunofluorescence. (G) Lung cryosections of uninfected and 56 days H37Rv-infected WT (littermate control) and *Sirt6* het mice

797

798

799

800

801

802

803

804

805

806

807

808

809

810

811

812

813

814

815

816

817

818

819

were assessed for the protein level of ABCA1 by immunofluorescence. *In vivo* data represents the mean  $\pm$  SEM from 2-3 mice. The MOI of infection is 1:10 (macrophage: mycobacteria) for all the *in vitro* experiments. All data represents the mean  $\pm$  SEM from 3 independent experiments, \*, P < 0.05; \*\*, P < 0.01; \*\*\*, P < 0.001 (One-way ANOVA for A-D). Med, medium; WT, wild type; het, heterozygous; inh., inhibitor; NT, non-targeting. Scale bar, 10µm.

Figure 4. Mtb-induced G9a and Sirt6 mediate fine-tuning of cholesterol accumulation. (A) SREBP2 was immunoprecipitated in whole cell lysates of BALB/c mouse peritoneal macrophages infected with H37Rv for 12 h to assess its interaction with G9a by immunoblotting. (B, C) BALB/c mouse peritoneal macrophages were transfected with Srebpf2 siRNA and (B) infected for 12 h with H37Rv to assess the expression of cholesterol accumulation genes by qRT-PCR, or (C) infected with H37Rv for 48 h to analyze the in vitro CFU. (D, E) ChIP assay was performed to affirm the (D) recruitment of G9a and (E) corresponding H3K9me1 mark, on the promoters of the indicated genes upon 12 h infection with H37Rv in BALB/c mouse peritoneal macrophages. (F) Sequential ChIP was conducted to assess the corecruitment of SREBP2 and G9a at the promoters of Lrp2, Aacs, Hmgcs1, Mvd and Dhcr24 in mouse peritoneal macrophages infected with H37Rv for 12 h. (G, H) BALB/c mouse peritoneal macrophages infected with H37Rv for 12 h were analyzed by ChIP for (G) SIRT6 recruitment and (H) H3K9Ac mark, on the promoters of Abca1 and Abcq1. The MOI of infection is 1:10 (macrophage: mycobacteria) for all the in vitro experiments. All data represent the mean ± SEM from 3 independent experiments. The blots are representative of 3 independent experiments. \*, P < 0.05;

821

822

823

824

825

826

827

828

829

830

831

832

833

834

835

836

837

838

839

840

841

842

843

844

\*\*, P < 0.01; \*\*\*, P < 0.001; ns, not significant (One-way ANOVA for B, Student's ttest for C-H). Med, Medium; NT, non-targeting. Figure 5. Cholesterol accumulation regulates the expression of anti-oxidant genes during mycobacterial infection. (A-I) BALB/c mouse peritoneal macrophages were transfected with NT or *Ehmt2* and *Sirt6* siRNA or *Nfe2l2* siRNA or cholesterol accumulation genes siRNA (Lrp2, Aacs, Hmgcs1, Mvd and Dhcr24 siRNAs) followed by 48 h of H37Rv infection. (A, B) The expression of the indicated molecules was assessed at the protein level by immunoblotting. (C-F) CellROX staining was performed to assess ROS levels in macrophages; (C, E) representative images and (D, F) respective quantification (n=200-300). (G, H, I) In vitro CFU was assessed. The MOI of infection is 1:10 (macrophage: mycobacteria) for all the in vitro experiments. All data represent the mean ± SEM from 3 independent experiments. The blots are representative of 3 independent experiments. \*, P < 0.05; \*\*, P < 0.01; \*\*\*, P < 0.001; ns, not significant (Student's t-test for C-E, G and I). Med, Medium; NT, non-targeting; chol. accum. genes, cholesterol accumulation genes. Scale bar, 10µm. Figure 6. Epigenetic modifiers G9a and Sirt6 aid in mycobacterial pathogenesis. (A-J) Mice were aerosolized with 200 CFU of H37Rv. (A) Schematic of in vivo mouse TB model for G9a inhibitor therapeutic treatment. (B) CFU of H37Rv from G9a inhibitor (40mg/kg) treated and untreated BALB/c mice in lung and spleen, after 56 days of total infection and therapeutic treatment. (C-D) Lungs of BALB/c mice from the indicated groups were analyzed for TB pathology by H & E staining; (C) representative image, (D) % of granulomatous area and (E)

aid its survival within the host.

corresponding histological evaluation for granuloma score. **(F)** Schematic of *in vivo* TB infection model for WT (littermate control) and *Sirt6* het mice. **(G)** CFU of H37Rv in lung and spleen of WT (littermate control) and *Sirt6* het mice after 56 days of H37Rv infection. Lungs from the indicated groups of mice were analyzed by H and E staining; **(H)** representative images, **(I)** % of granulomatous area and **(J)** corresponding histological evaluation for granuloma score. All data represents the mean ± SEM from 4-5 mice, \*, P < 0.05; \*\*, P < 0.01; \*\*\*, P < 0.001 (Student's *t*-test for B, D, G and I). WT, Wild type; Het, heterozygous; inh., inhibitor. **Figure 7. Schematic**, Mycobacteria utilizes the host epigenetic factors G9a and SIRT6 to augment cholesterol accumulation and antioxidant responses in order to

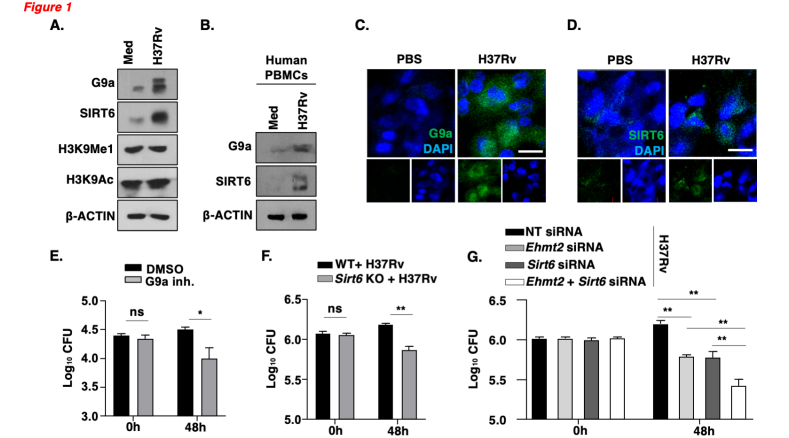


Figure 2

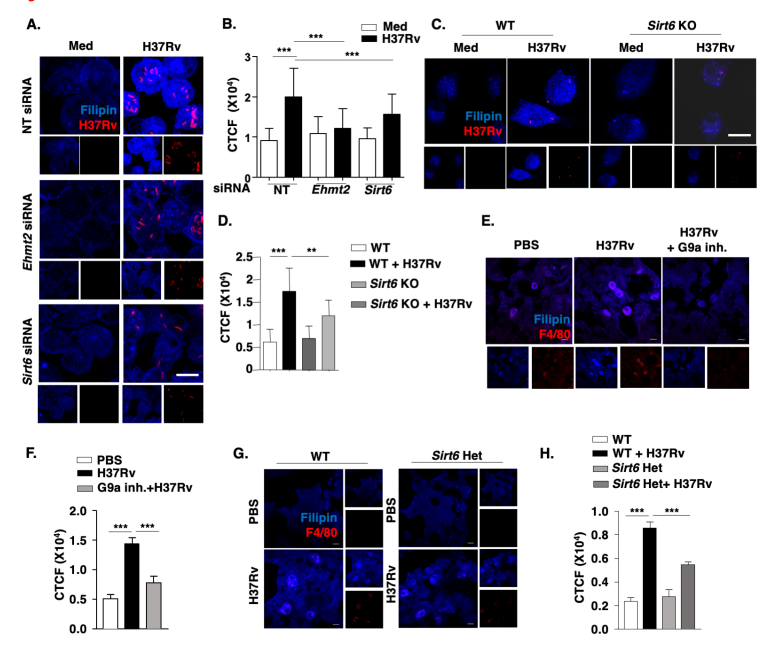


Figure 3

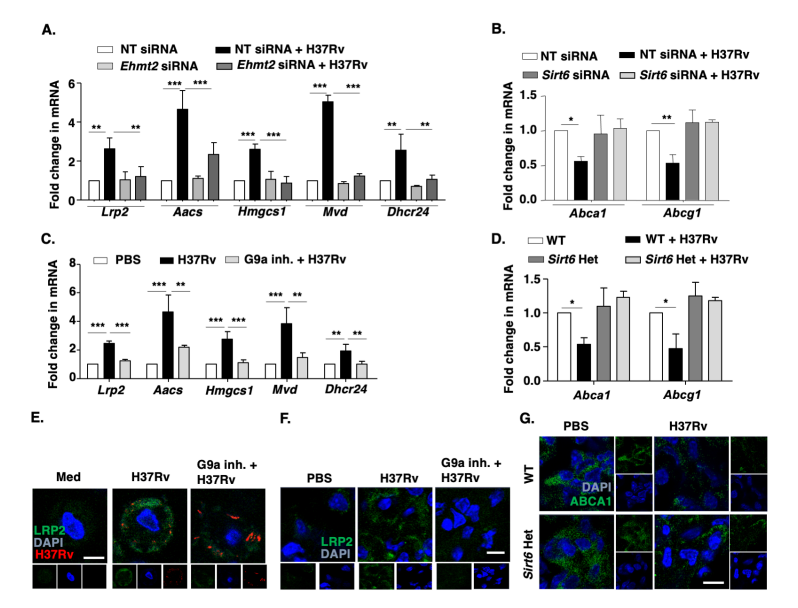


Figure 4

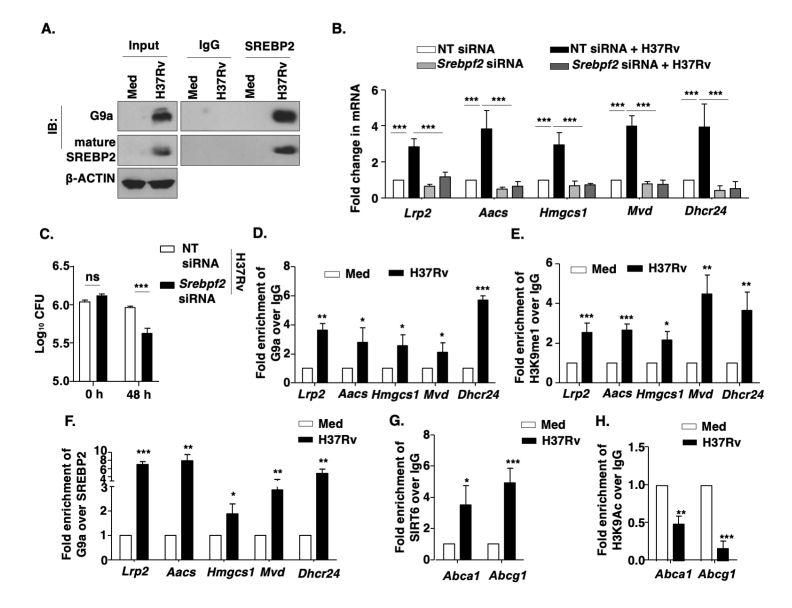


Figure 5

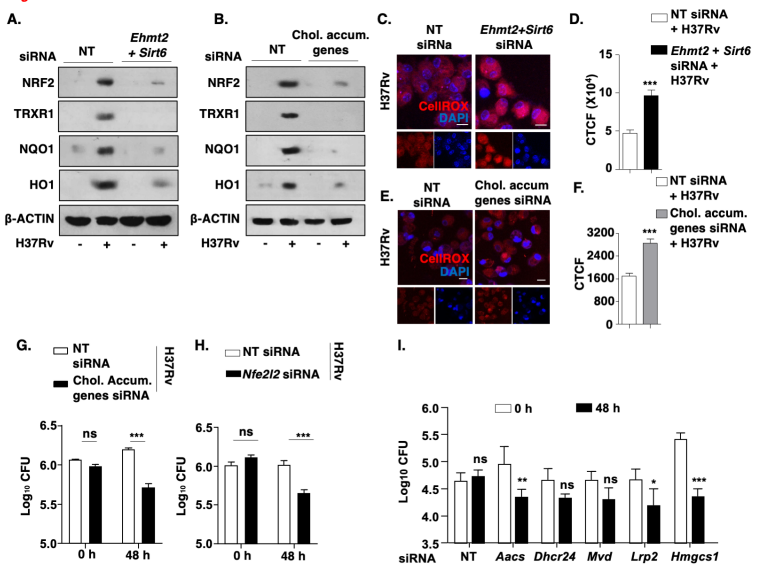
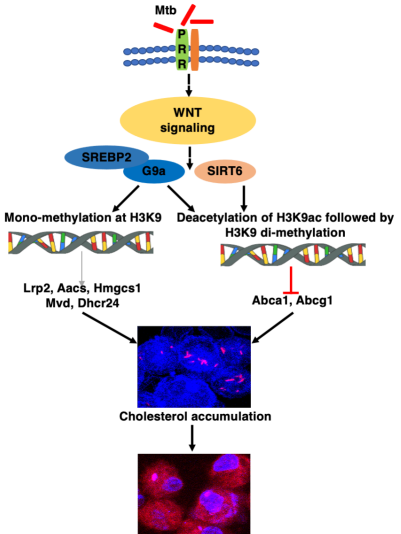


Figure 6 A. F. 56<sup>th</sup> day CFU and 0 day Aerosol infection with H37Rv 0 day Aerosol 28<sup>th</sup> day Inhibitor 56th day CFU and **Histo-pathology** infection with H37Rv treatment every day Histo-pathology G. В. Spleen Lung Lung Spleen 6.5 4.0 3.5-7.2 \*\*\* Log₁ 2.5-03.5 3.0 2.5 2.5 3.5 6.0 Log₁₀ CFU Log₁0 CFU 7.0 5.5 6.8 6.6 5.0 2.0 2.0 6.4 4.5 1.5 Sirt6 Het WT WT Sirt6 Het H37Rv G9a inh. H37Rv + + H. C. G9a inh. + H37Rv WT + H37Rv **PBS** H37Rv 5X D. Sirt6 Het Sirt6 Het+H37Rv % of area w.... Granulomatous lesion N P 9 8 0 **Granulomatous lesion** 80 % of area with 60-40-5X 20-Sirt6 Het H37Rv G9a inh. WT H37Rv E. J.

Group (BALB/c mice)	No. of Granuloma without necrosis Score (x2.5)	Total Granuloma Score
PBS	0	0
H37Rv	27	67.4
H37Rv + G9a inh	12	30

Group	No. of Granuloma without necrosis Score (x2.5)	Total Granuloma Score
WT, PBS	0	0
WT, H37Rv	25	62.5
Sirt6 Het, PBS	0	0
Sirt6 Het, H37Rv	14	35



Anti-oxidant responses-mediated enhanced TB progression

- 1 G9a and Sirtuin6 epigenetically modulate host cholesterol accumulation to
- 2 facilitate mycobacterial survival
- 3 Praveen Prakhar<sup>a,1</sup>, Bharat Bhatt<sup>a,1</sup>, Tanushree Mukherjee<sup>a,1</sup>, Gaurav Kumar Lohia<sup>a</sup>,
- 4 Ullas Kolthur-Seetharam<sup>b</sup>, Nagalingam Ravi Sundaresan<sup>a</sup>, R.S. Rajmani<sup>c</sup>, Kithiganahalli
- 5 Narayanaswamy Balaji<sup>a,c,2</sup>
- <sup>a</sup> Department of Microbiology and Cell Biology, Indian Institute of Science, Bangalore –
- 7 560012, Karnataka, India
- 8 b Department of Biological Sciences, Tata Institute of Fundamental Research, Mumbai –
- 9 400005, Maharashtra, India
- 10 <sup>c</sup> Centre for Infectious Disease Research, Indian Institute of Science, Bangalore -
- 11 560012, Karnataka, India

17

18

19

20

21

22

- <sup>1</sup> P.P., B.B., and T.M. contributed equally to this work.
- 13 <sup>2</sup> To whom correspondence may be addressed: Kithiganahalli Narayanaswamy Balaji,
- 14 Department of Microbiology and Cell Biology, Indian Institute of Science, Bangalore -
- 15 560012, Karnataka, India; Ph: +91-80-22933223; Email: balaji@iisc.ac.in

### 24 SUPPLEMENTARY FIGURES AND LEGENDS

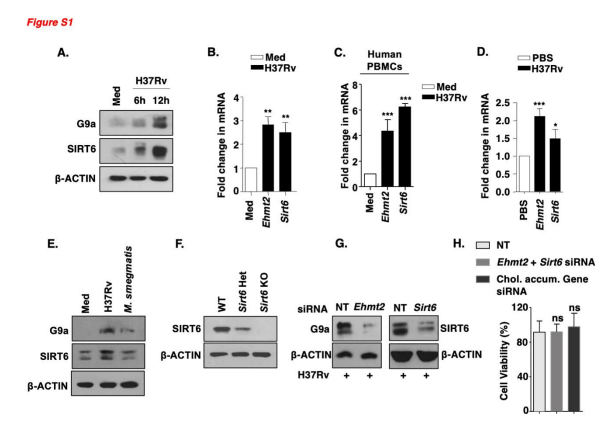


Figure S1. Mtb-triggered expression of epigenetic modifiers G9a and Sirt6 in host cells. (A) Immunoblot of G9a and SIRT6 in BALB/c mouse peritoneal macrophages infected with H37Rv for 6 h or 12 h. (B, D) Transcript level of the *Ehmt2* and *Sirt6* was analyzed by qRT-PCR (B) in BALB/c mouse peritoneal macrophages infected with H37Rv for 12 h or (C) in human PBMCs infected with H37Rv for 12 h or (D) in lung homogenates of mice infected with H37Rv for 56 days. (E) Protein level of G9a and SIRT6 was assessed in BALB/c macrophages infected with H37Rv or *M. smegmatis* for 12 h by immunoblotting. (F) The protein levels of SIRT6 was assessed in lung homogenates of WT (littermate control), *Sirt6* het and *Sirt6* KO mice by immunoblotting. (G) BALB/c mouse peritoneal macrophages were transfected with the indicated siRNAs and infected with H37Rv for 12 h. Whole cell lysates were assessed for the knock down

of G9a and SIRT6 by immunoblotting. **(H)** MTT assay was performed to assess cell viability of BALB/c macrophages transfected with NT or *Ehmt2* and *Sirt6* siRNA or chol. accum. genes siRNA (*Lrp2*, *Aacs*, *Hmgcs1*, *Mvd* and *Dhcr24*). The MOI of infection is 1:10 (macrophage: mycobacteria) for all the *in vitro* experiments. All data represents the mean ± SEM from 3 independent experiments. The blots are representative of 3 independent experiments. \*, P < 0.05; \*\*, P < 0.01; \*\*\*, P < 0.001 (Student's t-test for B, C, D and H). Med, Medium. NT, non-targeting; ns, not significant; WT, wild type; Het, heterozygous; KO, knock out; chol. accum. gene, cholesterol accumulation genes.

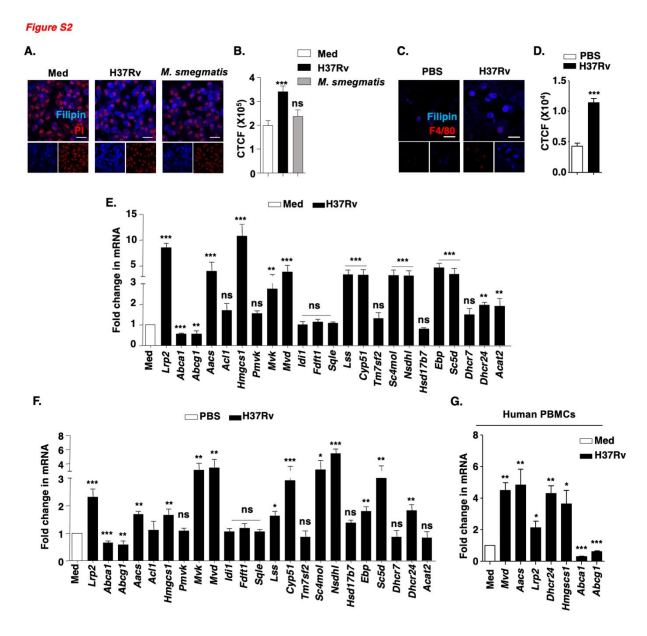


Figure S2. Mtb-driven free cholesterol accumulation in host cells. (A, B) BALB/c mouse peritoneal macrophages were infected with H37Rv or *M. smegmatis* for 48 h and assessed for cholesterol accumulation by Filipin staining; (A) representative image and (B) respective quantification (n=200-300). (C) Lung cryosections from BALB/c mice infected with H37Rv for 56 days was assessed for cholesterol accumulation by Filipin staining in macrophages stained with F4/80, (D) quantification of Filipin staining in F4/80 positive cells in lung cryosections. (E, F) Transcript level of the indicated set of genes

was analysed by qRT-PCR **(E)** in BALB/c mouse peritoneal macrophages infected with H37Rv for 12 h, **(F)** in lung homogenates of BALB/c mice infected with H37Rv for 56 days and **(G)** in human PBMCs infected with H37Rv for 12 h. The MOI of infection is 1:10 (macrophage:mycobacteria) for all the *in vitro* experiments. All data represents the mean ± SEM from 3 independent experiments. Confocal images were obtained from lungs of at least three groups of mice. \*, P < 0.05; \*\*, P < 0.01; \*\*\*, P < 0.001 (One-way ANOVA for B, Student's t-test for D-F). Med, Medium; CTCF; corrected total cell fluorescence; MFI, mean fluorescence intensity; PI, Propidium Iodide (nuclear stain); PBMC, peripheral blood mononuclear cells; Scale bar, 25 μm.



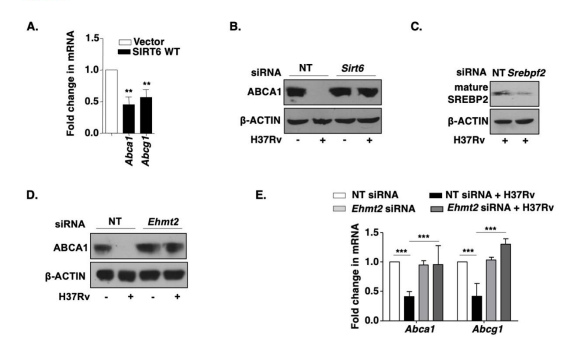


Figure S3. Interplay of G9a and Sirt6 in regulation of cholesterol efflux during Mtb infection. (A) RAW 264.7 macrophages were transfected with vector or SIRT6 WT construct and transcript levels of ABC transporters was analysed by qRT-PCR. (B-E) BALB/c mouse peritoneal macrophages were transfected with NT or *Ehmt2* or *Sirt6* or *Srebf2* siRNA as indicated, followed by 12 h infection with H37Rv. Whole cell lysates were assessed for (B) ABCA1 or (C) SREBP2 expression by immunoblotting. (D) ABCA1 was assessed by western blotting and (E) transcript level of the indicated genes were measured by qRT-PCR. The MOI of infection is 1:10 (macrophage: mycobacteria) for all the *in vitro* experiments. All data represents the mean ± SEM from 3 independent experiments. The blots are representative of 3 independent experiments. \*\*, P < 0.01; \*\*\*\*, P < 0.001 (Student's t-test for A, One-way ANOVA for D and E); Med, Medium; NT, non-targeting; ns, not significant.

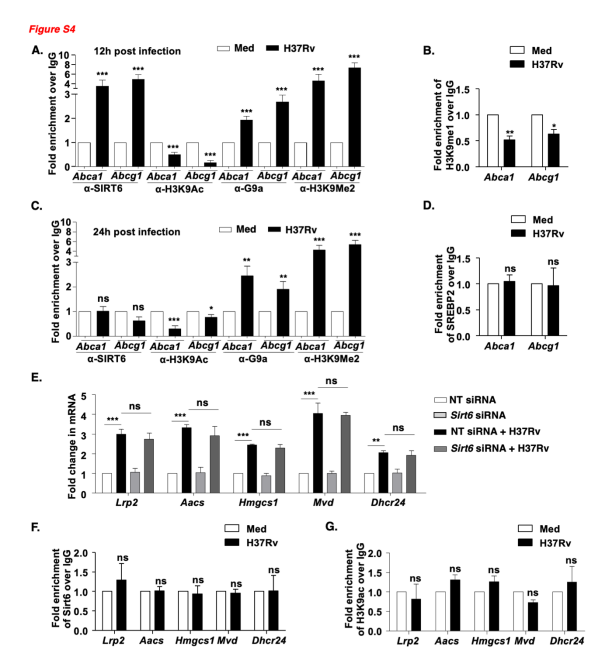


Figure S4. G9a and Sirt6 mediated transcriptional regulation of cholesterol efflux and synthesis genes. (A, B) BALB/c mouse peritoneal macrophages were infected with H37Rv for 12 h, and assessed for the recruitment of SIRT6, G9a and presence of H3K9Ac H3K9me2 and H3K9me1 (B), on the promoters of *Abca1* and *Abcg1*. (C) BALB/c mouse peritoneal macrophages were infected with H37Rv for 24h and assessed for recruitment of SIRT6, G9a, H3K9Ac and H3K9me2 on promoters of *Abca1* 

and *Abcg1* **(D)** ChIP analysis of SREBP2 on the promoters of *Abca1* and *Abcg1* in BALB/c mouse peritoneal macrophages after 12 h H37Rv infection. **(E)** Cholesterol accumulation genes were assessed in BALB/c mouse peritoneal macrophages transfected with NT or *Sirt6* siRNA post 12h of H37Rv infection. **(F, G)** Recruitment of SIRT6 and H3K9Ac on the promoters of cholesterol accumulation genes in BALB/c mouse peritoneal macrophages upon 12 h of H37Rv infection. The MOI of infection is 1:10 (macrophage: mycobacteria) for all the *in vitro* experiments. All data represents the mean ± SEM from 3 independent experiments, \*, P < 0.05; \*\*, P < 0.01; \*\*\*, P < 0.001 (Student's t-test in A-F). Med, Medium; NT, non-targeting; ns, not significant.

#### Figure S5

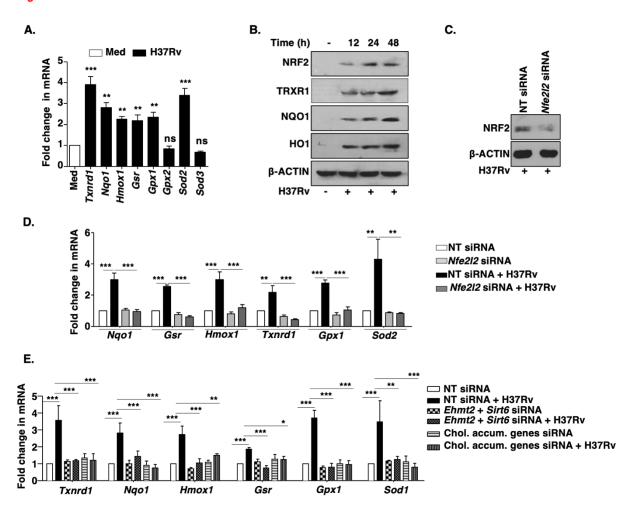


Figure S5. NRF2 and its target genes are expressed during Mtb infection. (A) BALB/c mouse peritoneal macrophages were infected with H37Rv for 48 h and the expression of NRF2 target genes was assessed by qRT-PCR. (B) BALB/c mouse peritoneal macrophages were infected with H37Rv for the indicated time points and whole cell lysates were assessed for the expression of NRF2 and its target genes. (C) Immunoblotting to validate NRF2 knockdown in murine macrophages transfected with Nfe2/2 siRNA. (D-E) BALB/c mouse peritoneal macrophages were transfected with NT or (D) Nfe2/2 siRNA or (E) Ehmt2 and Sirt6 siRNA or Chol accum genes siRNA (combination of Lrp2, Aacs, Hmgcs1, Mvd and Dhcr24 siRNAs) in the presence or

absence of 48 h infection with H37Rv and the transcript levels of NRF2 target genes were analysed by qRT-PCR. The MOI of infection is 1:10 (macrophage:mycobacteria) for all the *in vitro* experiments. All data represents the mean ± SEM from 3 independent experiments; \*, P < 0.05; \*\*, P < 0.01; \*\*\*, P < 0.001 (Student's t- test for A and Oneway ANOVA for D, E) and the blots are representative of 3 independent experiments. Med, Medium; NT, non-targeting; ns, not significant; chol. accum. genes, cholesterol accumulation genes.

#### Figure S6

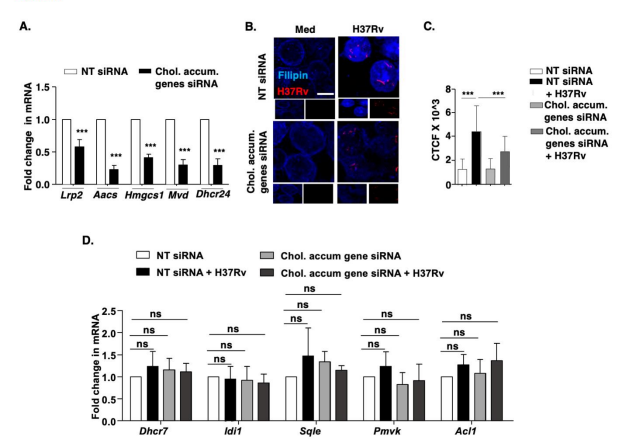
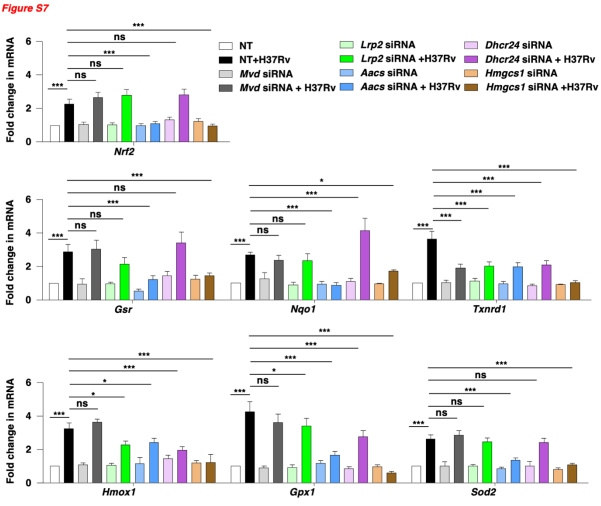


Figure S6. Validation of abrogated cholesterol accumulation in cells knocked down for genes involved in cholesterol biosynthesis and uptake. (A-D) BALB/c mouse peritoneal macrophages were transfected with siRNAs against the selected cholesterol accumulation genes (combination of *Lrp2*, *Aacs*, *Hmgcs1*, *Mvd* and *Dhcr24* siRNAs) or NT and (A) the expression of the concerned cholesterol genes was analysed by qRT-PCR; (B-D) followed by H37Rv infection for 48 h and cholesterol accumulation was confirmed by Filipin staining; (B) representative image, (C) respective quantification (n=200-300); (D) qRT-PCR to assess transcript levels of indicated genes. The MOI of infection is 1:10 (macrophage:mycobacteria) for all the *in vitro* experiments. All data represents the mean ± SEM from 3 independent experiments, \*\*\*, P < 0.001 (Student's t- test for A; and One-way ANOVA for C); CTCF, corrected total cell

- 125 fluorescence; NT, non-targeting; ns, not significant; chol. accum. gene, cholesterol
- 126 accumulation genes. Scale bar, 10 μm.



**Figure S7. Cholesterol biosynthesis/uptake pathway modulates NRF2 and its target genes.** BALB/c mouse peritoneal macrophages were transfected with NT or *Lrp2* or *Aacs* or *Hmgcs1* or *Mvd* or *Dhcr24* siRNAs and the transcript level of NRF2 and its target genes were assessed 48 h post Mtb infection. The MOI of infection is 1:10 (macrophage: mycobacteria) for all the *in vitro* experiments. Data represents the mean  $\pm$  SEM from 3 independent experiments; \*, P < 0.05; \*\*\*, P < 0.001 (Two-way ANOVA); NT, non-targeting; ns, not significant.

#### Figure S8

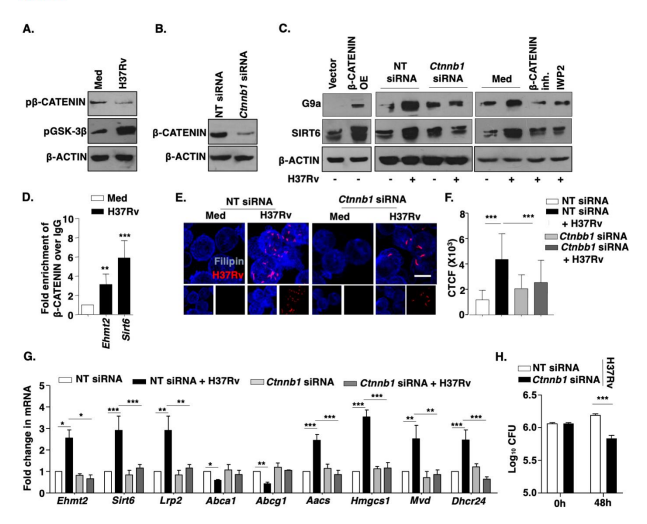


Figure S8: Contribution of WNT/β-CATENIN axis in mycobacterial infection. (A)

BALB/c mouse peritoneal macrophages were infected with H37Rv for 1 h and whole cell lysates were assessed for the activation of WNT pathway. **(B)** Immunoblotting to validate  $\beta$ -CATENIN knockdown in BALB/c mouse macrophages transfected with NT or *Ctnnb1* siRNA . **(C)** RAW 264.7 macrophages were transfected with  $\beta$ -CATENIN OE construct (**C**, left panel) or mouse peritoneal macrophages were transfected with NT or *Ctnnb1* siRNA (**C**, middle panel) or BALB/c mouse peritoneal macrophages were pretreated with  $\beta$ -CATENIN inhibitor (15  $\mu$ M) or IWP2 (5  $\mu$ M) for 1 h (**C**, right panel), followed by 12 h infection with H37Rv. Whole cell lysates were assessed for SIRT6 and

G9a expression by immunoblotting. (D) β-CATENIN recruitment at the promoter of Ehmt2 and Sirt6 was assessed by ChIP assay in BALB/c mouse primary macrophages infected with H37Rv for 12 h. (E, F) Free cholesterol was assessed by Filipin staining in BALB/c mouse peritoneal macrophages transfected with NT or Ctnnb1 siRNA followed by 48 h infection with tdTomato H37Rv, (E) representative image and (F) respective quantification (n=200-300). (G) Indicated genes were analyzed at transcript level by qRT- PCR in BALB/c mouse peritoneal macrophages that were transfected with NT or Ctnnb1 siRNA followed by infection with H37Rv for 12 h. (H) BALB/c mouse macrophages were transfected with NT or Ctnnb1 siRNA and in vitro CFU was assessed at the indicated time points post H37Rv infection. The MOI of infection is 1:10 (macrophage:mycobacteria) for all the in vitro experiments. All data represents the mean ± SEM from 3 independent experiments, \*, P < 0.05; \*\*, P < 0.01; \*\*\*, P < 0.001 (Student's t- test for D; One-way ANOVA for F-H). All blots are representative of 3 independent experiments. Med, medium;  $\beta$ -CATENIN OE,  $\beta$ -CATENIN over expression; NT, non-targeting; inh., inhibitor. Scale bar, 10 µm.

171

172

173

174

175

176

177

178

179

180

181

182

183

### MATERIALS AND METHODS

#### **Antibodies**

HRP-conjugated anti-β-ACTIN antibody, Filipin and 4',6-Diamidino-2phenylindole dihydrochloride (DAPI) were purchased from Sigma-Aldrich/ Merck Millipore. Alexa488-conjugated anti-rabbit IgG, HRP-conjugated anti-rabbit total IgG and light chain specific IgG antibodies were purchased from Jackson ImmunoResearch, USA; PE-conjugated F4/80 was procured from Tonbo Biosciences, USA. Anti-G9a, anti-SIRT6, anti-H3K9me1, anti-H3K9me2, anti-H3K9Ac, anti-Ser33/37/Thr41 phospho-β-CATENIN, anti-Ser9 phospho-GSK-3β, anti-β-CATENIN, anti-NRF2, anti-HO1 and anti-TRXR1 antibodies were obtained from Cell Signaling Technology, USA. Anti-LRP2 antibody was purchased from Santa Cruz Biotechnology, USA; anti-SREBP2 antibody was procured from Abcam, USA; and anti-NQO1 antibody was purchased from Calbiochem, USA.

184

185

186

187

188

189

190

191

192

#### Treatment with pharmacological reagents

Cells were treated with concerned pharmacological inhibitors for 1 h prior to the experiment at the following final concentrations: BIX-01294 (G9a inhibitor, 5  $\mu$ M),  $\beta$ -CATENIN inhibitor (15  $\mu$ M), IWP-2 (5  $\mu$ M). DMSO at 0.1% concentration was used as the vehicle control. In all experiments involving pharmacological reagents, a tested concentration was used after careful titration experiments assessing the viability of the macrophages using the MTT (3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) assay.

## MTT assay

siRNA transfected mouse peritoneal macrophages were treated with 3-(4, 5-Dimethylthiazol-2-yl)-2, 5-diphenyltetrazolium bromide (MTT) for 4 h at a final concentration of 0.5 mg/ml. Media was gently removed post incubation and 200 µL of DMSO was added. This solubilized purple formazan crystals were quantified by measuring absorbance at 550 nm in an 96-well plate reader. Viability of siRNA transfected macrophages were assessed relative to non-transfected macrophages.

### **Isolation of Human PBMCs**

Histopaque-1077 (Sigma-Aldrich, USA) polysucrose solution was utilized to isolate PBMCs from whole blood as per manufacturer's instruction. Briefly, 3 ml of whole blood was carefully layered onto 3 ml of Histopaque-1077 in a 15 ml conical centrifuge tube followed by centrifugation at  $400 \times g$  for 30 min at room temperature. Upper layer was carefully removed without disturbing the opaque interface of mononuclear cells. The interface was transferred into a fresh 15 ml conical centrifuge tube and resuspended in 10 ml isotonic phosphate buffered saline solution. The solution was centrifuged at  $250 \times g$  for 10 min, cell pellet was resuspended and cultured in RPMI supplemented with 10 % heat inactivated FBS (Gibco-Life Technologies) in the presence of 10 ng/ml M-CSF (PeproTech, USA) for 5 days at 37 °C in 5 % CO<sub>2</sub> incubator and utilized for experiments.

## RNA isolation and Real-Time qRT-PCR

Total RNA from treated, untreated and infected macrophages were isolated using TRI reagent (Sigma). 2 µg of total RNA was converted into cDNA using First Strand cDNA synthesis kit (Applied Biological Materials Inc.). Target gene expression was assessed by Real-Time quantitative Reverse Transcription-PCR (qRT-PCR) using SYBR Green PCR mix (Thermo Fisher Scientific). All the experiments were repeated at least 3 times independently to ensure the reproducibility of the results. *Gapdh* was used as internal control. The list of primers is detailed in Supplementary Tables 1 and 2.

## Immunoblotting analysis

Cells post treatment and/or infection were washed with 1X PBS. Whole cell lysate was prepared by lysing in RIPA buffer [50 mM Tris-HCl (pH 7.4), 1 % NP-40, 0.25 % sodium deoxycholate, 150 mM NaCl, 1 mM EDTA, 1 mM PMSF, 1 μg/ml each of aprotinin, leupeptin, pepstatin, 1 mM Na<sub>3</sub>VO<sub>4</sub>, 1 mM NaF] on ice for 30 min. Total protein from whole cell lysates was estimated by Bradford reagent. Equal amount of protein from each cell lysate was resolved on 12 % SDS-PAGE and transferred onto PVDF membranes (Millipore) by semi-dry immunoblotting method (Bio-Rad). 5 % non-fat dry milk powder in TBST [20 mM Tris-HCl (pH 7.4), 137 mM NaCl, and 0.1 % Tween 20] was used for blocking nonspecific binding for 60 min. After washing with TBST, the blots were incubated overnight at 4 °C with primary antibody diluted in TBST with 5 % BSA. After washing with TBST, blots were incubated with secondary antibody conjugated to HRP for 4 h at 4 °C. The immunoblots were developed with enhanced chemilluminescence detection system (PerkinElmer) as per manufacturer's instructions.

β-ACTIN was used as loading control.

## **Immunoprecipitation Assay**

Immunoprecipitation assays were carried out following a modified version of the protocol provided by Millipore, USA. In brief, macrophages were gently resuspended and lysed in ice-cold RIPA buffer. The cell lysates obtained were subjected to preclearing with BSA-blocked Protein A beads. The amount of protein was estimated in the supernatant and equal amount of protein was incubated with IgG or anti-SREBP2 antibody for 4 h at 4 °C. The immune complexes were captured on protein A agarose beads (Bangalore Genei, India) at 4 °C for 4 h. The beads were separated, washed and boiled in Laemmli buffer for 10 min. These bead free samples were analyzed for respective target molecules by immunoblotting. Light chain specific secondary antibody was used for immunoblotting after immunoprecipitation.

#### **Chromatin Immunoprecipitation (ChIP) Assay**

ChIP assays were carried out using a protocol provided by Upstate Biotechnology and Sigma-Aldrich with certain modifications. Briefly, macrophages were fixed with 3.6 % formaldehyde for 15 min at room temperature followed by inactivation of formaldehyde with addition of 125 mM glycine for 10 min. Nuclei were lysed in 0.1 % SDS lysis buffer [50 mM Tris-HCl (pH 8.0), 200 mM NaCl, 10 mM HEPES (pH 6.5), 0.1 % SDS, 10 mM EDTA, 0.5 mM EGTA, 1 mM PMSF, 1 μg/ml of each aprotinin, leupeptin, pepstatin, 1 mM Na<sub>3</sub>VO<sub>4</sub> and 1 mM NaF]. Chromatin was sheared using

Bioruptor Plus (Diagenode, Belgium) at high power for 70 rounds of 30 sec pulse ON and 45 sec pulse OFF. Chromatin extracts containing DNA fragments with an average size of 500 bp were immunoprecipitated with SIRT6 or G9a or H3K9Ac or H3K9me1 or H3K9me2 or β-CATENIN antibodies or rabbit preimmune sera complexed with Protein A agarose beads (Bangalore Genei, India). Immunoprecipitated complexes were sequentially washed with Wash Buffer A, B and TE [Wash Buffer A: 50 mM Tris-HCl (pH 8.0), 500 mM NaCl, 1 mM EDTA, 1 % Triton X-100, 0.1 % Sodium deoxycholate, 0.1 % SDS and protease/phosphatase inhibitors; Wash Buffer B: 50 mM Tris-HCl (pH 8.0), 1 mM EDTA, 250 mM LiCl, 0.5% NP-40, 0.5 % Sodium deoxycholate and protease/phosphatase inhibitors; TE: 10 mM Tris-HCl (pH 8.0), 1 mM EDTA] and eluted in elution buffer [1 % SDS, 0.1 M NaHCO3]. After treating the eluted samples with RNase A and Proteinase K, DNA was precipitated using phenol-chloroform-ethanol method. Purified DNA was analyzed by quantitative real time RT-PCR. All values in the test samples were normalized to amplification of the specific gene in Input and IgG pull down and represented as fold change in modification or enrichment. All ChIP experiments were repeated at least three times. The list of primers is detailed in Supplementary Table 3.

279

280

281

282

283

284

278

262

263

264

265

266

267

268

269

270

271

272

273

274

275

276

277

### Sequential ChIP Assay

The protocol for sequential ChIP was adopted from (de Medeiros, 2011; Truax and Greer, 2012). Briefly, the DNA fragments obtained following sonication [in lysis buffer; 1 % SDS, 10 mM EDTA, 50 mM Tris-HCl (pH 8.0)] were immunoprecipitated with SREBP2-complexed Protein A beads. After first pull down, beads were washed with Re-

ChIP Buffer [2 mM EDTA, 500 mM NaCl, 0.1 % SDS, 1 % NP40], followed by elution of DNA in Re-ChIP elution buffer [2 % SDS, 15 mM DTT in TE] at 37 °C for 30 min. The eluted DNA was subjected to subsequent round to immunoprecipitation with Protein-A beads pre-complexed with G9a or rabbit pre-immune sera. Immunoprecipitated complexes were sequentially washed with Wash Buffer A, B and TE [Wash Buffer A: 20 mM Tris-HCl (pH 8.0), 150 mM NaCl, 2 mM EDTA, 1% Triton X-100, 0.1% SDS and protease/phosphatase inhibitors; Wash Buffer B: 20 mM Tris-HCl (pH 8.0), 2 mM EDTA, 500 mM NaCl, 1 % Triton X-100, 0.1 % SDS and protease/phosphatase inhibitors; Wash Buffer C: 10 mM Tris-HCl (pH 8.0), 1 mM EDTA, 1 % sodium deoxycholate, 1 % NP40, 250 mM LiCl and protease/phosphatase inhibitors; TE: 10 mM Tris-HCl (pH 8.0), 1 mM EDTA and protease/phosphatase inhibitors] and eluted [0.1 M NaHCO<sub>3</sub>, 1 % SDS], purified and subjected to qRT-PCR (as described previously). The fold change of SREBP2-G9a versus SREBP2-IgG upon infection signified the co-occupancy of the two factors at concerned promoters. The list of primers is given in Supplementary Table 3.

## **Isolation and culture** of murine bone marrow derived macrophages

Mice tibia and femur were flushed with ice-cold DMEM containing 10 % fetal bovine serum from WT (littermate control) and *sirt6* KO mice. Bone marrow was collected in 50 ml tube and bone marrow clusters were disintegrated by vigorous pipetting. The cell suspension was centrifuged at 1500 rpm for 5 min at 4 °C followed by two washes with DMEM containing 10 % fetal bovine serum. Then the cells were suspended in DMEM containing 10 % fetal bovine serum and 20 % of L929 cell supernatant and seeded at 1 million cells per well and incubated at 37 °C, 5 % CO<sub>2</sub> and

95 % humidity in a  $CO_2$  incubator. The medium was supplemented on the 3<sup>rd</sup> and 5<sup>th</sup> day with DMEM containing 10 % fetal bovine serum and 20 % L929 cell supernatant. Post 7 days of differentiation, the cells were used for further experiments.

### In vitro Mtb CFU

BALB/c peritoneal macrophages transfected with *Ehmt2*, *Sirt6*, Srebf2, cholesterol genes (*Lrp2*, *Mvd*, *Aacs*, *Hmgcs*, *Dhcr24*), *Nfe2l2*, *Ctnnb1* or non-targeting siRNA for 24 h; or BMDMs obtained from *Sirt6* KO mice were infected with Mtb H37Rv at MOI 5 for 4 h. Post 4 h, the cells were thoroughly washed with PBS to remove any surface adhered bacteria and medium containing amikacin (0.2 mg/ml) was added for 2 h to kill any extracellular mycobacteria. After amikacin treatment, the cells thoroughly washed with PBS were taken as 0 h time point and a duplicate set was maintained in antibiotic free medium for next 48 h. Intracellular mycobacteria was enumerated by lysing macrophages with 0.06 % SDS in 7H9 Middlebrook medium. Appropriate dilutions were plated onto Middlebrook 7H11 agar plates supplemented with OADC (oleic acid, albumin, dextrose, catalase). Total colony forming units (CFUs) were counted after 21 days of plating.

# Microtomy and Hematoxylin and Eosin (H&E) staining

Microtome sections (5 μm) were obtained from formalin-fixed, paraffin-embedded mouse lung tissue samples using Leica RM2245 microtome. These sections were first deparaffinized and rehydrated. The rehydrated sections were subjected to Hematoxylin staining followed by Eosin staining as per manufacturer instructions. The sections were

then dehydrated and mounted with coverslip using permount. Sections were given to consultant pathologist for blinded analyses.

## **Cryosection preparation**

The excised and fixed lungs were placed in the optimal cutting temperature (OCT) media (Jung, Leica). Cryosections of 10-15  $\mu$ m were prepared using Leica CM 1510 S or Leica CM 3050 S cryostat with the tissue embedded in OCT being sectioned onto glass slides and then stored at -80  $^{\circ}$ C.

## Immunofluorescence (IF)

Treated/infected macrophages were fixed with 3.6 % formaldehyde for 30 min at room temperature. The cells were washed with PBS and blocked in 2 % BSA in PBST. After blocking, cells were stained with LRP2 overnight at 4 °C. Then they were incubated with DyLight 488-conjugated secondary antibody for 2 h and nuclei were stained with DAPI. The coverslips were mounted on a slide with glycerol. For IF of the cryosections, frozen sections were thawed to room temperature. After blocking with 2 % BSA containing saponin, the sections were stained with specific antibodies overnight at 4 °C. The sections were then incubated with DyLight 488-conjugated secondary antibody for 2 h and nuclei were stained with DAPI. A coverslip was mounted on the section with glycerol as the medium. Confocal images were taken with Zeiss LSM 710 Meta confocal laser scanning microscope (Carl Zeiss AG, Germany) using a plan-Apochromat 63X/1.4 Oil DIC objective (Carl Zeiss AG, Germany) and images were analyzed using ZEN black software. CTCF (corrected total cell fluorescence) was

calculated as (fluorescence observed in an area of a cell – fluorescence of background for the same area) using ImageJ. Cells boundaries were demarcated based on brightfield image and the fluorescence intensities of different channels were measured. Background fluorescence intensity was measured from a field devoid of cells.

## Filipin fluorescence staining for Free Cholesterol

Filipin complex (Sigma-Aldrich, USA) was utilized to assess free cholesterol following protocol from (Leventhal et al, 2001). Briefly, mouse peritoneal macrophages were fixed with 3.6 % paraformaldehyde for 1 h at room temperature. After incubation, cells were washed with 1X PBS followed by incubation in 1.5 mg glycine per ml PBS for 10 min at room temperature. Filipin staining was then performed at a final concentration of 0.05 mg/ml in PBS for 2 h at room temperature. Cell were washed thrice with 1X PBS and nuclei were stained with propidium iodide (PI). For Filipin staining of cryosections, frozen sections were thawed to room temperature. After blocking with 2 % BSA containing saponin, the sections were stained with Filipin (0.05 mg/ml in PBS) and PEconjugated F4/80 (macrophage marker) for 2 h at room temperature. The samples were mounted on glycerol. Images were captured in Zeiss LSM 710 confocal laser scanning microscope as described above.

#### **CellROX Oxidative Stress Reagent staining**

CellROX Deep Red Reagent (Thermo Fisher Scientific, USA) was utilized to measure oxidative stress in macrophages as per manufacturer's instructions. In brief, siRNA transfected mouse peritoneal macrophages were treated with CellROX Deep

Red Reagent at a final concentration of 5  $\mu$ M and incubated for 30 min at 37 °C. Cells were then washed with 1X PBS thrice followed by fixation with 3.6 % formaldehyde for 15 min. Nuclei were stained with DAPI and images were captured in Zeiss LSM 710 confocal laser scanning microscope.

## 

# **Supplementary Tables:**

# 

# **Supplementary Table 1: List of primers for mouse gene expression analyses**

SI. No.	Gene Name	Forward/ Reverse	Sequence (5'-3')
1.	Glyceraldehyde-3-phosphate	Forward	gagccaaacgggtcatcatct
	dehydrogenase ( <i>Gapdh</i> )	Reverse	gaggggccatccacagtctt
2.	Euchromatic histone lysine N-	Forward	agccaagaggggtctccaat
	methyltransferase 2 (Ehmt2/G9a)	Reverse	ctcgctgatgcggtcaatct
3.	Sirtuin 6 (Sirt6)	Forward	atgtcggtgaattatgcagca
		Reverse	gctggaggactgccacatta
4.	Low density lipoprotein	Forward	aaaatggaaacggggtgactt
	receptor-related protein 2 ( <i>Lrp2</i> )	Reverse	ggctgcatacattgggttttca
5.	ATP-binding cassette, sub-	Forward	aaaaccgcagacatccttcag
	family A (ABC1), member 1 (Abca1)	Reverse	cataccgaaactcgttcaccc
6.	ATP binding cassette	Forward	gtggatgaggttgagacagacc
	subfamily G member 1 (Abcg1)	Reverse	cctcgggtacagagtaggaaag
7.	Acetoacetyl-CoA synthetase	Forward	gtggaatcgtctactcacgca
	(Aacs)	Reverse	taaagggcgactctgtcgttc
8.	ATP citrate lyase (Acl1)	Forward	tggatgccacagctgactac
		Reverse	ggttcagcaaggtcagcttc
9.	3-hydroxy-3-methylglutaryl-	Forward	aactggtgcagaaatctctagc
	Coenzyme A synthase 1 (Hmgcs1)	Reverse	ggttgaatagctcagaactagcc
10.	Phosphomevalonate kinase	Forward	cctatggggctgtgatacaga
	(Pmvk)	Reverse	tctccgtggttctcaatgacc
11.	Mevalonate kinase ( <i>Mvk</i> )	Forward	ggtgtggtcggaacttccc
		Reverse	ccttgagcgggttggagac
12.	Mevalonate (diphospho)	Forward	ctcagcctcagctataaggtgc

decarboxylase ( <i>Mvd</i> )  13. Isopentenyl-diphosphate delta isomerase ( <i>Idi1</i> )  14. Farnesyl diphosphate farnesyl transferase 1 ( <i>Fdft1</i> )  15. Squalene epoxidase ( <i>Sqle</i> )  16. Lanosterol synthase ( <i>Lss</i> )  17. Cytochrome P450, family 51  Reverse gagccacttcggagaggg Forward agcttctagcggagatgt Reverse cagcaactattggtgaaa  Forward gtttgaagaccccatagtt Reverse atatccgagaaggcaga Reverse cgtagcagtaactcaacacacacacacacacacacacaca	gta acaacc tggtg tttag tgtcac cgaac cctat ggca cagaag
isomerase ( <i>Idi1</i> )  14. Farnesyl diphosphate farnesyl transferase 1 ( <i>Fdft1</i> )  15. Squalene epoxidase ( <i>Sqle</i> )  16. Lanosterol synthase ( <i>Lss</i> )  17. Cytochrome P450, family 51  Reverse cagcaactattggtgaaa geverse cacatctacgttctctggc ataacgaagtgagagagagagagagagagagagagagaga	acaacc tggtg ettag tgtcac egaac ectat ggca cagaag
14.Farnesyl diphosphate farnesyl transferase 1 (Fdft1)Forwardgtttgaagaccccatagtransferase 1 (Fdft1)15.Squalene epoxidase (Sqle)Forwardataagaaatgcggggatransferase 1 (Ess)16.Lanosterol synthase (Lss)Forwardgggaaggactcaacacacacacacacacacacacacacac	tggtg ettag tgtcac egaac ectat ggca eagaag
transferase 1 ( <i>Fdft1</i> )  15. Squalene epoxidase ( <i>Sqle</i> )  16. Lanosterol synthase ( <i>Lss</i> )  17. Cytochrome P450, family 51  Reverse cacatctacgttctctggc ataagaaatgcggggat Reverse cacatctacgttctctggc ataagaaatgcggggat Reverse cacatctacgtctctggc ataagaaatgcggggat Reverse cacatctacgtctctggc ataagaagactgaaggcaggat Reverse cgtagcagtaactcatgg	ettag tgtcac egaac ectat ggca eagaag
15. Squalene epoxidase ( <i>Sqle</i> )  Reverse atatccgagaaggcagd  16. Lanosterol synthase ( <i>Lss</i> )  Forward gggaaggactcaacacd Reverse cgtagcagtaactcatgg  17. Cytochrome P450, family 51  Forward aacgaagacctgaatgd	tgtcac cgaac cctat ggca cagaag
Reverse atatccgagaaggcagc  16. Lanosterol synthase ( <i>Lss</i> ) Forward gggaaggactcaacacc Reverse cgtagcagtaactcatgg  17. Cytochrome P450, family 51 Forward aacgaagacctgaatgc	cgaac cctat ggca cagaag
16. Lanosterol synthase ( <i>Lss</i> ) Forward gggaaggactcaacacacacacacacacacacacacacac	cctat ggca cagaag
Reverse cgtagcagtaactcatgg 17. Cytochrome P450, family 51 Forward aacgaagacctgaatgd	ggca cagaag
17. Cytochrome P450, family 51 Forward aacgaagacctgaatgo	cagaag
(Cyp51) Reverse gtgggctatgttaaggcc	
18. Transmembrane 7 superfamily Forward ggcctttgcgaccactctd	
member 2( <i>Tm7sf2</i> ) Reverse gttcagctcccgtccaag	aaa
19. Methylsterol monooxygenase 1 Forward tcatcggaattgtgcttttg	tgt
(Sc4mol) Reverse cagcgggttgagaggaa	atatc
20. NAD(P) dependent steroid dehydrogenase-like ( <i>Nsdhl</i> )	gact
Reverse ggtcccttgggccgaaaa	at
21. Hydroxysteroid (17-beta) Forward tctctgccatgtggataac	ccc
dehydrogenase 7 ( <i>Hsd17b7</i> ) Reverse ggtcggtagcgtatttgga	aag
22. Phenylalkylamine Ca2 <sup>+</sup> Forward actggccttgtgctggttt	
antagonist (emopamil) binding Reverse tccatacagacgacgaa protein ( <i>Ebp</i> )	agctg
23. Sterol-C5-desaturase ( <i>Sc5d</i> ) Forward ggggttacagcaaacto	ctaca
Reverse ggtgcaggcccctatga	
24. 7-dehydrocholesterol Forward cagatttctgccaggtta	
reductase ( <i>Dhcr7</i> ) Reverse agaaccaggataagag	* * * *
25. 24-dehydrocholesterol Forward gcacaggcatcgagtca	
reductase ( <i>Dhcr24</i> ) Reverse cagggcacggcataga:	
26. Acetyl-Coenzyme A Forward tccattcaaaacatgggg	
acetyltransferase 2 ( <i>Acat2</i> ) Reverse tcagcctggaagaggtc	
27. Thioredoxin reductase 1 Forward cccacttgccccaactgt	
(Txnrd1/TRXR1) Reverse gggagtgtcttggaggga	
28. NAD(P)H dehydrogenase, Forward ttctctggccgattcagag	
quinone 1 (Nqo1) Reverse ggctgcttggagcaaaa	
29. Heme oxygenase 1 Forward cacgcatatacccgctad	•
(Hmox1/HO-1) Reverse ccagagtgttcattcgag	
30. Glutathione reductase (Gsr) Forward gacacctcttccttcgact	
Reverse cccagcttgtgactctcca	ас
31. Glutathione peroxidase 1 Forward gtccaccgtgtatgccttc	et
(Gpx1) Reverse tctgcagatcgttcatctcg	
32. Glutathione peroxidase 2 Forward gcctcaagtatgtccgad	ectg
( <i>Gpx2</i> ) Reverse ggagaacgggtcatcat	
33. Superoxide dismutase 2 Forward gcggtcgtgtaaacctca	

	(Sod2)	Reverse	ccagagcctcgtggtacttc
34.	Superoxide dismutase 3	Forward	ctgaggacttcccagtgac
	(Sod3)	Reverse	ggtgagggtgtcagagtgt

# Supplementary Table 2: List of primers for human gene expression analyses

SI. No.	Gene Name	Forward/ Reverse	Sequence (5'-3')
1.	Glyceraldehyde-3-phosphate	Forward	ggagcgagatccctccaaaat
	dehydrogenase(GAPDH)	Reverse	ggctgttgtcatacttctcatgg
2.	Euchromatic histone lysine N-	Forward	gggcgggaaaatcacctcc
	methyltransferase 2 (EHMT2/G9a)	Reverse	cactcatgcggaaatgctgtat
3.	Sirtuin 6 (SIRT6)	Forward	cccacggagtctggaccat
		Reverse	ctctgccagtttgtccctg
4.	Low density lipoprotein	Forward	gttcagatgacgcggatgaaa
	receptor-related protein 2 ( <i>LRP2</i> )	Reverse	tcacagtcttgatcttggtcaca
5.	ATP-binding cassette, sub-	Forward	ttcccgcattatctggaaagc
	family A (ABC1), member 1 (ABCA1)	Reverse	caaggtccatttcttggctgt
6.	ATP binding cassette	Forward	cgtgcgctttgtgctgttt
	subfamily G member 1 (ABCG1)	Reverse	ccactgtaggtacgtggggat
7.	Acetoacetyl-CoA synthetase	Forward	ggcagtcggctcaactatg
	(AACS)	Reverse	acaacccgatctcctttcttca
8.	3-hydroxy-3-methylglutaryl-	Forward	gatgtgggaattgttgccctt
	Coenzyme A synthase 1 (HMGCS1)	Reverse	attgtctctgttccaacttccag
9.	Mevalonate (diphospho)	Forward	ggaccggatttggctgaatg
	decarboxylase (MVD)	Reverse	cccatcccgtgagttcctc
10.	24-dehydrocholesterol	Forward	cactgtctcactacgtgtcgg
	reductase (DHCR24)	Reverse	ccagccaatggaggtcagc

# **Supplementary Table 3: List of primers for ChIP assays**

SI. No.	Gene Name		Sequence
	For G9a	(or Sirt6) binding a	and Sequential ChIP
1	l m2	Forward	aggcacaggtcgaggatct
1.	Lrp2	Reverse	ctgccctccagtctcagttt
2	4000	Forward	tgctaccgtttcgttcactg
2.	Aacs	Reverse	gcaggtttccgaacaaagag
2	Hmgcs1	Forward	cattggcaggcttgttctc
3.		Reverse	gatccgctttcagccaatg
4	Mvd	Forward	aaaagcaactcccattcactg
4.		Reverse	tggctgttgaatggcttagag
_	Dhcr24	Forward	ctcccactctagggaatcca
5.		Reverse	cagtgctattgcaggtgttca
	For S	Sirt6 binding and Ti	me Kinetics ChIP
6.	About	Forward	agtccggagtttcccgttt
О.	Abca1	Reverse	agcagaaagcacgtggagac
7.	Abcg1	Forward	ccgactaggccatcttttga
		Reverse	agctaatggatggatcacagg
		For β-CATENIN	binding
8.	Ehmt2	Forward	atgtcctcatccgctgaaag
Ο.		Reverse	gtctccggctccatcttttt
9.	Sirt6	Forward	agttcagcagctcacacagg
9.		Reverse	gaacttggaagctccgtttg

### References

409 1. R. B. de Medeiros. Sequential chromatin immunoprecipitation assay and 410 analysis. *Methods Mol Biol* **791**:225-37 (2011).

2. A. D. Truax, and S. F. Greer. ChIP and Re-ChIP assays: investigating interactions between regulatory proteins, histone modifications, and the DNA sequences to which they bind. *Methods Mol Biol* **809**:175-88 (2012).

- 416 3. A. R. Leventhal, W. Chen, A. R. Tall, & I. Tabas. Acid Sphingomyelinase-417 deficient Macrophages Have Defective Cholesterol Trafficking and Efflux. *J Biol Chem*
- 418 **276**(48):44976-83 (2001).