

Rifabutin is bactericidal against intracellular and extracellular forms of Mycobacterium abscessus

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29 ABSTRACT

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31 Mycobacterium abscessus is increasingly recognized as an emerging opportunistic pathogen causing 32 severe lung diseases. As it is intrinsically resistant to most conventional antibiotics, there is an unmet medical need for effective treatments. Repurposing of clinically validated pharmaceuticals represents 33 34 an attractive option for the development of chemotherapeutic alternatives against *M. abscessus* infections. In this context, rifabutin (RFB) has been shown to be active against *M. abscessus* and has 35 raised renewed interest in using rifamycins for the treatment of *M. abscessus* pulmonary diseases. 36 37 Herein, we compared the *in vitro* and *in vivo* activity of RFB against the smooth and rough variants of M. abscessus, differing in their susceptibility profile to several drugs and physiopathologial 38 characteristics. While the activity of RFB is greater against rough strains than in smooth strains in 39 40 vitro, suggesting a role of the glycopeptidolipid layer in susceptibility to RFB, both variants were equally susceptible to RFB inside human macrophages. RFB treatment also led to a reduction in the 41 42 number and size of intracellular and extracelluar mycobacterial cords. Furthermore, RFB was highly effective in a zebrafish model of infection and protected the infected larvae from M. abscessus-43 44 induced killing. This was corroborated with a significant reduction in the overall bacterial burden, as well as decreased numbers of abscesses and cords, two major pathophysiological traits in infected 45 zebrafish. This study indicates that RFB is active against *M. abscessus* both in vitro and in vivo, further 46 47 supporting its potential usefulness as part of combination regimens targeting this difficult-to-treat mycobacterium. 48

49 INTRODUCTION

50

51 Nontuberculous mycobacteria (NTM) are environmental mycobacteria. Among all NTM, Mycobacterium avium and Mycobacterium abscessus represent the most frequent pathogens 52 associated with pulmonary disease (1). *M. abscessus* is a rapidly growing NTM of increasing clinical 53 54 significance, particularly in cystic fibrosis (CF) patients (2). In CF patients, infection with *M. abscessus* correlates with a more rapid decline in lung function and can represent an obstacle to subsequent 55 56 lung transplantation (3–5). From a taxonomical view, the species currently comprises three 57 subspecies: *M. abscessus* subsp. *abscessus* (designated hereafter *M. abscessus*), *M. abscessus* subsp. bolletii (designated hereafter M. bolletii) and M. abscessus subsp. massiliense (designated hereafter 58 59 *M. massiliense*) (6). These subspecies exhibit different clinical outcomes and drug susceptible profiles 60 to antibiotic treatments (7).

M. abscessus strains can exhibit either a smooth (S) or rough (R) morphotype as a consequence 61 62 of the presence or absence, respectively, of bacterial surface glycopeptidolipids (GPL) (1, 8–10). These morphological distinctions are associated with important physiological differences. S variants 63 64 are more hydrophilic than R variants, enabling increased sliding motility and the capacity to form biofilms (8, 9, 11), while the aggregative R variants possess a high propensity to produce large 65 bacterial cords (11, 12). While S and R variants can be viewed as two representatives of the same 66 67 isolate, which can co-exist and evolve differently in response to host immunity, they express different pathophysiological traits (10). S variants are typically less virulent than the R variants (11, 13, 14), the 68 latter being more frequently associated with severe lung diseases and persisting for years in CF 69 70 patients (3, 5). Importantly, an S-to-R transition within the colonized host (5, 15) is linked to genetic 71 polymorphisms within the GPL biosynthetic/transport locus (15, 16). Moreover, differences in the 72 susceptibility to drug candidates have been identified between S and R variants (17, 18), highlighting the need for the improved evaluation of new compounds/drug regimens against both morphotypes. 73

Treatment of *M. abscessus* lung disease remains particularly challenging, largely due to intrinsic resistance to wide panel of antimicrobial agents, including most antitubercular drugs such as rifampicin (RIF) (19–22). The extensive resistome of *M. abscessus* results from a low permeability of the cell wall, absence of drug-activating systems, induction of efflux pumps and production of a wide panel of drug-modifying enzymes (19, 22, 23). In addition, mutations in genes encoding drug targets 79 can result in acquired drug resistance further complicating therapy (1, 24). Treatment of infections 80 caused by *M. abscessus* require prolonged courses of multiple antibiotics, usually combining a macrolide (azithromycin or clarithomycin), a β -lactam (imipenem or cefoxitin) and an aminoglycoside 81 (amikacin) (25, 26, 27). Additional drugs, such as tigecycline or clofazimine, are often added to 82 83 strengthen the regimen, particularly in response to toxic side effects or unsatisfactory clinical response (28). Despite intensive chemotherapy, treatment success rates typically remain around 25-84 40% in the case of macrolide resistance, which occurs in at least 40-60% of clinical isolates (29). 85 86 Therefore, there is an urgent clinical need for new drug regimens with improved efficacy (30). While the current drug pipeline against *M. abscessus* remains poor, it has recently been fueled with the 87 88 discovery of several active hits and the development of repurposed drugs (24). Among the latter, screening of libraries of approved pharmaceuticals revealed that rifabutin (RFB), a rifamycin related 89 to the poorly active rifampicin (RIF), shows activity against *M. abscessus* (31, 32). RIF, along with 90 many other rifamycins, is inactivated by the ADP-ribosyltransferase (Arr_{Mab}) encoded by MAB_0591, 91 which ribosylates the drug at the C23 hydroxyl position (33). RFB has also been reported to be as 92 93 active as clarithromycin in immune-compromized NOD/SCID mice infected with *M. abscessus* (34). However, most studies on RFB have been carried out on either S or R variants (when reported), 94 rendering results sometimes difficult to interpret and/or to compare. Due to the co-existence of S 95 96 and R variants in patients (15) and the presence of each variant in different compartments (S residing 97 mostly in macrophages and R growing also in the form of intra- or extracellular cords), it is essential 98 to address the activity of RFB on isogenic S/R pairs in both *in vitro* and *in vivo* studies.

99 The present study aimed to describe and compare the activity of RFB against S and R *M*. 100 *abscessus* complex strains *in vitro* and *ex vivo* in a macrophage infection model. Due to the 101 importance of cording, considered as a marker of severity of the infection with the R variant, we also 102 investigated the efficacy of RFB in a zebrafish model of infection.

103 MATERIALS AND METHODS

104

Mycobacterial strains and growth conditions. *M. abscessus* CIP104536^T, *M. bolletii* CIP108541^T and 105 *M. massiliense* CIP108297^T reference strains and clinical isolates from CF and non-CF patients were 106 reported previously (35, 36). Strains were routinely grown and maintained at 30°C in Middlebrook 107 7H9 broth (BD Difco) supplemented with 0.05% Tween 80 (Sigma-Aldrich) and 10% oleic acid, 108 albumin, dextrose, catalase (OADC enrichment; BD Difco) (7H9^{T/OADC}) or on Middlebrook 7H10 agar 109 (BD Difco) containing 10% OADC enrichment (7H10^{OADC}) and in the presence of antibiotics, when 110 required. For drug susceptibility testing, bacteria were grown in Cation-Adjusted Mueller-Hinton 111 Broth (CaMHB; Sigma-Aldrich). RFB was purchased from two independent commercial sources 112 (Adoog Bioscience and Selleckchem) and dissolved in DMSO. 113

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115 Drug susceptibility testing. The minimal inhibitory concentrations (MIC) were determined according to the CLSI guidelines (37). The broth micro-dilution method was used in CaMHB with an inoculum of 116 5x10⁶ CFU/mL in exponential growth phase. The bacterial suspension was seeded in 100 µL volumes 117 118 in all of the wells of a 96-well plate, except for the first column, to which 198 µL of the bacterial suspension was added. In the first column, 2 μL of drug at its highest concentration was added to the 119 first well containing 198 µL of bacterial suspension. Two-fold serial dilutions were then carried out 120 121 and the plates were incubated for 3-5 days at 30°C. MICs were recorded by visual inspection. Assays 122 were completed in triplicate in three independent experiments.

123

Growth inhibition kinetics. To monitor growth inhibition of *M. abscessus* CIP104536^T S and R, 96-well plates were set-up as for MIC determination and serial dilutions of the bacterial suspensions exposed to increasing concentrations of RFB were plated on LB agar plates after 0, 24, 48 and 72 hrs. Colonyforming units (CFUs) were counted after 4 days of incubation at 30°C. Results from each drug concentration are representative of at least 2 independent experiments.

129

130 **Cytotoxicity assay.** THP-1 cells were differentiated with PMA for 48 hrs and exposed to decreasing 131 concentrations of either RFB or RIF (starting at 200 μ g/mL) for an additional 72 hrs at 37°C with 5% 132 CO₂. Following incubation, 10% (vol/vol) resazurin dye was added to each well and left to incubate for 4 hrs at 37°C and 5% CO₂. Data was acquired using a fluorescent plate reader (excitation 540 nm,
emission 590 nm). DMSO was included as a negative control, while SDS was included as a positive
control.

136

Intracellular killing assay. Human THP-1 monocytes were grown in RPMI medium supplemented with 137 10% Fetal bovine serum (Sigma Aldrich) (RPMI^{FBS}) and incubated at 37°C in the presence of 5% CO₂. 138 Cells were differentiated into macrophages in the presence of 20 ng/mL Phorbol Myristate Acetate 139 (PMA) in 24-well flat-bottom tissue culture microplates (10⁵ cells/mL) and incubated for 48 hrs at 140 37°C with 5% CO₂. Infection with clinical isolates or *M. abscessus* harbouring pTEC27 fluorescent 141 tdTomato was carried out at 37°C in the presence of 5% CO₂ for 3 hrs at a MOI 2:1. After extensive 142 washing with 1X phosphate buffered saline (PBS), cells were incubated with RPMI^{FBS} containing 250 143 μg/mL amikacin for 2 hrs and washed again with PBS prior to the addition of 500 μL RPMI^{FBS} 144 containing DMSO (negative control) or 500 µL RPMI^{FBS} containing 50 µg/mL of RIF or AMK, or 12.5 145 μg/mL of RFB. Macrophages were washed with PBS and lysed with 100 μL of 1% Triton X-100 at 146 required time points. Serial dilutions of macrophage lysates were plated onto LB agar plates and 147 148 colonies were counted to determine intracellular CFUs.

149

150 **Microscopy-based infectivity assays.** Monocytes were differentiated into macrophages (THP-1) in the presence of PMA and were grown on coverslips in 24-well plates at a density of 10⁵ cells/mL for 151 48 hrs at 37°C with 5% CO₂ prior to infection with Tdtomato expressing *M. abscessus* for 3 hrs at a 152 MOI of 2:1. After washing and AMK treatment to remove the extracellular bacilli, macrophages were 153 154 exposed to DMSO (negative control), or 50 µg/mL RIF or AMK, or 12.5 µg/mL RFB, and fixed at 0, 1 155 and 3 days post-infection with 4% paraformaldehyde in PBS for 20 min. Cells were then 156 permeabilized using 0.2% Triton X-100 for 20 min, blocked with 2% BSA in PBS supplemented with 0.2% Triton X-100 for 20 min, incubated with anti-CD63 antibodies (Becton Dickinson); dilution 157 1:1000) for 1 hr and with an Alexa Fluor 488-conjugated anti-mouse secondary antibody (Molecular 158 Probes, Invitrogen). After 5 min of incubation with DAPI (dilution 1:1000), cells were mounted onto 159 160 microscope slides using Immu-mount (Calbiochem) and examined with an epifluorescence 161 microscope using a 63X objective. The average proportion of macrophages containing fewer than <5, 162 5-10, or >10 bacilli were quantified using Zeiss Axio-vision software. Images were acquired by

163 focusing on combined signals (CD63 in green and red fluorescent M. abscessus) and captured on a Zeiss Axio-imager confocal microscope equipped with a 63X oil objective and processed using Zeiss 164 165 Axiovision software. Quantification and scoring of the numbers of bacilli present within macrophages 166 were performed using ImageJ. Equal parameters for the capture and scoring of images were consistently applied to all samples. For each condition, approximately 1000 infected macrophages 167 168 were analyzed. The presence of the intra- or extracellular cords within or among the macrophages 169 infected with the R morphotype strain were treated in the presence of DMSO, RIF, RFB or AMK at the 170 concentrations previously described, counted and imaged using confocal microscopy.

171

Assessment of RFB efficacy in infected zebrafish. Experiments in zebrafish were conducted according 172 to the Comité d'Ethique pour l'Expérimentation Animale de la Région Languedoc Roussillon under the 173 174 reference CEEALR36-1145. Experiments were performed using the *golden* mutant (38). Embryos were obtained and maintained as described (14). Embryo age is expressed as hours post fertilisation (hpf). 175 Red fluorescent *M. abscessus* CIP104536^T (R) expressing tdTomato were prepared and microinjected 176 in the caudal vein (2-3 nL containing ≈100 bacteria/nL) in 30 hpf embryos previously dechorionated 177 178 and anesthetized with tricaine, as described earlier (39). The bacterial inoculum was checked a*posteriori* by injection of 2 nL in sterile PBS^T and plating on 7H10^{OADC}. Infected embryos were 179 transferred into 24-well plates (2 embryos/well) and incubated at 28.5°C to monitor kinetics of 180 181 infection and embryo survival. Survival curves were determined by counting dead larvae daily for up to 12 days, with the experiment concluded when uninfected embryos started to die. RFB treatment 182 of infected embryos and uninfected embryos was commenced at 24 hpi (hours post-infection) for 4 183 184 days. The drug-containing solution was renewed daily. Bacterial loads in live embryos were 185 determined by anesthetising embryos in tricaine as previously described (40), mounting on 3% (w/v) 186 methylcellulose solution and taking fluorescent images using a Zeiss Axio Zoom.V16 coupled with an 187 Axiocam 503 mono (Zeiss). Fluorescence Pixel Count (FPC) measurements were determined using the 'Analyse particles' function in ImageJ (39). Bacterial cords were identified based on the size and 188 shape of fluorescent bacteria within the live zebrafish embryo, vastly exceeding the surrounding size 189 190 and shape of neighbouring cells. All experiments were completed at least three times independently.

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- 192

193 **Overexpression of** *MAB***<u>1409c in </u>***M. abscessus***.**

Overexpression was achieved by PCR amplification of MAB 1409c (tap) in fusion with an HA tag using 194 195 genomic DNA and the forward primer (5'- gaga<u>CAATTG</u>CCATGTCCACTCCGACGGCGGATTC-3'; MfeI) 196 (5'and reverse primer gagaGTTAACCTAAGCGTAATCTGGAACATCGTATGGGTACCGAGTTGGTTCCTTGTCGGGCT-3'; Hpal). The 197 198 amplified product was digested with Mfel/Hpal and ligated into the Mfel/Hpal-restricted pMV306 199 integrative vector to generate pMV306-MAB 1409c-HA where MAB 1409c-HA is under the control of 200 the *hsp60* promoter. The construct was sequenced and electroporated in *M. abscessus* S and R.

201

Selection of resistant *M. abscessus* mutants and target identification. Exponentially growing *M. abscessus* CIP104536^T R cultures were plated on LB agar containing either 25 or 50 μ g/mL RFB. After one week of incubation at 37°C, four individual colonies from each RFB concentration were selected, grown in CaMHB, individually assessed for MIC determination and scored for resistance to RFB. Identification of SNPs in the resistant strains was completed by PCR amplification using *rpoB_f* 5'-TCAGTGGGGCTGGTTAG -3' and *rpoB_r* 5'-AAAACATCGCAGATGCGC-3' to produce a 3541 bp amplicon for full coverage sequencing of the *rpoB* gene.

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Western blotting. Bacteria were harvested, resuspended in PBS, and disrupted by bead-beating with 1-mm diameter glass beads. The protein concentration in the lysates was determined and equal amounts of proteins (100 μ g) were subjected to SDS/PAGE. Proteins were transferred to a nitrocellulose membrane. For detection of Tap-HA and KasA (loading control), the membranes were incubated for 1 hr with either the rat anti-HA or rat anti-KasA antibodies (dilution 1:2000), washed, and subsequently incubated with goat anti-rat antibodies conjugated to HRP (Abcam, dilution 1:5000). The signal was revealed using the ChemiDoc MP system (Bio-Rad).

217

Statistical analyses. Statistical analyses were performed on Prism 5.0 (Graphpad) and detailed for each figure legend. * $P \le 0.05$, ** $P \le 0.01$, *** $P \le 0.001$, **** $P \le 0.0001$.

220 **RESULTS**

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222 Rough *M. abscessus* is more susceptible to RFB treatment than smooth *M. abscessus in vitro*. Exposure of exponentially-growing *M. abscessus* CIP104536^T S and R isogenic variants to increasing 223 concentrations of RFB, starting at 25 µg/mL for S and 6.25 µg/mL for R, resulted in a noticeable 224 225 growth inhibition (Fig. 1). At the lowest concentration, the CFUs at 72 hrs post-treatment remained 226 comparable to those of the inoculum, suggestive of a bacteriostatic effect. However, the highest RFB 227 concentrations for both S (200 µg/mL) and R (50 µg/mL) variants were accompanied by 1.81 and 2.47 Log reduction in the CFU counts at 72 hrs post-treatment, respectively (Fig. 1). While similar 228 229 bactericidal effects of RFB were observed against both variants, this was achieved with lower 230 concentrations of RFB against the R variant relative to the isogenic S variant. Overall, RFB at concentratios of 12.50 µg/mL (For R) resulted in a killing effect comparable to the one of imipenem 231 (IPM) used at the MIC (16 μ g/mL), known as an active β -lactam drug against *M. abscessus* (41) (Fig. 232 233 1).

To confirm the differences in the susceptibility to RFB, we determined the MIC of the CIP104536^T 234 S and R variants in CaMHB. Table 1 clearly shows that the S strain is 4-fold more resistant than its R 235 236 counterpart. However, both variants were similarly resistant to other rifamycins (RIF, RPT and RFX), in 237 agreement with previous studies (32, 34). Our MIC values, obtained in repetitive experiments with 238 two different commercial sources of RFB, were higher than those reported earlier (32, 34), but 239 comparable to values reported in another study (42). Consistently with other studies (31), we also noticed that the MIC values were dependent on the culture medium (Table S1). Interestingly, MICs of 240 RFB against S and R strains were lower in Middlebrook 7H9 as compared to CaMHB but this effect 241 242 was lost when supplementing the medium with OADC enrichment. In contrast the S and R strains 243 displayed equal susceptibility levels to RFB in Sauton's medium.

To investigate the relationship between RFB activity and GPL production, drug susceptibility was assessed in CaMHB using the GPL-deficient $\Delta mmpL4b$ mutant, generated in the S background of the type strain CIP104536^T, and its complemented counterpart (14, 43, 44). The *mmpL4b* gene encodes the MmpL4b transporter which participates in the translocation of GPL across the inner membrane (43, 45). Mutations in this gene are associated with loss of GPL and acquisition of a R morphotype

249 (13, 43). The parental S strain and, to a lesser extent the $\Delta mmpL4b$ -complemented strain, showed 250 reduced susceptibility to RFB (MIC 32-64 µg/mL) than the *M. abscessus* R strain and the GPL-deficient 251 $\Delta mmpL4b$ mutant (MIC 16 µg/mL) (Table 1). The MIC results are in agreement with the growth 252 inhibition kinetics (Fig. 1) and suggest that the outer GPL layer influences the activity of RFB.

253 *M. abscessus* possesses numerous potential drug efflux systems (45), including MAB 1409c, a 254 homolog of Rv1258c, previouly reported to mediate efflux of RIF in *M. tuberculosis* (46). We thus 255 addressed whether overexpression of MAB 1409c induces resistance to RFB in *M. abscessus*. 256 MAB 1409c was cloned in frame with a HA-tag in the integrative pMV306. The resulting construct pMV306-MAB 1409c-HA was introduced in both S and R variants and the expression of MAB 1409c 257 258 was confirmed by Western blot analysis using anti-HA antibodies (Fig. S1). Drug susceptibility 259 assessment indicated a 4-fold upshift in the MIC of RFB against the R strain carrying pMV306-MAB 1409c-HA while no changes in the MIC were observed with the S strain overproducing 260 261 MAB 1409c (Table 1). This suggests that increasing expression of MAB 1409c in the R variant is likely 262 to mediate efflux of RFB, leading to reduced susceptibility to the drug.

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264 RFB is active against S and R M. abscessus isolates in vitro. The activity of RFB was next tested using a set of clinical strains isolated from CF patients or non-CF patients. In general, the MIC of R strains 265 were 2 to 4 times lower than those of S strains, altough there were variations among the strains 266 (Table 2). Some R strains (10, 112, 179, 210) exhibited higher MIC values (100 µg/mL) than the 267 reference CIP104536^T R strain, while one S strain appeared particularly susceptible to RFB (*M*. 268 massiliense 120 with a MIC of 6.25 µg/mL). These differences between strains and S/R morphotypes 269 270 were not observed previously with BDQ and S and R variants were also equally sensitive to BDQ (47). 271 Overall, these results demonstrate that RFB is active against *M. abscessus*, including isolates from CF 272 patients, while R variants appear in general more susceptible to RFB than S variants, supporting previous findings (48). 273

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275 **Mutations in** *rpoB* confer resistance to RFB. Although rifamycin resistance mechanisms mediated by 276 mutations in the *rpoB* gene coding for the β -subunit of RNA polymerase have been widely described 277 for *M. tuberculosis*, this is not the case for *M. abscessus*. Therefore, to identify the mechanism of 278 resistance of RFB, a genetic approach involving the selection of spontaneous RFB-resistant mutants of *M. abscessus* followed by *rpoB* sequencing was applied. Four spontaneous strains were isolated in 279 280 the presence of 25 or 50 µg/mL RFB, exhibiting 4- to 8-fold increased resistance levels as compared 281 to the parental strain, respectively (**Table 3**). Sequencing analyses of *rpoB* identified several single nucleotide polymorphisms (SNPs) across four resistor mutants. In mutants 25.1 and 50.1, a C1339T 282 283 substitution was identified at position 447 (H447Y). Similarly, in mutant 25.2, a C1339G replacement 284 was found, resulting in an amino acid substitution at position 447 (H447D). Comparatively, in mutant 285 strain 50.2, another SNP (C1355T) occurred, leading to an amino acid change at position 452 (S452L). 286 A comparison of growth of different resistant strains on agar plates containing increasing 287 concentrations of RFB is shown in Fig. S2. Whereas RFB abrogated growth of the wild-type S and R strains, growth of all four resistors harbouring mutations at either positions 447 or 452 sustained 288 289 bacterial growth at 50 μ g/mL, confirming that mutations in *rpoB* confer resistance to RFB.

290

291 M. abscessus S and R strains are equally susceptible to RFB in macrophages. While RFB has been 292 shown to be active against *M. tuberculosis* in a macrophage infection model (49), this has not been 293 thoroughly investigated for *M. abscessus*. We thus compared the intracellular efficacy of RFB in THP-294 1 macrophages infected with either S or R variants. Firstly, the cytotoxicity of RFB and RIF against 295 THP-1 cells was investigated over a 3-days exposure period to either drug. Fig. S3, clearly shows that 296 RFB exerts significant cytotoxicity at concentration >25 μ g/ml and that the kinetic of macrophage 297 killing was more rapid with RFB than with RIF. Based on these results, all subsequent macrophage 298 studies were treated with 50 µg/mL RIF or 12.5 µg/mL RFB. AMK at 50 µg/mL was added as a positive 299 control. DMSO-treated macrophages were included as a negative control for intracellular bacterial 300 replication. At 0, 1 and 3 days post-infection (dpi), macrophages were lysed and plated to determine 301 the intracellular bacterial loads following drug treatment. Whereas the presence of DMSO or RIF failed to inhibit intramacrophage growth of *M. abscessus* S, exposure to RFB strongly decreased the 302 intracellular bacterial loads at 1 dpi, with this effect further exacerbated at 3 dpi (Fig. 2A). As 303 304 anticipated, treatment with RIF did not show any effect, in agreement with the poor activity of this 305 compounds in vitro (Table 1). Comparatively, AMK treatment resulted in a significantly reduced 306 intracellular growth rate in both *M. abscessus* S and R variants between 1 and 3dpi. Interestingly, the

307 RFB susceptibility profile for the S variant at 1 and 3 dpi was comparable to that of the R variant, with
308 a ~3 Log reduction in the CFU counts (Fig. 2A and B, respectively).

Macrophages were next infected with *M. abscessus* strains expressing Tdtomato and exposed to either DMSO, AMK, RIF or RFB, followed by staining with anti-CD63 and DAPI and observed under a confocal microscope. A quantitative analysis confirmed the marked reduction in the number of *M. abscessus* S-infected THP-1 cells treated with AMK and RFB at 1 and 3 dpi compared to RIF-treated cells or untreated control cells (Fig. 2C). A similar trend was observed when macrophages were infected with *M. abscessus* R (Fig. 2D).

Macrophages infected with the S variant were then classified into three categories based on their 315 bacterial burden: poorly infected (<5 bacilli), moderately infected (5-10 bacilli) and heavily infected 316 (>10 bacilli) macrophages. Cells containing bacilli were then individually observed under the 317 318 microscope and scored to one of the three categories. The quantitative analysis indicates that 319 exposure to RFB significantly reduces the percentage of S variant heavily infected THP-1 cells while 320 increasing the proportion of the poorly infected category, as compared to the untreated cells at 1 dpi (Fig. 2E). At 3 dpi, the effect of RFB was even more pronounced with 10% of the infected bacilli 321 322 belonging to the heavily infected category and more than 50% associated with the poorly infected category. Analysis performed on cells infected with the R variant generated a similar category profile, 323 although treatment with RFB was associated with a higher proportion of heavily infected 324 325 macrophages at 3 dpi with the R variant than with the S variant (Fig. 2F). Fig. 2G illustrates the 326 reduced number of *M. abscessus* S in infected THP-1 cells treated with RFB at 1 dpi, as compared to the untreated control cells (DMSO) or those treated with RIF or AMK. Collectively, these results 327 328 indicate that RFB enters THP-1 macrophages and similarly impedes bacterial replication of both M. 329 abscessus S and R variants.

330

RFB reduces the intramacrophage growth of clinical isolates. RFB has recently shown vast potential as an effective antibiotic for the treatment of *M. abscessus* infection in a NOD/SCID murine model (34). However, to date the efficacy of RFB has only been evaluated against a limited panel of *M. abscessus* clinical isolates within an infection setting. As such, we explored the activity of RFB against S and R clinical isolates of the *M. abscessus* complex with varying MIC values against RFB within THP-1 macrophages. In support of our previous findings in infected macrophages, RFB treatment (12.5 or 337 25 μg/mL) was very active against all *M. abscessus* subspecies within macrophages at 1 and 3 dpi 338 when compared to Day 0 and DMSO treatment (Fig. 3), irrespective of S and R morphotypes and the 339 corresponding MIC values (Table 2). Overall, these findings suggest that RFB is very effective against 340 intracellular clinical isolates and highlights the lack of direct correlation between MICs determined *in* 341 *vitro* and the intracellular activity of RFB.

342

Reduced intra- and extracellular cording by RFB treatment. An important phenotypic difference 343 between S and R morphotypes is that R morphotypes display increased bacterial aggregation. R bacilli 344 345 remain attached during replication, forming compact colonies containing structures that resemble cords on agar and in broth medium (8, 12, 14). Fig. 4A clearly shows that, upon infection with M. 346 abscessus R expressing TdTomato, the total number of cords per field was significantly reduced in the 347 348 presence of 50 µg/mL AMK or 12.5 µg/mL RFB when compared to 50 µg/mL RIF or DMSO alone. 349 Moreover, we observed intracellular cords that are capable of growing inside the macrophage as well 350 as in the extracellular milieu, which were easily observable at 3 dpi (Fig. 4B). As illustrated in Fig. 4C, treatment with AMK or RFB strongly impacted on both intra- and extracellular cords. While AMK 351 352 treatment severely reduced the number of both intra- and extracellular cords, this effect was almost completely abrogated with RFB at 3 dpi. Together, these results indicate that RFB is highly effective in 353 354 reducing *M. abscessus* cords, thought to affect the outcome of the infection.

355

356 **RFB treatment enhances protection of zebrafish infected with** *M. abscessus.* In vivo drug efficacy has previously been well described using the zebrafish model of infection (40, 47, 50). Initial 357 358 experiments indicated that RFB concentrations $\leq 100 \ \mu g/mL$ (final concentration in fish water) did not 359 interfere with larval development and was well tolerated in embryos when treatment was applied for 360 4 days with daily drug renewal (Fig. 5A). Higher concentrations of RFB, however, were associated 361 with rapid larval death. As such, only lower RFB doses ($\leq 100 \ \mu g/mL$) were used in subsequent studies. Red fluorescent tdTomato-expressing M. abscessus (R variant) was microinjected in the 362 caudal vein of embryos at 30 hrs post-fertilisation (hpf). RFB was directly added at 1 dpi to the water 363 containing the infected embryos, with RFB-supplemented water changed on a daily basis for 4 days. 364 365 Embryo survival was monitored and recorded daily for 12 days. No decrease in the survival rate was 366 observed in the presence of 5 μ g/mL RFB, however, a significant dose-dependent increase in the 367 survival of embryos exposed to 25 or 50 μ g/mL RFB was observed as compared to the untreated 368 group (Fig. 5B). When exposed to 50 μ g/mL RFB, the highest dose examined in this setting, nearly 369 80% of the treated embryos survived at 12 dpi, as compared to 40% of the untreated group. This 370 clearly indicates that RFB protects zebrafish from *M. abscessus* infection.

To test whether RFB exerts an effect on the bacterial burden in zebrafish, we quantified fluorescent pixel counts (FPC) (39). As expected, embryos treated with 50 µg/mL RFB had significantly decreased bacterial burdens at 2, 4 and 6 dpi when compared to the untreated group (**Fig. 5C**). These results were corroborated by imaging whole embryos, characterised by the presence of large abscesses and cords in the brain when left untreated and which were observed much less frequently in the RFB-treated animals despite the presence of single bacilli or small aggregated bacteria (**Fig. 5D**).

378

379 RFB treatment reduces abscess formation by M. abscessus in zebrafish. Virulence of M. abscessus R 380 variants in zebrafish are correlated with the presence of abscesses, particularly in the central nervous system (14, 39). To address whether the enhanced survival of RFB-treated fish is associated with 381 382 decreased abscess formation, the percentage of abscesses and cords were determined by monitoring abscesses and cords in whole embryos, as reported previously (14, 39). Extracellular cords can be 383 384 easily distinguished based on their serpentine-like shape and by their size, often greater as compared 385 to the size of the surrounding macrophages and neutrophils. Exposure of infected embryos to 50 386 µg/mL RFB was accompanied by a significant decrease in the proportion of embryos with cords (Fig. 6A) at 4 dpi, and the number of embryos with abscesses (Fig. 6B) at 4 and 6 dpi. This decrease in the 387 388 physiopathological signs of RFB-treated larvae correlates also with the FPC analysis and whole 389 embryo imaging (Fig. 6C and 6D). Overall, these results demonstrate that RFB reduces the 390 pathophysiology of *M. abscessus* infection in zebrafish larvae and protects them from bacterial 391 killing.

392 DISCUSSION

393

394 Treatment success of infections caused by M. abscessus is unacceptably low even upon prolonged, multidrug chemotherapy with a significant risk of severe toxic side effects. Although RIF is used as a 395 first-line drug against M. tuberculosis, it has no activity against M. abscessus. While ADP 396 397 ribosyltransferases can utilise both RIF and RFB as substrates (51), a lower catalytic efficiency with 398 RFB may explain its greater potency against *M. abscessus*. Our study supports and extends previous 399 investigations highlighting the potential of RFB against *M. abscessus in vitro* against a wide panel of 400 *M. abscessus* complex clinical isolates (31, 32, 52, 53). We found, however, that our MIC values were 401 higher than those observed in previous investigations (31, 32). In our study, following the Clinical and Laboratory Standard Institute (CLSI) guidelines, MIC were determined in CaMHB while Aziz et al. 402 403 showed that MIC values were 2- to 3-fold higher in CaMHB as compared to Middlebrook 7H9 (31), clearly implicating an effect of medium on RFB susceptibility testing. In line with these results, we 404 405 noticed important variations in the MIC values depending on the culture medium used for RFB susceptibility assessments. It is also noteworthy that the growth curve of the untreated S strain is 406 407 different from the one of the R strain, which is very likely linked to the highly aggregative surface properties typifying the R strain which, in contrast to the S strain, produces very clumpy and corded 408 409 cultures in broth medium (10, 39, 54). As a consequence, colonies on agar plates are very likely 410 emerging from aggregated bacteria rather than individual bacilli, explaining why the CFU counts were significantly lower in both cultures. Thus, the CFU counts of the R strain does not accurately reflect 411 the absolute number of living bacilli in the culture. We also selected RFB-resistant mutants and 412 413 identified mutations in rpoB, known as the primary target of rifampicin in M. tuberculosis (55). 414 Interestingly, the mutations identified are part of the rifampicin-resistance-determining region 415 (RRDR), a 81-bp central segment corresponding to codons 426 to 452 in *M. tuberculsosis* that 416 harbours the vast majority of *rpoB* mutations associated with resistance to RIF (55). Noteworthingly, S452L corresponds to one of the most frequently mutated coding region in the rpoB gene in M. 417 418 tuberculosis (S450L replacement) (55). Together, these results suggest RpoB is very likely the target of RFB in M. abscessus. 419

420 Among the various studies reporting the activity of RFB against *M. abscessus in vitro*, very few 421 discriminated the activity of RFB against the S or R morphotypes. Herein, we found that the type

strain CIP104536^T S was reproducibly more resistant to RFB than its R counterpart. Supporting these 422 results, deletion of *mmpL4b* in the S genetic background, resulting into an R morphotype lacking GPL 423 (13, 43), increased susceptibility to RFB. Conversely, functional complementation of the mmpL4b 424 mutant, restoring the S morphotype and GPL production (13, 43), partially rescued the higher MIC. 425 426 This highlights the influence of the outermost GPL layer on susceptility to RFB. Previously, the activity 427 of other inhibitors have been shown to be dependent on the presence or absence of GPL in M. 428 abscessus (17, 18). A logical explanation is that the GPL layer protects the bacilli from the penetration 429 of drugs. The absence of GPL may enhance the permeability of the cell wall and accumulation of the 430 drug inside the bacteria. However, one cannot exclude the possibility that MmpL4b, like other MmpL transporters, can act as an efflux pump (56–58) and may participate in the extrusion of RFB in M. 431 abscessus S, resulting in higher MIC. The implication of efflux pumps in resistance to RFB has been 432 433 investigated, whereby the overexpression of MAB 1409c (a homologue of the *M. tuberculosis* 434 Rv1258c) resulted in increased resistance to RFB in the R variant of *M. abscessus*. This effect was not 435 observed in the S strain overexpressing MAB 1409c, presumably because of the already elevated MIC of the parental S strain towards RFB. However, while the increased susceptibility of the R strain 436 437 as compared to the S strain was true with respect to the type strain, this was not observed for all clinical strains tested. The heterogeneity of the clinical strains in response to RFB treatment cannot 438 be simply explained by the presence or absence of GPL, but may also include additional determinants 439 of resistance to RFB (52), such as differences in the expression level of Arr_{Mab} or the expression of Rox 440 441 monooxygenases, known to inactivate RIF in other bacterial species as proposed earlier (59). This, 442 however, requires further investigation in follow-up studies.

443 One unanticipated finding from this study relies on the fact that, although S and R variants 444 respond differently to RFB treatment in vitro, this was not the case against the intracellularly-residing 445 *M. abscessus.* We found that, using a macrophage model of infection, the isogenic S and R type 446 strains responded equally well to treatment with 12.5 μ g/mL RFB, largely exceeding the results obtained with AMK, a drug displaying weak intracellular activity (60). These observations are 447 448 reminiscent of other studies indicating that various naphtalenic ansamycins, including RIF, differ profoundly in their capacity to kill extracellular Staphylococcus aureus, albeit there were few 449 450 differences observed between them in promoting human macrophages to kill phagocytosed bacteria 451 (61). There is no simple explanation as why *M. abscessus* S is as efficiently killed as *M. abscessus* R

452 inside the cells. A plausible explanation may be that the stress response inside macrophages alters the composition/architecture of the cell wall of *M. abscessus*, thereby affecting the GPL layer and/or 453 454 permeability of the S variant. It has been shown that the GPL layer significantly influences the 455 hydrophobic surface properties (62), potentially impacting on the adhesion and the uptake of the 456 bacilli. Furthermore, electron microscopy observations revealed that the electron translucent zone 457 (ETZ) that fills the entire space between the phagosome and the bacterial surface relies on GPL 458 production in the S variant (11). Alternatively, RFB may directly induce the antimycobacterial activity 459 of the macrophage, which in turns translates into a rapid killing of the phagocytosed bacteria, 460 regardless of their morphotype. Overall, these results suggest that the MIC values of RFB are not indicative of the intraphagocytic killing of *M. abscessus* and highlights the importance of testing the 461 efficacy of drugs in a macrophage infection model. 462

Cords and abscesses are pathophysiological markers of *M. abscessus* infection, as revealed using 463 the zebrafish model of infection (40). In particular, extracellular cords, due to their size, prevent the 464 465 bacilli from being phagocytosed by macrophages and neutrophils, representing an important mechanism of immune evasion (14, 39). We demonstrate here that treatment of infected 466 467 macrophages was associated with reduced intra- and extracellular cording of the R variant. It is very likely that RFB prevents cording, as a consequence of the inhibition of bacterial replication/killing. 468 Cords are a hallmark of virulence of the R variant of *M. abscessus*, as emphasized by a deletion 469 470 mutant of MAB 4780, encoding a dehydratase, displaying a pronounced defect in cording and a highly attenuated phenotype in macrophages (63). Importantly, we observed also a significant 471 472 decrease in the number of embryos with cords following RFB treatment in infected zebrafish. It is 473 worth highlighting that in the presence of RFB, there is no change in the number of embryos with 474 cords between 2 and 4 dpi, implying that while RFB does not degrade or modify the bacterial cord 475 structure, it likely prevents the formation of additional cords. Moreover, the effect of RFB on cord 476 reduction is particularly interesting as it may prevent the subsequent formation of abscesses (14), considered as a marker of severity of the disease. Consistent with this hypothesis, a marked decrease 477 478 in abscess formation was observed in RFB-treated zebrafish embryos. Overall, this work supports the practicality of zebrafish as a pre-clinical model to evaluate in real-time the bactericidal efficacy of RFB 479 480 against *M. abscessus* infection in the sole context of innate immunity.

In summary, although there is a clear lack of bactericidal activity of drugs against *M. abscessus* (64), these findings support the high activity of RFB against *M. abscessus in vivo* and *in vitro*. Our results further emphasize the efficacy of RFB against both extracellular and intracellular forms of *M. abscessus*, both co-existing in infected patients, as well as a protective effect in an animal model of *M. abscessus* infection. In addition, we have provided further evidence that S and R variants are differentially susceptible to RFB, likely due to the GPL layer, however the MIC values are not predictive of intracellular drug efficacy.

488 Together with the fact that RFB is an FDA-approved drug that is already used to treat tuberculosis (66) and *M. avium* infections (67) with favourable pharmacological properties (68), our 489 490 data strengthen the view that RFB should be considered as a repurposing drug candidate for the 491 treatment of *M. abscessus* infections. Importantly, recent work has shown that RFB is synergistic in combinations with other antimicrobials such as clarithromycin, imipenem and tigecycline, and 492 493 significantly improves the activity of imipenem-tedizolid drug combinations (32, 48, 53, 65). Future studies are required to test whether these RFB combinations are effective against M. abscessus 494 pulmonary infections. 495

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- 502

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Table 1. Drug susceptibility/resistance profile of smooth and rough variants derived from the reference *M. abscessus* 104536^{T} strain to various rifamycins in CaMBH. MIC (µg/mL) were determined following the CLSI guidelines.

				MIC (μg/	mL)
Strain	Morphotype	RFB	RIF	RPT	RFX
CIP104536 (S)	S	64	>128	>128	>128
CIP104536 (R)	R	16	>128	>128	>128
ΔMAB_mmpL4b	R	16	>128	>128	>128
ΔMAB_mmpL4b_C	S	32	>128	>128	>128
CIP104536 (S) + pMV306- <i>MA</i>	B_1409c-HA S	64	>128	>128	>128
CIP104536 (R) + pMV306-MA	<i>B_1409c-HA</i> R	64	>128	>128	>128

736 RFB, Rifabutin; RIF, rifampicin; RPT, rifapentine; RFX, rifaximin.

737 Table 2. Comparison of the activity of RFB against clinical isolates from CF and non-CF patients. The

MIC (µg/mL) was determined in Cation-Adjusted Mueller-Hinton broth for different subspecies
 belonging to the *M. abscessus* complex. Results are from 3 independent experiments. RFB, rifabutin.

Strain	Morphotype	Source	RFB
M. abscessus			
CIP104536	S	Non-CF	50
3321	S	Non-CF	50
1298	S	CF	50
2587	S	CF	100
2069	S	Non-CF	100
CF	S	CF	25
2524	R	CF	25
2648	R	CF	25
3022	R	Non-CF	50
5175	R	CF	25
CIP104536	R	Non-CF	25
M. massiliense			
CIP108297	R	Addison Disease	50
210	R	CF	100
179	R	CF	100
CIP108297	S	Addison Disease	100
140	S	CF	50
185	S	CF	100
107	S	CF	50
122	S	CF	100
120	S	CF	6.25
212	S	CF	100
100	S	CF	100
111	S	CF	100
M. bolletii			
CIP108541	S	Non reported	100
114	S	CF	100
17	S	CF	50
116	S	CF	100
97	S	CF	100
112	R	CF	100
19	R	Non-CF	50
10	R	Non reported	100
108	R	CF	25

Table 3. Characterisitics of spontaneous RFB-resistant mutants of *M. abscessus*. MIC (μ g/mL) were determined in Cation-adjusted Mueller-Hinton broth. Resistant strains were derived from the rough *M. abscessus* CIP104536^T parental strain on Middlbrook 7H10 supplemented with either 25 or 50 μ g/mL RFB. Single nucleotide polymorphism identification in *rpoB* (*MAB_3869c*) and corresponding amino acid changes are also indicated. RFB, rifabutin.

	MIC (µg/ml)	Mutation in <i>rpoB</i>	
		SNP	AA change
CIP104536 ^T (R)	12.5	-	-
25.1	50	C1339T	H447Y
25.2	100	C1339G	H447D
50.1	50	C1339T	H447Y
50.2	50	C1355T	S452L

754 FIGURE LEGENDS

Figure 1. *In vitro* activity of rifabutin. *M. abscessus* CIP104536^T S (left panel) or R (right panel) was exposed either to 200, 100, 50, 25, 12.5 or 6.25 μ g/mL RFB or 16 μ g/mL IPM in CaMHB at 30°C. At various time points, bacteria were plated on LB agar and further incubated at 30°C for 4 days prior to CFU counting. Results are expressed as the mean of triplicates ± SD and are representative of two independent experiments. **P* ≤ 0.05, ** *P* ≤ 0.01.

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Figure 2. Intracellular activity of RFB on M. abscessus-infected THP-1 cells. (A) Macrophages were 761 infected with *M. abscessus* S-morphotype and (B) R-morphotype expressing tdTomato (MOI of 2:1) 762 763 for 3 hrs prior to treatment with RIF (50 μg/mL), AMK (50 μg/mL), RFB (12.5 μg/mL) or DMSO. CFU 764 were determined at 0, 1 and 3 dpi. Data are mean values \pm SD for three independent experiments. Data were analysed using a one-way ANOVA Kruskal-Wallis test. (C) Percentage of infected THP-1 765 766 macrophages at 0, 1 and 3 days post-infection after infection with *M. abscessus* S or (D) *M. abscessus* R. Data are mean values ± SD for three independent experiments. Data were analysed using a one-767 way ANOVA Kruskal-Wallis test. (E) Percentage of S-infected macrophage categories and (F) 768 percentage of R-infected macrophage categories infected with different numbers of bacilli (<5 bacilli; 769 5-10 bacilli and >10 bacilli). The categories were counted at 0 or at 1 and 3 days post-infection in the 770 absence of antibiotics or in the presence of RIF or AMK at 50 µg/mL, or RFB at 12.5 µg/mL. Values are 771 772 means ± SD from three independent experiments performed in triplicate. (G) Four immuno-773 fluorescent fields were taken at 1 day post-infection showing macrophages infected with M. 774 abscessus expressing Tdtomato (red). The surface and the endolysosomal system of the macrophages 775 were detected using anti-CD63 antibodies (green). The nuclei were stained with DAPI (blue). White arrows indicate individual or aggregate mycobacteria. Scale bar, 20 μ m. ** *P* \leq 0.01, *** *P* \leq 0.001. 776

777

Figure 3. Intracellular activity of RFB on S and R clinical isolates. CFU counts of clinical isolates
exposed to 25 and 12.5 μg/mL RFB. Macrophages were infected with *M. abscessus* (A-B) *M. bolletii*(C-D) or *M. massiliense* (E-G) clinical strains belonging to S or R morphotypes at MOI of 2:1 for 3 hrs
prior to treatment with 250 μg/mL AMK for 2 hrs to kill extracellular bacteria. Following extensive
PBS washes, cells were exposed to 50 μg/mL RIF, 50 μg/mL AMK, 25 or 12.5 μg/mL RFB. CFU were

determined at 0, 1 and 3 days post-infection. Data are mean values \pm SD for two independent experiments. Data were analysed using the *t*-test. **P* \leq 0.05, ** *P* \leq 0.01, *** *P* \leq 0.001.

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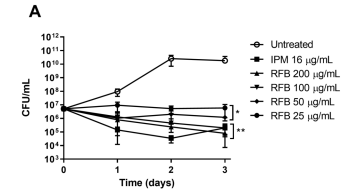
Figure 4. Activity of RFB on extracellular and intracellular cords. (A) Total number of cords 786 displayed in 20 fields at 3 days post-infection after infection of macrophages with M. abscessus R 787 788 variant. Data are mean values ± SD for three independent experiments performed in triplicate. Data 789 were analysed using one tailed Mann Whitney's t-test. (B) Percentage of cords formed either 790 extracellularly or intracellularly. The two categories were counted at 3 days post-infection in the absence of antibiotics or in the presence of 50 µg/mL RIF, 50 µg/mL AMK or 12.5 µg/mL RFB. 791 792 Extracellular or intracellular cords are highlighted using the indicated colour codes. Values are means 793 ± SD for two independent experiments performed each time in triplicate. (C) Four immuno-794 fluorescent fields were taken at 3 days post-infection showing the cords formed extracellularly or 795 within macrophages infected with *M. abscessus* R variant expressing Tdtomato (red). Macrophages were infected for 3 days in the presence of DMSO, RIF (50 µg/mL), AMK (50 µg/mL) or RFB (12.5 796 µg/mL). The macrophage surface was stained using anti-CD63 antibodies (green). The nuclei were 797 798 stained with DAPI (blue). White arrows indicate intracellular cords, while red arrows indicate extracellular cords. Scale bars represent 20 µm. Results represent the average of a total of 120 fields 799 per condition. **** $P \leq 0.0001$. 800

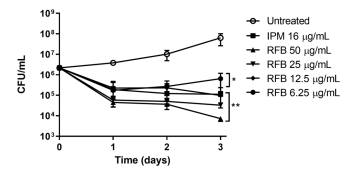
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802 Figure 5. RFB displays high bactericidal activity against *M. abscessus* in an embryonic zebrafish 803 infection model. (A) Groups of uninfected embryos were immersed in water containing increasing 804 concentrations of RFB (ranging from 3.125 to 250 µg/mL) for 4 days. The red bar indicates the 805 duration of treatment. The graph shows the survival of the RFB-treated and untreated (DMSO) embryos over a 12-days period. (B) Zebrafish embryos at 30 hrs post-fertilisation were intravenously 806 infected with approximately 250-300 CFU of *M. abscessus* CIP104536^T (R variant) expressing 807 tdTomato (n=20-25). A standard PBS injection control was included for each experiment. At 1 dpi, 808 embryos were randomly split into equal groups of approximately 20 embryos per group, and varying 809 810 concentrations of RFB (5 to 50 μ g/mL) were added to the water. DMSO was included as a positive 811 control group. RFB was changed daily after which, embryos were washed twice in fresh embryo 812 water, maintained in embryo water and monitored daily over a 12-days period. Each treatment group 813 was compared against the untreated infected group with significant differences calculated using the 814 log-rank (Mantel-Cox) statistical test for survival curves. Data shown is the merge of three 815 independent experiments (C) Bacterial burden was determined at 2, 4 and 6 days post-infection following treatment with either DMSO or 50 µg/mL RFB. Bacteria were quantified by fluorescent 816 817 pixel count determination using ImageJ software, with each data point representing a single embryo. 818 Error bars represent standard deviations. Statistical significance was determined by Student's *t*-test. 819 The plots represent a pool of 2 independent experiments containing approximately 20-25 embryos 820 per group. (D) Representative embryos from the untreated group (WT) (upper panel) and from the treated group with 50 µg/mL RFB at 6 days post-infection. White arrowheads show tdTomato-821 expressing bacteria. Scale bars represent 1 mm. $*P \le 0.05$, $***P \le 0.001$, $****P \le 0.0001$. 822

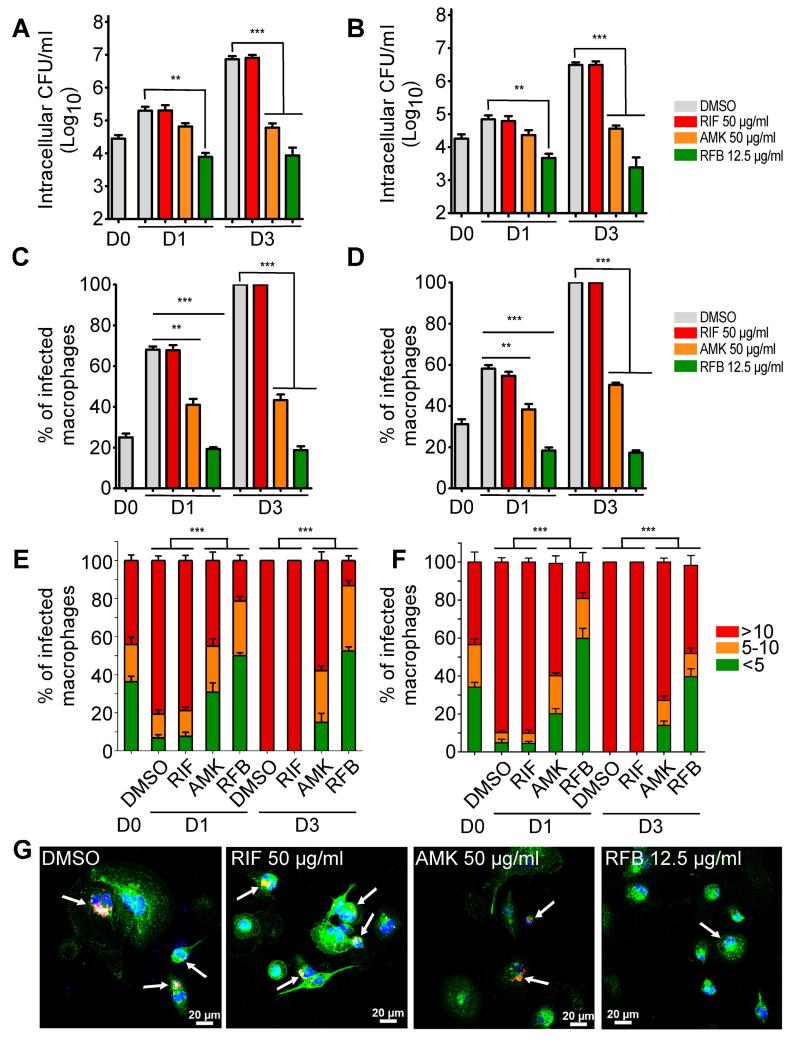
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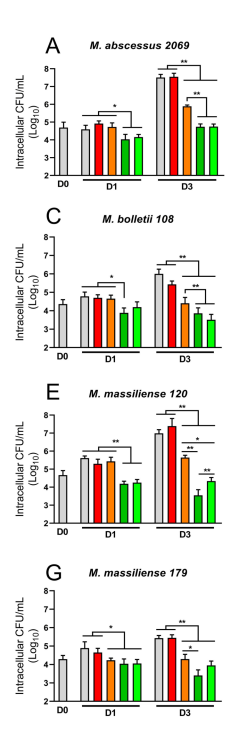
Figure 6. RFB reduces the pathophysiological traits of *M. abscessus* infection in zebrafish embryos. 824 825 (A) Proportion of embryos with cords at 2 and 4 days post-infection in infected embryos that were 826 either untreated or treated with 50 μ g/mL RFB (250-300 CFU, n=30). Data were analysed using an unpaired student's t-test. Data shown is the mean of three independent experiments \pm SD. (B) Total 827 828 percentage of embryos with abscesses at 4 and 6 dpi in infected embryos that were either untreated or treated with 50 µg/mL RFB (250-300 CFU, n=30). Data were analysed using an unpaired student's 829 830 t-test. Data shown is the mean of three independent experiments \pm SD. (C-D) Representative 831 zebrafish images of untreated (WT) embryos and those treated with treated with 50 μ g/mL RFB at 6 dpi. Scale bar represents 0.5 mm. White arrows indicate extracellular cords. The white box highlights 832 a large extracellular cord based on the size and morphology, with the scale bar representing 100 µm. 833 834 Red overlay represents *M. abscessus* expressing tdTomato. $*P \le 0.05$.

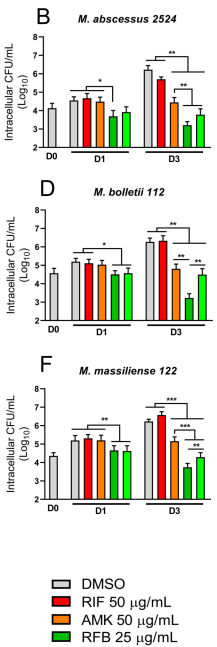




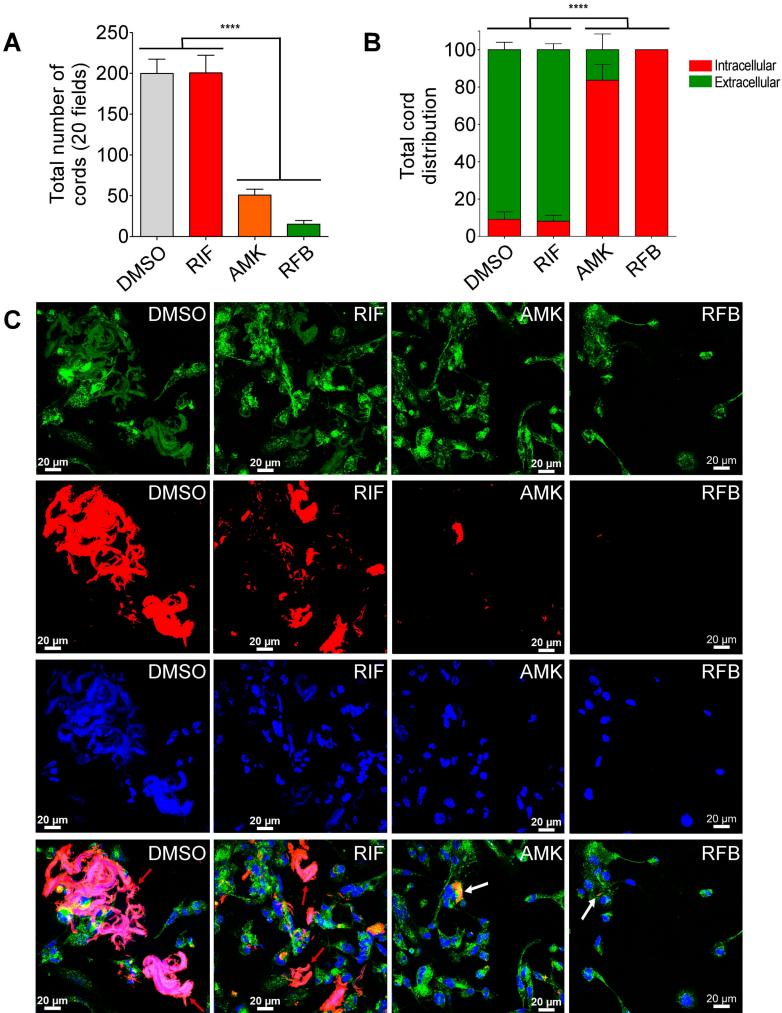








RFB 12.5 μg/mL



С

