1	Chronic infection of adult zebrafish with rough or smooth variant Mycobacterium abscessus		
2	causes necrotising inflammation and differential activation of host immunity		
3			
4	Elinor Hortle <sup>1,2</sup> *, Julia Y Kam <sup>1</sup> *, Elizabeth Krogman <sup>1</sup> *, Sherridan E Warner <sup>1,2</sup> , Pradeep		
5	Manuneedhi Cholan <sup>1</sup> , Kazu Kikuchi <sup>3,4</sup> , James A Triccas <sup>2</sup> , Warwick J Britton <sup>1,2</sup> , Matt D Johansen <sup>5</sup> ,		
6	Laurent Kremer <sup>5,6</sup> , Stefan H Oehlers <sup>1,2#</sup>		
7			
8	1 Tuberculosis Research Program at Centenary Institute, The University of Sydney, Camperdown		
9	NSW Australia		
10	2 The University of Sydney, Faculty of Medicine and Health & Marie Bashir Institute,		
11	Camperdown NSW Australia		
12	3 Developmental and Stem Cell Biology Division, Victor Chang Cardiac Research Institute,		
13	Darlinghurst, NSW Australia		
14	4 St. Vincent's Clinical School, University of New South Wales, Kensington, NSW 2052, Australia		
15	5 Centre National de la Recherche Scientifique UMR 9004, Institut de Recherche en Infectiologie		
16	de Montpellier (IRIM), Université de Montpellier, 1919 Route de Mende, 34293, Montpellier,		
17	France.		
18	6 INSERM, IRIM, 34293 Montpellier, France.		
19			
20	*These authors contributed equally and are listed in alphabetical order.		
21			
22	<sup>#</sup> Corresponding author: Dr Stefan Oehlers, <u>stefan.oehlers@sydney.edu.au</u>		
23			
24	Author contributions		
25	EH: qPCR analyses; JK: histological analysis; EK: dose finding, survival experiments, histological		
26	analysis; SW: CFU recovery assays; PMC: histological analysis; KK: provided reagents; JAT &		
27	WJB: supervision of study; MDJ: conceived study, supervision of study, wrote manuscript; LK:		
28	conceived study, provided reagents, supervision of study, wrote manuscript; SHO: conceived study,		
29	performed experiments, supervision of study, wrote manuscript.		
30			
31	Keywords: non-tuberculous mycobacterium, rapid-growing mycobacteria, animal model, zebrafish,		
32	glycopeptidolipids		
33			
34	Abstract		

35 Infections caused by *Mycobacterium abscessus* are increasing in prevalence within patient groups 36 with respiratory comorbidities including Cystic Fibrosis or Chronic Obstructive Pulmonary Disease. 37 Initial colonisation by the smooth colony *M. abscessus* (S) can be followed by an irreversible 38 genetic switch into a highly inflammatory rough colony M. abscessus (R), often associated with a 39 decline in pulmonary function. Currently available animal models such as the embryonic zebrafish, 40 have largely explored the role of innate immunity in the pathogenesis of *M. abscessus*, and 41 demonstrated that infection with the R variant produces a hyperinflammatory infection due to the 42 presence of large extracellular cords, whereas the S variant produces a chronic persistent infection. 43 However, our understanding of the role of adaptive immunity in *M. abscessus* pathogenesis is 44 largely unknown. Here, we have used intraperitoneal infection of adult zebrafish to model M. 45 *abscessus* pathogenesis in the context of fully functioning host immunity. We find infection with 46 the R variant penetrates host organs causing an inflammatory immune response leading to necrotic 47 granuloma and abscess formation within 2 weeks. The R bacilli are targeted by T cell-mediated 48 immunity and burden is progressively reduced. Strikingly, the S variant colonises host internal 49 surfaces at high loads and is met with a robust innate immune response. Invasive granuloma 50 formation is delayed in S variant infection compared to R variant infection. In mixed infections, the 51 S variant outcompetes the R variant in an adaptive-immunity dependent manner. We also find the R 52 variant activates innate immunity to detriment of S variant M. abscessus in mixed infections. These 53 findings demonstrate the applicability of the adult zebrafish to model persistent M. abscessus 54 infection.

55

### 56 Introduction

57 Mycobacterium abscessus is an increasingly recognized human pathogen responsible for a wide 58 array of clinical manifestations including muco-cutaneous infections, disseminated or chronic 59 pulmonary diseases. The latter is mostly encountered in patients with underlying lung disorders, such as bronchiectasis or cystic fibrosis (CF). Irrespective of being a rapid-growing mycobacteria 60 61 (RGM), M. abscessus displays many pathophysiological traits with slow-growing mycobacteria 62 (SGM), such as *Mycobacterium tuberculosis*. These include the capacity to persist silently within 63 granulomatous structures and to produce pulmonary caseous lesions (1, 2). In addition, M. 64 abscessus is notorious for being one of the most-drug resistant mycobacterial species, characterized 65 by a wide panel of acquired and innate drug resistance mechanisms against nearly all anti-tubercular 66 drugs, as well as many classes of antibiotics (3). Consequently, this explains the complexity and 67 duration of the treatments and the high level of therapeutic failure (4).

68 *M. abscessus* exists either as a smooth (S) or a rough (R) colony morphotype associated with 69 distinct clinical outcomes (5). Previous epidemiological studies have highlighted the association of

the R variant, persisting for many years in the infected host, with a rapid decline in the pulmonary functions (6-8). It is well established that these morphological differences between S and R variants are dependent on the presence or absence of surface-exposed glycopeptidolipids (GPL), respectively (5, 9, 10). However, our knowledge of the pathophysiological characteristics and interactions between R or S variants with the host immune cells remains largely incomplete and is hampered by the lack of animal models that are permissive to persistent *M. abscessus* infection.

Intravenous injection or aerosol administration of *M. abscessus* in immunocompetent BALB/c mice fails to establish a persistent infection, typified by a rapid clearance of the bacilli from the liver, spleen and lungs within 4 weeks (11). Immunosuppression is required to produce a progressive high level of infection with *M. abscessus* in mice, as shown in nude, SCID (severe combined immunodeficiency), interferon-gamma (GKO) and granulocyte-macrophage colonystimulating factor (GM-CSF) knock-out mice (12).

82 The contribution of B and T cells in the control of M. abscessus infection has been studied in C57BL/6 mice with Rag2<sup>-/-</sup>, Cd3e<sup>-/-</sup> and  $\mu$ MT<sup>-/-</sup> knockouts. These studies indicated that infection 83 control was primarily T cell dependent in the spleen, and both B and T cell dependent in the liver 84 (13). In addition, IFNg-receptor KO mice (ifngr1<sup>-/-</sup>) were significantly impaired in their control of 85 M. abscessus both in the spleen and in the liver, with markedly different granulomas and more 86 87 pronounced in TNF<sup>-/-</sup> mice (13). This points to the central role of T-cell immunity, IFNg and TNF for the control of M. abscessus in C57BL/6 mice, similarly to the control of M. tuberculosis 88 infection. 89

90 In recent years, alternative non-mammalian models, such as Drosophila (14), Galleria larvae 91 (15), and zebrafish embryos (16) have been developed to study the chronology and pathology of M. 92 abscessus infection and for in vivo therapeutic assessment of drugs active against M. abscessus. In 93 particular, zebrafish embryos have delivered important insights into the pathogenesis of M. 94 abscessus and the participation of innate immunity in controlling infection. The optical 95 transparency of zebrafish embryos has been used to visualise the formation of large extracellular 96 cords by the R form in vivo, representing a mechanism of immune subversion by preventing 97 phagocytic destruction and highlighting the importance bacterial virulence factors such as the 98 dehydratase MAB\_4780 and the MmpL8<sub>MAB</sub> lipid transporter (9, 17, 18). Other studies in zebrafish 99 embryos have demonstrated the contribution of host TNF signalling and IL8-mediated neutrophil 100 recruitment for protective granulomatous immunity against *M. abscessus* (19), and the link 101 between dysfunctional CFTR and vulnerability to M. abscessus infection via the macrophage 102 oxidative response (20).

103 Adult zebrafish models have been well-described for the study of mycobacterial pathogenesis 104 by *Mycobacterium marinum*, used as a surrogate for the closely related *M. tuberculosis*, and the

105 human pathogen *Mycobacterium leprae* (21-23). Encompassing a fully functional immune system, 106 previous studies in adult zebrafish with pathogenic mycobacteria such as *M. marinum* have 107 unravelled the interplay between innate and adaptive immunity in mycobacterial granuloma 108 formation and function.

Herein, we addressed whether adult zebrafish may be a useful host to analyse and compare the chronology of infection with *M. abscessus* S and R variants and to study the contribution of the T cell-mediated immunity and granulomatous response in *M. abscessus* infection.

112

113 **Results** 

114 Adult zebrafish can be chronically infected with *M. abscessus*.

115 To establish the susceptibility of adult zebrafish to infection by *M. abscessus*, we performed a dose 116 escalation experiment up to  $10^6$  CFU per animal with the rough (R) and smooth (S) variants of the reference strain CIP104536<sup>T</sup>. While animals underwent a period of sickness behaviour within the 117 118 first week of infection, we did not observe mortality for up to 4 weeks post infection (wpi) (data not 119 shown). To determine if *M. abscessus* produces a persistent infection in adult zebrafish, we performed CFU recovery on animals infected with a standard dose of 10<sup>5</sup> CFU (Figure 1A). We 120 121 observed a progressive clearance of the R variant with a 1-log reduction in burden from 1 day post 122 infection (dpi) to 4 wpi (P=0.018). Conversely, the S variant was recovered at a consistent burden 123 across the 4 week duration of the experiment (1 dpi vs 4 wpi, P=0.12). We hypothesised that the 124 better survival of the S variant compared to the R variant could be attributed to reduced 125 immunogenicity of the CIP104536 S cell surface and different modes of growth in macrophages 126 (24). To test this hypothesis, we performed qPCR detection of zebrafish immune gene expression 127 from infected adult homogenates (25). We observed higher innate immunity-associated *illb* 128 transcription in the *M. abscessus* S-infected group from early in infection and a late trend to 129 increases in *illb* transcription in 4 wpi *M. abscessus* R-infected animals (Figure 1B). Expression of cd3, indicative of total T cell numbers, was significantly decreased in all M. abscessus-infected 130 131 animals at 2 wpi but returned to uninfected levels by 4 wpi with higher cd3 expression in M. 132 abscessus R-infected compared to M. abscessus S-infected animals (Figure 1C). There was a trend 133 towards increased transcription of the Th1 activation marker *ifng* in R compared to S or uninfected 134 fish at 4 wpi (Figure 1D).

135

136 Adult zebrafish contain *M. abscessus* R within granulomas.

We next performed histology on adult zebrafish infected with fluorescent *M. abscessus* R. At 3 dpi,
bacteria were diffusely spread through the peritoneal cavity with occasional foci of infection located
external to peritoneal organs. From 10 dpi to 2 wpi we noted a heterogeneous mix of unorganised

140 lesions (Figure 2A) and organised lesions with stereotypical concentric rings of nuclei around a 141 central focus of bacteria and necrotic debris in all animals (Figure 2B). We also observed the 142 appearance of very large abscess-like granulomas filled with fluorescent bacteria and necrotic 143 debris measuring over 600 µm at a rate of no more than 1 per infected animal from 2 wpi onwards 144 (Figure 2C). Oil red O staining revealed the accumulation of foam cells in cellular rim of M. 145 abscessus R granulomas (Figure 2D), consistent with immunopathology seen in 146 immunocompromised mice infected with *M. abscessus* (12). Although we observed a fairly stable 147 proportion of organised granulomas in *M. abscessus* R-infected adult zebrafish from 10 to 28 dpi 148 (Figure 2E), the proportion of fluorescent bacteria associated with organised granulomas 149 significantly increased from 10 to 28 dpi corresponding to the appearance of abscesses at 14 dpi 150 (Figure 2F).

151

### 152 Adult zebrafish contain *M. abscessus* S infection prior to delayed granuloma formation.

Histological analysis of adult zebrafish infected with fluorescent *M. abscessus* S revealed significantly less tissue damage than the equivalent *M. abscessus* R infection up to 2 wpi. *M. abscessus* S was observed to grow freely in mesenteric spaces and form poorly organised cellular granulomas at 2 wpi (Figure 3A). We observed the appearance of tissue-invasive organised granulomas at 4 wpi (Figure 3B). Although these granulomas could reach similar size to the very large abscess-like granulomas seen in *M. abscessus* R infection, there was little Oil Red O staining in *M. abscessus* S granulomas indicating a lack of foam cell formation (Figure 3C).

We observed a significant increase in the proportion of organised granulomas in S-infected adult zebrafish from 2 to 4 wpi (Figure 3D). Similarly, the proportion of fluorescent bacteria associated with organised granulomas significantly also increased from 2 to 4 wpi corresponding to the consolidation of *M. abscessus* S into granulomas (Figure 3E).

164 The cytokine *tumour necrosis factor* (*tnf*) is essential for the containment of *M. abscessus* in 165 zebrafish embryos (19). We next took advantage of the progressive pathology observed in 4 wpi *M.* 166 *abscessus* S-infected  $TgBAC(tnfa:GFP)^{pd1028}$  animals, where GFP expression is driven by the *tnfa* 167 promoter, to investigate if *tnf* expression is linked to granuloma formation. Expression of GFP was 168 concentrated to tightly organised necrotic granulomas demonstrating a conserved induction of *tnf* 169 expression during *M. abscessus* granuloma formation in adult zebrafish (Figure 3F).

170

# 171 Adaptive immunity is necessary for control of *M. abscessus* infection in adult zebrafish.

Given the requirement for T cells to maintain granuloma structure in adult zebrafish *M. marinum* infection (23), we next asked if there was T cell involvement around *M. abscessus* granulomas using the recently described  $TgBAC(lck:GFP)^{vcc4}$  zebrafish line (23, 26). We observed T cell

175 association and penetration throughout unorganised and organised M. abscessus R granulomas, but 176 T cells were largely excluded from the cores of the very large abscess-like lesions (Figure 4A). 177 Conversely, we did not observe T cell interaction with M. abscessus S growing around peritoneal 178 organs until invasive granuloma formation after 2 wpi, which was indistinguishable from the T cell 179 response to M. abscessus R infection (Figure 4B). To directly test the requirement of T cells in containing *M. abscessus* we next utilised the  $lck^{-/-sa410}$  mutant line which is T cell-deficient. We 180 infected wild type (WT) control and  $lck^{-/-sa410}$  mutant adult zebrafish with both the S and R variants. 181 T cell-deficient adult zebrafish were significantly more susceptible to M. abscessus R infection with 182 183 reduced survival over 4 weeks of infection (P = 0.0005, Log-rank test) (Figure 4C). T cell 184 deficiency had a less pronounced effect on the survival of animals infected with M. abscessus S 185 compared to *M. abscessus* R infection (WT S vs lck-/- S P = 0.03, Log-rank test), although both 186 groups eventually succumbed to infection at the same rate after 5 wpi (P = 0.78, Log-rank test) (Figure 4C). Consequently, bacterial burden was significantly increased in 2 wpi *lck*<sup>-/-sa410</sup> mutants 187 188 infected with the R (Figure 4D), but not the S variant (Figure 4E). These observations confirm the 189 highly inflammatory nature of *M. abscessus* R infection is conserved in the adult zebrafish model 190 and mediated by T-cells. Conversely, colonisation by M. abscessus S is contained by sufficient 191 activation of the innate immune response until invasive infection between 2 and 4 wpi.

192

## 193 <u>M. abscessus S has a survival advantage over M. abscessus R.</u>

194 To examine our hypothesis that *M. abscessus* S has a survival advantage over *M. abscessus* R in the 195 adult zebrafish infection model, we performed co-infection of adult zebrafish with equal numbers of 196 each variant expressing either Wasabi or Tdtomato fluorescent proteins to enable simple tracking 197 (Figure 5A). Analysis of the ratio of R:S colonies recovered revealed a clear and rapid shift in 198 population proportions from 1:1 at 1 dpi to 0.5 rough:1 smooth ratio that remained stable at 2 wpi 199 (Figure 5B). Coinfection did not affect the progressive reduction in *M. abscessus* R burden as near 200 identical *M. abscessus* R CFUs were recovered from single and mixed-infected animals at 1 and 2 201 wpi (Figure 5C). Coinfection did cause a decrease in the number of recoverable *M. abscessus* S that 202 was not observed in single variant infections at 1 and 2 wpi demonstrating a negative effect of R 203 granuloma formation on the survival of *M. abscessus* S (Figure 5C). Whole mount and histological 204 examination of bacterial distribution in animals infected with M. abscessus R expressing Wasabi 205 and M. abscessus S expressing Tdtomato revealed co-mingling of S and R variants within 206 granulomas (Figure 5D).

We hypothesised that the T cell response induced by the R variant to antigens shared with the S strain could be responsible for the clearance of S strain in mixed infections. To test this hypothesis we infected T-cell deficient  $lck^{-/-sa410}$  mutant animals with a mixed infection of R and S

210 variants of *M. abscessus*. CFU recovery from these mixed infections in the T-cell deficient animals

211 revealed a similar decrease in *M. abscessus* S burden as observed in WT animals (Figure 5E).

212 Therefore the clearance of *M. abscessus* S strain is mediated by innate immune mechanisms.

213 Interestingly,  $lck^{-/-sa410}$  zebrafish had a higher proportion of rough to smooth than in comparable WT

animals, confirming the importance of the adaptive immune response for controlling *M. abscessus* 

215 R strain burden (Figure 5F).

216

## 217 Discussion

In this study, we report for the use of adult zebrafish to probe both host and mycobacterial determinants of pathogenesis during persistent infection with *M. abscessus*. Infection with the R and S variants was maintained over months of infection in genetically intact animals, a major improvement on existing mouse models of *M. abscessus* infection.

222 While the R variant induces a more robust and aggressive infection than the S morphotype in 223 zebrafish embryos (9), this appears to not the case in the adult fish. We observed better clearance of 224 the R variant and establishment of a higher burden of persistent infection with the S variant. One 225 possible explanation for the better survival of the S compared to the R is that, in the presence of 226 effective adaptive immunity, R infection is better controlled due to the induction of a potent Th1 227 cell-mediated response, as evidenced by the increased expression of the CD3 marker and IFN-228 gamma response at later time points. This contribution of the T cell response was further substantiated using T cell-deficient fish, where infection of  $lck^{-/-}$  fish with the R bacilli resulted in a 229 230 higher bacterial burden than in WT fish, which was not observed with S bacilli. These observations 231 provide insight into the clinical observation that AIDS patients are not at increased risk of M. 232 abscessus infection to the same degree that AIDS is a risk factor for *M. tuberculosis* and other non-233 tuberculous mycobacterium infections such as Mycobacterium avium.

234 It is well known that the intracellular lifestyle of S and R M. abscessus variants differ 235 significantly, resulting in entirely distinct infection scenarios (27). The absence of GPL on the outer 236 mycomembrane causes corded growth of R variants, resulting in multiple bacilli being 237 simultaneously phagocytosed by macrophages and overloaded phagosomes that rapidly activate 238 autophagy pathways (27). Comparatively, the S variant is able to survive for an extended period of 239 time within the phagosome, producing a chronic and persistent infection (28, 29). As such, these 240 polar infection responses may explain why the R displays increased granuloma formation at 2 wpi, 241 compared to S which shows a significantly delayed onset of granuloma formation. Moreover, this 242 observation matches the superior *in vivo* growth performance of S bacilli compared to R (Fig 1A), 243 suggesting that the R variant is at an overall disadvantage because of its intrinsic hyper-244 inflammatory status and the activation of effective adaptive immunity that results in granuloma formation. Taken together, our data provides additional evidence for the distinct intracellular fates of both S and R variants *in vivo*, and further implicates the role of adaptive immunity in granuloma formation and control of *M. abscessus* infection in an adult zebrafish model.

Our qPCR analysis surprisingly demonstrated that the S variant elicits greater production of *il1b*; a key inflammatory cytokine, during the first 2 wpi compared to the R variant (Fig 1B). It is likely that this observation is the nett effect of progressive increase in bacterial burden of S variants over time, coupled with the presence of extracellular bacilli and unorganised granuloma formation observed at 2 wpi, compared to the containment and clearance of the R variant in organised granulomas.

254 T cells are critical host determinants in the control of mycobacterial infection (30). 255 Recruitment of T cells into granulomas are thought to be essential in containing persistent infection, 256 while T cell deficiencies are associated with greater mycobacterial infection severities (21, 30-32). 257 Recently, an adult zebrafish infection model for *M. leprae* demonstrated that T cells are essential 258 for containment of infection (29). We examined the recruitment of T cells within granulomas and 259 identified that S variant granulomas were marked by the absence of T cell infiltration at 2 wpi, 260 highlighting the fact that T cells may play a less significant role in S variant infections than those with R variants. Using the  $lck^{-}$  mutants, we have shown that adult zebrafish are highly susceptible 261 262 to *M. abscessus* infection and succumb to intraperitoneal infection within 40 days in the absence of 263 T cells, irrespective of bacterial morphotype. Importantly, the R variant displayed an improved in 264 vivo growth performance in the absence of T cells when compared to wild-type zebrafish, 265 highlighting the role of T cells in the control of R variants. However, this observation was not 266 maintained with the S variant, which showed no increase in bacterial growth in vivo irrespective of 267 the absence of T cells early in infection.

Our co-infection experiments further support the theory that tissue destruction caused by R variant activates protective trans-acting host innate immunity that impairs bacterial growth, thereby restricting S growth (Graphical Abstract). This suggests *M. abscessus* must balance the benefits of R variant pathogenicity allowing individuals to kill and escape macrophage containment, with the need to avoid activation of host-protective immunity at a population level.

To date, our understanding of the diverse immune responses between S and R variants have only been thoroughly described with respect to innate immunity, and currently our knowledge pertaining to adaptive immunity in *M. abscessus* infection has been poorly characterised. Using this new adult zebrafish *M. abscessus* infection model, we have shown that S and R variants produce strikingly different disease phenotypes, which was further exemplified in the absence of a functioning adaptive immune response. Consequently, these results suggest that the host-pathogen interactions dictating *M. abscessus* pathogenesis are complex and may implicate adaptive immunity

to a greater extent than originally anticipated. Future work should exploit this new animal model in

281 combination with zebrafish lacking the Cystic fibrosis transmembrane conductance regulator gene,

and for the development and testing of novel antibiotics and vaccine candidates that may be used

- 283 for the treatment of *M. abscessus* infection.
- 284

### 285 Methods

## 286 Zebrafish strains and handling

Zebrafish strains used in this study are AB strain wildtype,  $TgBAC(tnfa:GFP)^{pd1028}$ ,  $TgBAC(lck:EGFP)^{vcc4}$ , lck-/- <sup>sa410</sup> (26, 33). Animals were held in a 28°C incubator with a 14:10 hour light:dark cycle. Animals were infected by intraperitoneal injection with approximately 10<sup>5</sup> CFU *M*. *abscessus*, unless otherwise stated, using a 31 G insulin needle and syringe as previously described (34). Infected zebrafish were recovered into system water and held in 1 L beakers with daily feeding for the duration of the experiment. Infection experiments were carried out with ethical approval from the Sydney Local Health District Animal Welfare Committee approval 16-037.

294

# 295 <u>M. abscessus strains and handling</u>

Rough (R) and smooth (S) variants of *M. abscessus* strain CIP104536<sup>T</sup> were grown at 30°C in Middlebrook 7H9 broth supplemented with 10% Oleic acid/Albumin/Dextrose/Catalase (OADC) enrichment and 0.05% Tween 80 or on Middlebrook 7H10 agar containing 10% OADC (7H10 OADC). Recombinant *M. abscessus* strains expressing Tdtomato or Wasabi were grown in the presence of 500  $\mu$ g/ml hygromycin (9, 19). Homogenous bacterial suspensions for intraperitoneal injection in adult fish were prepared as previously reported (35).

302

## 303 <u>Bacterial recovery</u>

Animals were euthanised by tricaine anaesthetic overdose and rinsed in sterile water. Individual carcasses were homogenised and serially diluted into sterile water. Homogenates were plated onto 7H10 OADC supplemented with 300  $\mu$ g/ml hygromycin. Plates were grown for at least 4 days at 37 degrees.

308

# 309 Zebrafish and *M. abscessus* gene expression analysis by qPCR

RNA was extracted from whole fish homogenates and was reverse transcribed with the Applied Biosystems High Capacity cDNA kit and qPCR was carried out on an Agilent Technologies Stratagene Mx3005P. Zebrafish gene expression primers were previously described by Hammaren *et al.* (25).

314

# 315 <u>Histology</u>

316 Animals subjected to cryosectioning as previously described (34). Briefly, euthanasia was 317 performed by tricaine anaesthetic overdose and specimens were fixed for 2-4 days in 10% neutral 318 buffered formalin at 4°. Specimens were then rinsed in PBS, incubated overnight in 30% sucrose, 319 incubated overnight in 50/50 30% sucrose and OCT, and finally incubated overnight in OCT prior 320 to freezing at  $-80^{\circ}$ . Cryosectioning was performed to produce 20  $\mu$ m thick sections. Sections were 321 post-fixed for 1-2 minutes in 10% neutral buffered formalin and rinsed in PBS prior to further 322 processing. Slides for fluorescent imaging were mounted with coverslips using Fluoromount G 323 containing DAPI. Oil Red O staining was performed as previously described (34, 36). T cells were detected in sections from  $TgBAC(lck:GFP)^{vcc4}$  zebrafish by anti-GFP staining (primary antibody: 324 325 ab13970, Abcam; secondary antibody: ab150173, Abcam), stained slides were then mounted with 326 coverslips using Fluoromount G containing DAPI. All imaging was carried out on a Leica 327 DM6000B microscope.

328

329 Statistics

All statistical testing was carried out using Graphpad Prism. Each data point indicates a singleanimal unless otherwise stated.

332

## 333 Acknowledgements

We thank the Centenary imaging facility core and Sydney Cytometry staff Drs Kristina Jahn,Angela Kurz, and David Liu, for their assistance.

Funding: Australian National Health and Medical Research Council CJ Martin Early Career
Fellowship APP1053407 and Project Grant APP1099912; The University of Sydney Fellowship
G197581; NSW Ministry of Health under the NSW Health Early-Mid Career Fellowships Scheme
H18/31086; the Kenyon Family Foundation Inflammation Award; Australian-French Association
for Research and Innovation (AFRAN) Initiative to SHO. The Fondation pour la Recherche
Médicale DEQ20150331719 to LK. Post-doctoral fellowship granted by Labex EpiGenMed, an
"Investissements d'avenir" program ANR-10-LABX-12-01 to MDJ.

343

## 344 **References**

- H. Medjahed, J. L. Gaillard, J. M. Reyrat, Mycobacterium abscessus: a new player in the
   mycobacterial field. *Trends Microbiol* 18, 117-123 (2010).
- J. F. Tomashefski, Jr., R. C. Stern, C. A. Demko, C. F. Doershuk, Nontuberculous
  mycobacteria in cystic fibrosis. An autopsy study. *Am J Respir Crit Care Med* 154, 523-528
  (1996).

350	3.	R. Nessar, E. Cambau, J. M. Reyrat, A. Murray, B. Gicquel, Mycobacterium abscessus: a new
351		antibiotic nightmare. The Journal of antimicrobial chemotherapy 67, 810-818 (2012).
352	4.	B. E. Ferro et al., Failure of the Amikacin, Cefoxitin, and Clarithromycin Combination
353		Regimen for Treating Pulmonary Mycobacterium abscessus Infection. Antimicrob Agents
354		<i>Chemother</i> <b>60</b> , 6374-6376 (2016).
355	5.	S. T. Howard et al., Spontaneous reversion of Mycobacterium abscessus from a smooth to a
356		rough morphotype is associated with reduced expression of glycopeptidolipid and
357		reacquisition of an invasive phenotype. <i>Microbiology</i> 152, 1581-1590 (2006).
358	6.	E. Catherinot et al., Acute respiratory failure involving an R variant of Mycobacterium
359		abscessus. J Clin Microbiol 47, 271-274 (2009).
360	7.	C. R. Esther, Jr., D. A. Esserman, P. Gilligan, A. Kerr, P. G. Noone, Chronic Mycobacterium
361		abscessus infection and lung function decline in cystic fibrosis. J Cyst Fibros 9, 117-123
362		(2010).
363	8.	B. E. Jonsson et al., Molecular epidemiology of Mycobacterium abscessus, with focus on
364		cystic fibrosis. J Clin Microbiol 45, 1497-1504 (2007).
365	9.	A. Bernut et al., Mycobacterium abscessus cording prevents phagocytosis and promotes
366		abscess formation. Proc Natl Acad Sci USA 111, E943-952 (2014).
367	10.	H. Medjahed, J. M. Reyrat, Construction of Mycobacterium abscessus defined
368		glycopeptidolipid mutants: comparison of genetic tools. Appl Environ Microbiol 75, 1331-
369		1338 (2009).
370	11.	A. Bernut et al., In vivo assessment of drug efficacy against Mycobacterium abscessus using
371		the embryonic zebrafish test system. Antimicrob Agents Chemother 58, 4054-4063 (2014).
372	12.	A. Obregon-Henao et al., Susceptibility of Mycobacterium abscessus to antimycobacterial
373		drugs in preclinical models. Antimicrob Agents Chemother 59, 6904-6912 (2015).
374	13.	M. Rottman et al., Importance of T cells, gamma interferon, and tumor necrosis factor in
375		immune control of the rapid grower Mycobacterium abscessus in C57BL/6 mice. Infect
376		Immun <b>75</b> , 5898-5907 (2007).
377	14.	C. T. Oh, C. Moon, M. S. Jeong, S. H. Kwon, J. Jang, Drosophila melanogaster model for
378		Mycobacterium abscessus infection. Microbes Infect 15, 788-795 (2013).
379	15.	M. Meir, T. Grosfeld, D. Barkan, Establishment and Validation of Galleria mellonella as a
380		Novel Model Organism To Study Mycobacterium abscessus Infection, Pathogenesis, and
381		Treatment. Antimicrob Agents Chemother 62 (2018).
382	16.	A. Bernut, J. L. Herrmann, D. Ordway, L. Kremer, The Diverse Cellular and Animal Models
383		to Decipher the Physiopathological Traits of Mycobacterium abscessus Infection. Front Cell
384		Infect Microbiol <b>7</b> , 100 (2017).

- 17. V. Dubois *et al.*, MmpL8MAB controls Mycobacterium abscessus virulence and production
  of a previously unknown glycolipid family. *Proc Natl Acad Sci U S A* 115, E10147-E10156
- 387 (2018).
- 18. I. Halloum *et al.*, Deletion of a dehydratase important for intracellular growth and cording
  renders rough Mycobacterium abscessus avirulent. *Proc Natl Acad Sci U S A* 113, E42284237 (2016).
- A. Bernut *et al.*, Mycobacterium abscessus-Induced Granuloma Formation Is Strictly
   Dependent on TNF Signaling and Neutrophil Trafficking. *PLoS Pathog* 12, e1005986 (2016).
- A. Bernut *et al.*, CFTR Protects against Mycobacterium abscessus Infection by Fine-Tuning
  Host Oxidative Defenses. *Cell reports* 26, 1828-1840 e1824 (2019).
- 21. C. A. Madigan, J. Cameron, L. Ramakrishnan, A Zebrafish Model of Mycobacterium leprae
  Granulomatous Infection. *J Infect Dis* 216, 776-779 (2017).
- 397 22. S. H. Oehlers *et al.*, Interception of host angiogenic signalling limits mycobacterial growth.
   398 *Nature* 517, 612-615 (2015).
- L. E. Swaim *et al.*, Mycobacterium marinum infection of adult zebrafish causes caseating
  granulomatous tuberculosis and is moderated by adaptive immunity. *Infect Immun* 74, 61086117 (2006).
- 402 24. A. L. Roux *et al.*, The distinct fate of smooth and rough Mycobacterium abscessus variants
  403 inside macrophages. *Open Biol* 6 (2016).
- 404 25. M. M. Hammaren *et al.*, Adequate th2-type response associates with restricted bacterial 405 growth in latent mycobacterial infection of zebrafish. *PLoS Pathog* **10**, e1004190 (2014).
- 406 26. K. Sugimoto, S. P. Hui, D. Z. Sheng, M. Nakayama, K. Kikuchi, Zebrafish FOXP3 is required
  407 for the maintenance of immune tolerance. *Dev Comp Immunol* **73**, 156-162 (2017).
- 408 27. A.-L. Roux *et al.*, The distinct fate of smooth and rough Mycobacterium abscessus variants
  409 inside macrophages. *Open biology* 6 (2016).
- 410 28. A. Bernut *et al.*, *Mycobacterium abscessus* cording prevents phagocytosis and promotes
  411 abscess formation. *Proceedings of the National Academy of Sciences* **111**, 943-952 (2014).
- 412 29. L. Ramakrishnan, Revisiting the role of the granuloma in tuberculosis. *Nature Reviews*413 *Immunology* 12, 352-366 (2012).
- 414 30. F. M. Collins, Mycobacterial disease, immunosuppression, and acquired immunodeficiency
  415 syndrome. *Clin Microbiol Rev* 2, 360-377 (1989).
- T. Mogues, M. E. Goodrich, L. Ryan, R. LaCourse, R. J. North, The relative importance of T
  cell subsets in immunity and immunopathology of airborne Mycobacterium tuberculosis
  infection in mice. *J Exp Med* 193, 271-280 (2001).

419 32. J. D. Yang et al., Mycobacterium tuberculosis-specific CD4+ and CD8+ T cells differ in their 420 capacity to recognize infected macrophages. PLoS Pathog 14, e1007060 (2018). 421 33. L. Marjoram et al., Epigenetic control of intestinal barrier function and inflammation in 422 zebrafish. Proc Natl Acad Sci U S A 112, 2770-2775 (2015). 423 34. T. Cheng, J. Y. Kam, M. D. Johansen, S. H. Oehlers, High content analysis of granuloma 424 histology and neutrophilic inflammation in adult zebrafish infected with Mycobacterium 425 marinum. Micron 10.1016/j.micron.2019.102782 (2019). 426 35. A. Bernut et al., Deciphering and Imaging Pathogenesis and Cording of Mycobacterium 427 abscessus in Zebrafish Embryos. J Vis Exp 10.3791/53130 (2015). 428 36. M. D. Johansen *et al.*, Mycobacterium marinum infection drives foam cell differentiation in 429 zebrafish infection models. Dev Comp Immunol 88, 169-172 (2018). 430 431 **Figure Legends** 432 433 Figure 1: *M. abscessus* establishes chronic infection in adult zebrafish. 434 A. Enumeration of CFUs from adult zebrafish infected with either the R or the S morphotype of M. 435 abscessus. 436 B. Gene expression analysis of zebrafish interleukin 1b. 437 C. Gene expression analysis of zebrafish *cd3*. 438 D. Gene expression analysis of zebrafish *interferon gamma*. 439 Statistical tests by one-way ANOVA with Tukey's multiple comparisons test. Data is pooled from 440 10 animals across 2 independent experiments. 441 442 Figure 2: *M. abscessus* R infection causes progressive granulomatous pathology. 443 A. Stereotypical unorganised granuloma in a 2 wpi adult zebrafish infected with the R variant of M. 444 abscessus expressing Tdtomato. Scale bar indicates 100 µm. 445 B. i. Stereotypical organised granuloma in a 2 wpi adult zebrafish infected with R M. abscessus-446 Tdtomato and ii. Higher magnification image of granuloma wall. Arrowheads indicate epithelised 447 macrophage nuclei surrounding the mycobacterial core. Scale bar indicates 100 µm. 448 C. Example of a very large abscess-like granuloma in a 2 wpi adult zebrafish infected with R M. 449 abscessus-Tdtomato measuring approximately 600 µm in diameter. Scale bar indicates 200 µm. 450 D. Example of Oil Red O-stained very large abscess-like granuloma in a 2 wpi adult zebrafish 451 infected with R M. abscessus. Arrowheads indicate Oil Red O-positive foamy macrophages 452 surrounding the mycobacterial core. Scale bar indicates 200 µm. 13

- 453 E. Quantification of granuloma organisation in adult zebrafish infected with R M. abscessus. Data is
- 454 pooled from 2 animals per timepoint.
- 455 F. Quantification of bacterial burden stratified by granuloma organisation in adult zebrafish infected
- 456 with R *M. abscessus*. Data is pooled from 2 animals per timepoint, statistical testing by Chi-squared 457 test.
- 458
- 459 Figure 3: *M. abscessus* S infection causes delayed progressive granulomatous pathology.
- 460 A. Examples S M. abscessus-Tdtomato lesions from 2 wpi adult zebrafish. i. M. abscessus growing
- 461 free, external to peritoneal organs. ii and iii. Stereotypical examples of unorganised cellular 462 granulomas around sites of sparse *M. abscessus* infection.
- 463 B. Stereotypical large granuloma found in 4 wpi adult zebrafish infected with S M. abscessus-
- 464 Tdtomato. Note lack of punctate bacterial fluorescence in the core of the granuloma compared to 465 second granuloma to the right.
- 466 C. Example of Oil Red O-stained large granuloma in 4 wpi adult zebrafish infected with S M.
- 467 *abscessus*. Note lack of lipid staining in the rim of the granuloma compared to R infection.
- 468 D. Quantification of granuloma organisation in adult zebrafish infected with S *M. abscessus*. Data is
- 469 pooled from at least 2 animals per timepoint, statistical testing by Chi-squared test.
- 470 E. Quantification of bacterial burden stratified by granuloma organisation in adult zebrafish infected
- with S *M. abscessus*. Data is pooled from at least 2 animals per timepoint, statistical testing by Chi-squared test.
- 473 F. Example of S *M. abscessus*-Tdtomato lesions in 4 wpi  $TgBAC(tnfa:GFP)^{pd1028}$  adult zebrafish.
- 474 Arrowheads indicate organised necrotic granulomas with strong *tnfa* expression marked by GFP.
- 475 Scale bars indicate 200 μm.
- 476
- 477 Figure 4: T cells are necessary to control R but not S *M. abscessus* infection.
- 478 A. Examples of T cell recruitment to granulomas in 2 wpi  $TgBAC(lck:EGFP)^{vcc4}$  adult zebrafish
- 479 infected with R M. abscessus-Tdtomato . i. Example of an unorganised granuloma. ii. Example of
- 480 an organised granuloma. iii. Example of a very large abscess-like granuloma. Scale bars indicate
- 481 100 μm.
- 482 B. Example of lack of T cell recruitment to S *M. abscessus*-Tdtomato in 2 wpi 483  $TgBAC(lck:EGFP)^{vcc4}$  adult zebrafish. Scale bar indicates 100 µm.
- 484 C. Survival analysis of WT and lck-/-<sup>sa410</sup> adult zebrafish infected with R or S *M. abscessus*.
- 485 D. Enumeration of CFUs from 2 wpi WT and lck-/- sa410 adult zebrafish infected with R M.
- 486 *abscessus*. Statistical testing by T-test.

487 E. Enumeration of CFUs from 2 wpi WT and lck-/- sa410 adult zebrafish infected with S M.

488 *abscessus*. Statistical testing by T-test.

489

- 490 Figure 5: S *M. abscessus* has a survival advantage over R *M. abscessus*.
- 491 A. Schema outlining mixed infection experiment.
- 492 B. Ratio of R:S CFUs recovered from WT adult zebrafish infected with a mixture of differentially
- 493 labelled R and S variant *M. abscessus*.
- 494 C. Enumeration of CFUs from the three groups of WT adult zebrafish outlined in panel A. Data is
- 495 pooled from two replicates, statistical testing by ANOVA.
- 496 D. Representative images of granuloma from adult zebrafish infected with a mixture of R
- 497 expressing Wasabi and S expressing Tdtomato *M. abscessus*.
- 498 E. Enumeration of CFUs from the three groups of lck-/- <sup>sa410</sup> adult zebrafish outlined in panel A.
- 499 Data is pooled from two replicates, statistical testing by ANOVA.
- 500 F. Ratio of R:S CFUs recovered from lck-/- sa410 adult zebrafish infected with a mixture of
- 501 differentially labelled R and S variants.

502

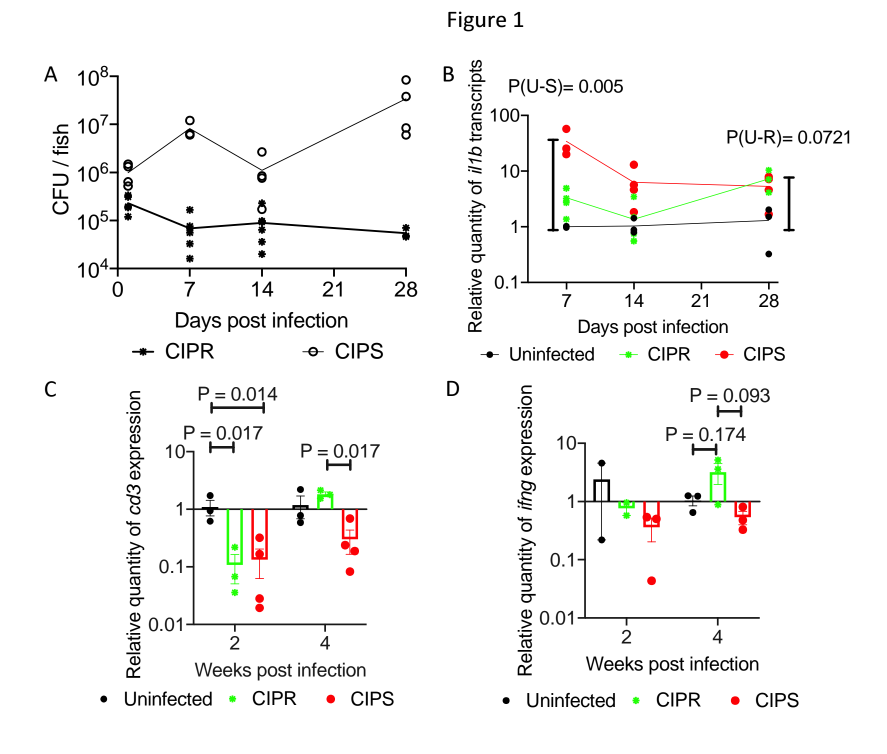
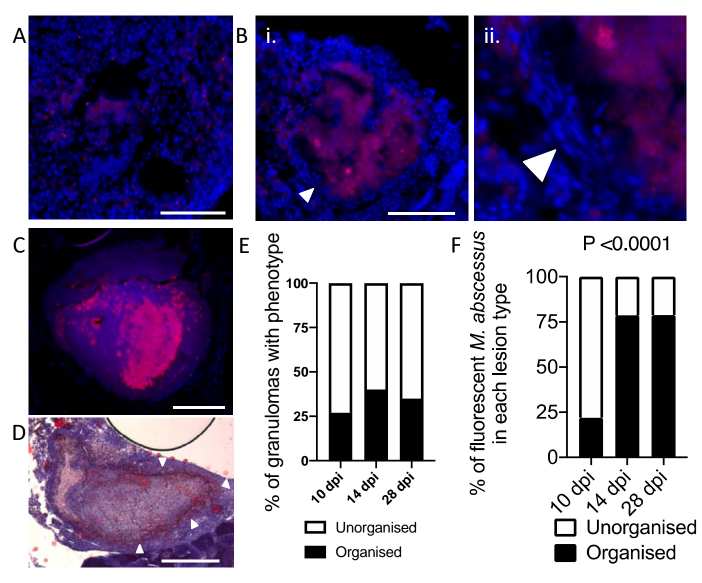
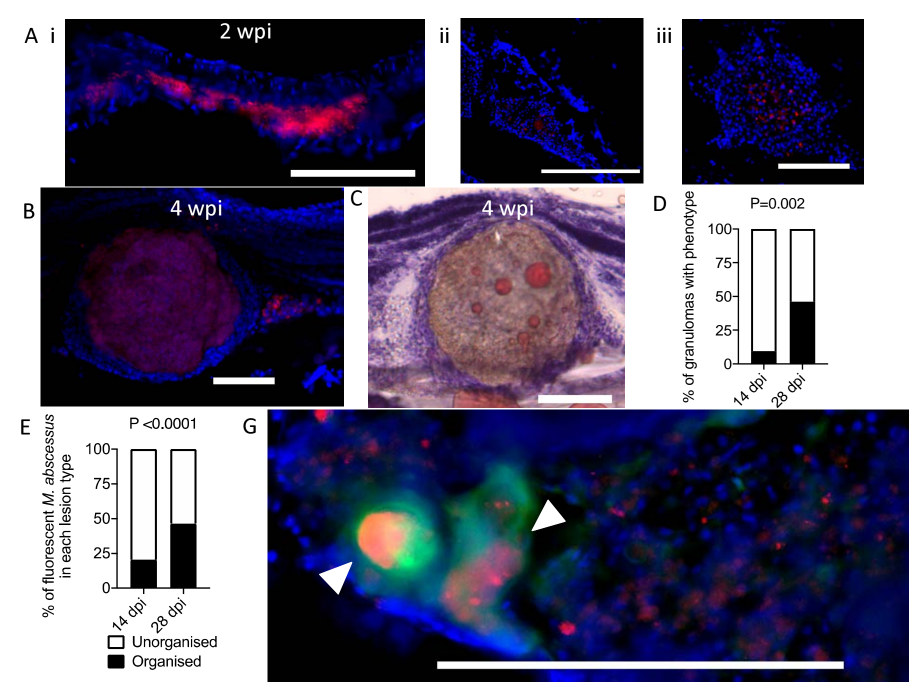


Figure 2







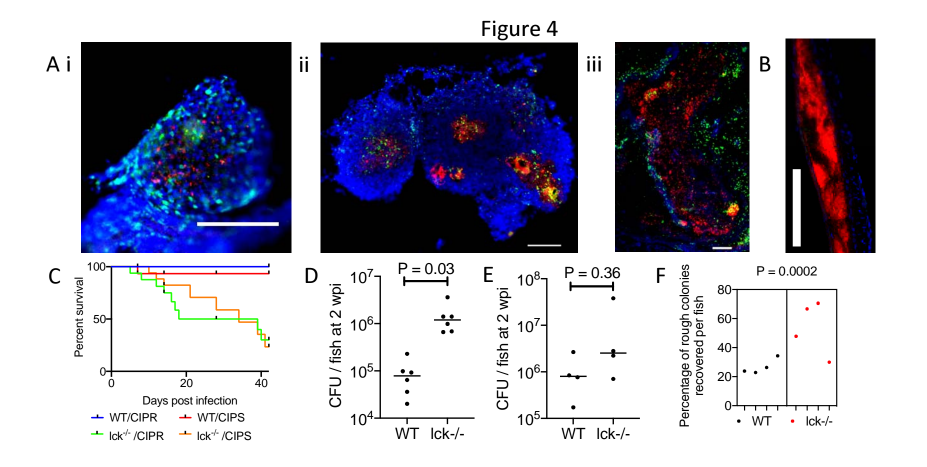
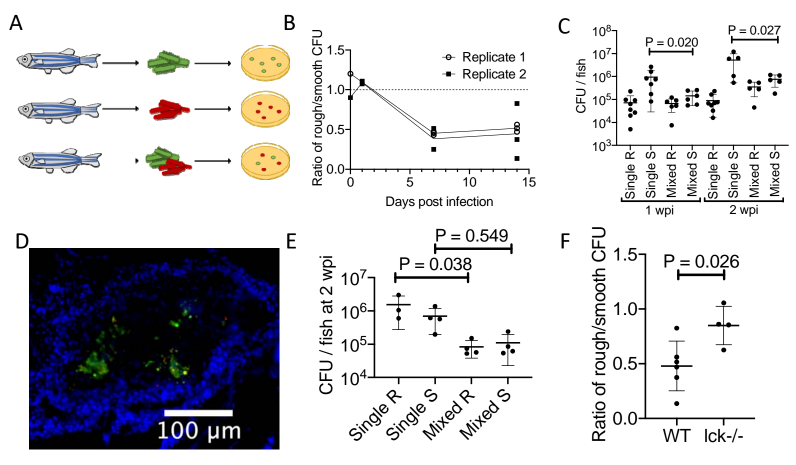


Figure 5



# **Graphical Abstract**

