

1 ***Mycobacterium abscessus*: environmental bacterium turned clinical nightmare**

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14 **Key Words:**

15 *Mycobacterium abscessus*; Non-tuberculous mycobacteria; Antimicrobial drug discovery; Cystic

16 fibrosis

17

18 **Abstract:**

19 Mycobacteria are a large family of over 100 species, most of which do not cause diseases in humans.

20 The majority of the mycobacterial species are referred to as nontuberculous mycobacteria (NTM),

21 meaning they are not the causative agent of tuberculous (TB) or leprosy, i.e. *Mycobacterium*

22 *tuberculosis* complex and *Mycobacterium leprae*, respectively. The latter group is undoubtedly the

23 most infamous, with TB infecting an estimated 10 million people and causing over 1.2 million deaths
24 in 2017 alone (1). TB and leprosy also differ from NTM in that they are only transmitted from person
25 to person and have no environmental reservoir, whereas NTM infections are commonly acquired from
26 the environment (2) (3). It took until the 1950's for NTM to be recognised as a potential lung pathogen
27 in people with underlying pulmonary disease and another 3 decades for NTM to be widely recognised
28 by the medical community when NTM, particularly *Mycobacterium avium* complex (MAC) was
29 recognised as the most common group of opportunistic pathogens in AIDS patients (4). This review
30 focusses on an emerging NTM called *Mycobacterium abscessus* (*M. abs*). *M. abs* is a rapidly growing
31 (RGM) NTM that is responsible for opportunistic pulmonary infections in patients with structural lung
32 disorders such as cystic fibrosis (CF) and bronchiectasis (5), as well as a wide range of skin and soft
33 tissue infections (SSTIs) in humans (6) (7). *M. abs* is a weakly staining Gram-positive mycobacterium
34 that is neverand is, like other NTM, most often seen in soil and aquatic environments (8). The bacillus-
35 shaped bacterium is 1-6µm long and 0.2-0.5µm in diameter, with curved ends and the presence of
36 cord factor, or trehalose 6-6'-dimycolate, a glycolipid found in the cell wall of virulent species of
37 mycobacteria that results in "serpentine cord" cell morphology is sometimes observed (8) (9). On solid
38 growth medium, *M. abs* can display either a rough (*M. abs*-R) or smooth (*M. abs*-S) morphotype, with
39 the rough morphotype displaying a more virulent phenotype than its smooth variant (10). The rough
40 morphotype is characterised by irregular parallel filaments that form ridges across the colony,
41 whereas a smooth morphology is displays a wet, smooth colony with no filaments or ridges (79). This
42 morphology is driven by cell wall glycopeptidolipid (GPL); a loss of GPL results in the reversion from
43 rough to smooth morphotype (80) (81). Moreover, it has been shown using human tissue culture
44 models of infection that *M. abs*-R is able to persist and multiply within the host macrophage whereas
45 *M. abs*-S lacks this capacity, hence its role in virulence (82). Like all other mycobacteria, *M. abs* are
46 aerobic, non-motile and acid-fast organisms with a characteristically thick, lipid-rich cell wall that is
47 hydrophobic. Due to their unusually impermeable, thick cell wall, mycobacteria are notoriously
48 resistant to many antibiotics, disinfectants and heavy metals (11). When the genome of *M. abs*

49 became available in 2009, elucidation of the resistance mechanisms of *M. abs* became an area of focus
50 for scientific research, as the considerable threat it poses to public health became more apparent (12)
51 (13) (14). In this review we will discuss how we came to understand the pathogen, how it is currently
52 treated, as well as a discussion of drug resistance mechanisms and novel treatments currently in
53 development.

54

55 **Introduction:**

56 *M. abs* was first isolated in 1952 by Moore and Frerichs from a 63-year-old woman's knee abscess (6)
57 and since then, our understanding of the pathogen has rapidly and somewhat turbulently expanded.
58 When it was first isolated, it was suggested by the authors that *M. abs* was an entirely new species of
59 NTM and was given its name due to its ability to produce subcutaneous abscesses. Interestingly, at
60 this point, *M. abs* was considered to be a pathogen of low virulence due to the perception that it was
61 primarily a pathogen causing cutaneous infections that appeared transient and self-limiting (6). 40
62 years after its discovery, *M. abs* was first implicated in pulmonary infections after an analysis of 154
63 patients with RGM pulmonary infections revealed that 82% of the isolates were *M. abs*; the disease
64 was considered to be slowly progressive but virulent nonetheless (15). Since its first identification, *M.*
65 *abs* nomenclature and species/subspecies identification have undergone many changes.

66 In 1952 (6), *M. abs* was believed to be identical to *Mycobacterium chelonae*, another RGM that infects
67 fish and amphibians, as it presented identical biochemical features (16). Then, in 1972, following an
68 international collaborative study by the International Working Group on Mycobacterial Taxonomy,
69 *M.abs* was designated subspecies status (16). 20 years later, in 1992, Kusunoki and Ezaki used DNA
70 hybridisation to establish that there is only 35% DNA relatedness between *M. chelonae* subsp.
71 *chelonae* and *M. chelonae* subsp. *abscessus*. In light of this, *M. abs* was finally re-elevated to species
72 status.

73 However, in 2004, an unusual *Mycobacterium* was isolated from a patient with hemoptoic pneumonia,
74 and researchers were unable to accurately identify the species using the techniques described above.
75 They developed partial PCR sequencing of the *rpoB* gene and were able to demonstrate that the
76 isolate shared 96.0% partial *rpoB* sequence similarity and a 98.0% *recA* gene sequence similarity with
77 only the *M. abs* type strain. They had previously proposed that *rpoB* gene sequence difference of >3%
78 and a *recA* gene sequence difference of >2% was sufficient to differentiate between different NTM
79 species. Using this new *rpoB* gene sequencing technique aided with the more traditional biochemical
80 assays and 16S rRNA gene sequencing, the authors were able to produce an accurate phylogenetic
81 tree of various NTM. They concluded that this novel isolate was a new species closely related to and
82 likely recently derived from *M. abs*. This was subsequently named *Mycobacterium massiliense* (17).

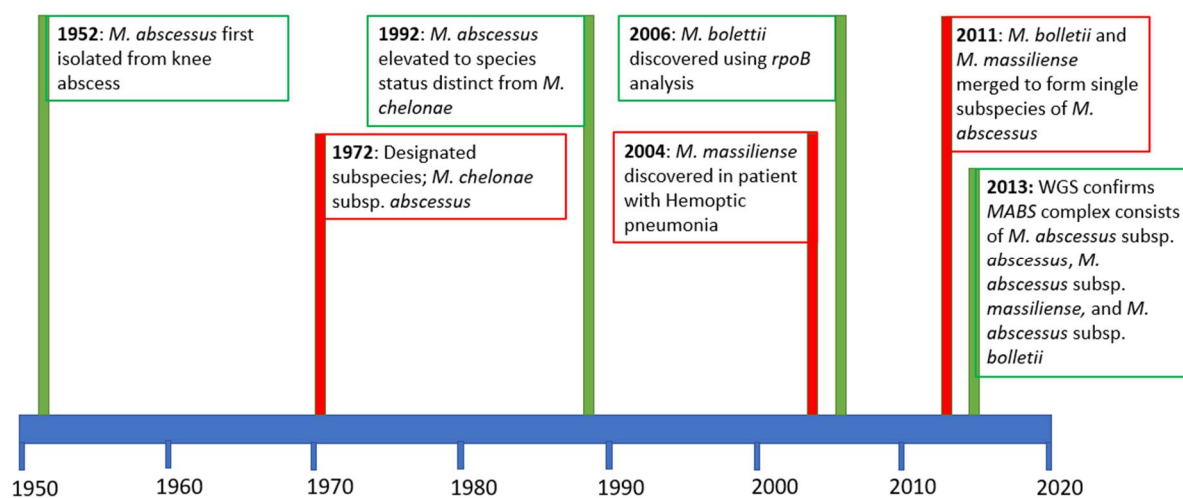
83 In 2006, *rpoB* gene sequencing was used on 59 clinical isolates of RGM (18), and they found that 15.3%
84 of these isolates were novel, corresponding to 3 new species of mycobacteria. One of these species,
85 named *Mycobacterium bolletii* by the authors, was found to share 100% 16S gene similarity and 95.6%
86 *rpoB* gene sequence similarity with *M. abs*.

87 In 2011 it was proposed by Leao *et. al.*, (19) that the *M. abscessus* complex (*MABS* complex) should
88 be amended to include *M. abscessus* subsp. *abscessus* (as before) and to combine the two subspecies
89 to form one single subspecies, *M. abscessus* subsp. *bolletii*. Finally, in 2013, whole-genome sequencing
90 (WGS) was used by Bryant *et. al.*, to identify transmission between patients with CF (20). The authors
91 subjected 168 clinical isolates of *M. abs* to WGS and a phylogenetic tree produced from the isolates
92 showed clearly, for the first time, that *M. abscessus* subsp. *abscessus*, *M. abscessus* subsp. *bolletii*, and
93 *M. abscessus* subsp. *massiliense* are three distinct subspecies belonging to the *MABS* complex. The
94 idea that *MABS* is a complex that contains 3 subspecies that are genetically very similar, but
95 phenotypically divergent was given more traction in 2016 when Tortoli *et. al.*, (21) published an
96 amended description of the *MABS* complex that highlighted the importance of subspecies
97 differentiation. The authors argued that the criteria for subspecies as proposed by Wayne *et. al.*, (22)

98 i.e. “genetically close organisms that diverge in phenotype” is appropriate in this case, considering the
 99 genetic similarity and the presence of an inducible and functional *erm*(41) gene conferring macrolide
 100 resistance in only *M. abscessus* subsp. *bolletii* and *M. abscessus* subsp. *abscessus* isolates whereas *M.*
 101 *abscessus* subsp. *massiliense* has a non-functional *erm*(41) gene.

102

103



104

105 **Figure 1:** Timeline of *Mycobacterium abscessus* taxonomy from 1950 through to the present day. The
 106 first 50 years since its discovery, no congruent terminology was in widespread use to accurately
 107 describe and differentiate *M. abs* from other NTM. In the mid-2000s improved molecular technology
 108 resulted in the discovery of the two *M. abscessus* subspecies; *M. abscessus* subsp. *massiliense* and
 109 *M. abscessus* subsp. *bolletii* in 2004 and 2006, respectively. Then in 2011 it was proposed that *M.*
 110 *abscessus* subsp. *massiliense* and *M. abscessus* subsp. *bolletii* should be merged into one subspecies,
 111 *M. abscessus* subsp. *massiliense*. This caused some confusion within the medical community, until in
 112 2013, when whole genome sequencing (WGS) showed genetic divisions that clearly identified the
 113 three subspecies within the *M. abs* complex.

114

115 **Speciation of the *M. abscessus* complex:**

116 Over the years, many different biochemical and molecular techniques have been employed to identify
117 NTM species. Up until the early 2000's, the sodium chloride tolerance test was used to identify species
118 of RGM, particularly in distinguishing between *M. abs* and *M. chelonae* species, as *M. abs* is able to
119 grow on Löwenstein-Jensen medium with 5% sodium chloride but *M. chelonae* is not (23). However
120 several investigators reported that this method is unreliable, likely due vague criteria and the cross-
121 over of biochemical features between differing species of RGM (23) (24) (25). The citrate utilization
122 assay perhaps provides more reliability, the premise being that *M. abs* is unable to use citrate as a
123 carbon source whereas other RGM such as *M. chelonae* are (26). As is also the case with the sodium
124 chloride test, this assay takes up to 8 weeks to complete and therefore is losing traction in the clinical
125 setting (23). High Performance Liquid Chromatography (HPLC) has also been used to generate mycolic
126 acid patterns and thus distinguish between RGM species, however this technique has limitations as
127 several RGM have similar mycolic acid profiles (27). Despite its widespread use in species
128 identification, 16S rRNA sequencing has been shown to be inadequate for species identification of
129 mycobacteria (17). An assay with superior specificity was needed to differentiate between NTM
130 species and subspecies.

131

132 ***M. abscessus* and Cystic Fibrosis:**

133 NTM species are ubiquitous in the environment (unlike *M. tuberculosis* and *M. leprae* which require a
134 living host and are transmitted patient to patient or zoonotically), suggesting that NTM exposure is
135 extremely common, whereas NTM disease is still relatively rare. Those with pre-existing lung diseases
136 undoubtedly have some predisposition to NTM infection, leading some to describe a "two-hit" theory
137 of NTM disease acquisition (28). Undoubtedly, the leading population affected by *M. abs* is the CF
138 population. However, there have also been incidences of *M. abs* infections in non-CF populations.

139 CF is an autosomal recessive disorder caused by mutations in the CF transmembrane conductance
 140 regulator gene (CFTR). Despite being a multi-organ disease, one of the most prominent features in CF
 141 is chronic pulmonary infection. The major pathogen associated with lung infection in CF is
 142 *Pseudomonas aeruginosa*, and unfortunately, 80 to 90% of patients with CF die from respiratory
 143 failure as a result of chronic bacterial infection (29). Even from infancy, the lungs of CF patients are
 144 already commonly colonised with a variety of organisms such as *Staphylococcus aureus* and
 145 *Haemophilus influenzae*. Before 1990, NTM infection was not often associated with CF. However, since
 146 then, reports of *M. abs* infection (along with other NTM species) have been increasingly common.
 147 Several large-scale studies have been performed over the past decade or so, revealing an NTM
 148 prevalence in CF patients in some areas as high as 20% (table 1).

Study	Location	Sample size	NTM prevalence in CF
Oliver, KN (2004) (30)	USA	750	13% (majority <i>M. avium</i> complex)
Roux, AL, et. al. (2009) (31)	France	1582	6.6% (<i>M. abs</i> most common)
Seddon, P, et. al. (2013) (32)	UK	3805 adults 3317 children	5% adults 3.3% children
Adjemian, J, et. al. (2014) (33)	USA	18,003	10-20%; depending on area
Mussaffi, H, et. al. (2005) (34)	Israel	139	8.6%

149

150 **Table 1:** Prevalence of non-tuberculous mycobacterial lung disease in cystic fibrosis patients in
 151 differing geographical areas between 2004 and 2014.

152

153 Age is a strong correlator of NTM infection in this group, with 40% of CF patients over the age of 40
 154 having NTM smear positive results, as opposed to 4-20% in the under 40s population (35). Other risk
 155 factors for NTM infection in CF patients appears to be lower body mass index (BMI) values, worse
 156 forced expiratory volume (FEV₁), current infection with *Pseudomonas aeruginosa* and
 157 *Stenotrophomonas maltophilia*, experience of pneumothorax requiring chest drain, the use of inhaled

158 antibiotics and other medical interventions. (36). One study performed in Israel found a significant
159 association between *Aspergillus* species and NTM species in sputum cultures of CF patients (37).

160

161 ***M. abscessus* infection in non-CF populations**

162 It is well documented that a risk factor for NTM pulmonary disease is patients with low body fat. The
163 mechanisms behind this are not well understood, however it is possible that leptin plays a role in NTM
164 predisposition (38).

165 Aside from pulmonary infections, *M. abs* is also able to produce skin and soft tissue infections (SSTIs)
166 in otherwise healthy hosts. There have been cases of *M. abs* outbreaks following the use of
167 contaminated needles and other surgical instruments (39) and even, as was the case in a cohort of
168 'lipotourists' (i.e., people who travel abroad for cosmetic surgery for fat removal), severe outbreaks
169 following cosmetic surgery (40). Interestingly, *M. abs* has also been linked to late-onset wound
170 infections following crush trauma sustained by Swedish survivors of the 2004 tsunami that killed over
171 200,000 people and caused serious crush injuries in another >2000 (41)

172 *M. abs* also causes serious disseminated infections following transplantation (42). A single case study
173 involving post-transplant *M. abs* SSTI resulted in disseminated pulmonary infection and eventually the
174 death of the patient, despite aggressive pre- and peri-operative anti-mycobacterial therapy (43). For
175 this reason, many have recommended that *M. abs* colonisation should be viewed as a contraindication
176 to lung transplantation. This suggestion, however, has been met with criticism. Some studies have
177 shown that it is possible to perform a lung transplant on patients with *M. abs* colonisation and that
178 subsequent clearance of infection is possible, albeit with a strong possibility of severe complications
179 (44) (45). Despite this uncertainty surrounding the outcome of lung transplantation in patients
180 colonised with *M. abs*, it is increasingly clear that effective treatments for *M. abs* lung infection must
181 be developed, as lung transplantation is a potentially life-saving therapy for end-stage lung disease
182 caused by CF and other lung disorders.

183

184 **Environmental reservoirs and transmission:**

185 NTM are ubiquitous in the environment; especially water sources and soil (4). They are prone to
186 biofilm formation and this contributes to their ability to persist in harsh environments (46). NTM can
187 persist in environments that are in close proximity to human populations, particularly human water
188 sources, hospital water supplies (sinks, showerheads), and homes.

189 *M. abs*, like other NTM, is able to survive in harsh, nutrient-starved environments where other
190 competing microorganisms would not survive, such as in chlorinated water (47). The presence of the
191 lipid-rich cell wall results in a hydrophilic cell surface, which facilitates the formation of biofilms, their
192 slow growth and adherence to surfaces, thus aiding their survival and providing them with a selective
193 advantage (48) (49) (50). Furthermore, many RGM are oligotrophic, requiring low levels of two carbon
194 sources and minimal amounts of metal ions (51), further indicating their hardiness and persistence in
195 harsh environments. The impenetrable nature of the *M. abs* cell wall in comparison to other non-
196 mycobacterial pathogens also contributes to its resistance to many antibiotics and disinfectants (52)
197 (12). The ability of *M. abs* to survive in the human environment presents a huge problem for human
198 health, with most studies up until this point suggesting that patients with CF predominately acquire
199 NTM infection from the environment (20). This long-held belief was called into question in 2013 when
200 Floto and his team used WGS to show possible patient to patient transmission of *M. abs* within a CF
201 clinic in the UK (53).

202 In 2009, Feazel *et al* demonstrated that showerheads provide an enriched environment for NTM
203 biofilm formation; the presence of human pathogens including NTM were >100 fold higher in
204 showerhead biofilms compared to the background water contents (54). A study in Hawaii investigated
205 the prevalence of NTM in household plumbing; areas such as showerheads, sinks, taps, shower drains,
206 and refrigerator water dispensers were sampled. The authors found that 69% of households surveyed
207 had clinically significant NTM colonisation, of which 10% was *M. abs* (55). Another 2018 study revealed

208 an outbreak of *M. abs* skin infections in children who were exposed to the same indoor wading pool
209 (56). This study demonstrates the importance of identifying *M. abs* environmental reservoirs,
210 reporting *M. abs* cases and subsequent environmental remediation in order to reduce the risk of
211 infection.

212 The persistence and spread of NTM species within healthcare environments is fast becoming a serious
213 problem and a significant threat to human health (57). It was a long-held belief in the scientific
214 community that NTM is transmitted to humans from the environment, and that patient to patient
215 transmission is unlikely. Resulting in a clinical focus on reducing the risk of environmental transmission
216 using effective sterilising techniques and other hygiene practices. Such as it is, the CF Trust published
217 *M. abs* infection control recommendations that include general infection control measures such as
218 hand washing and more specific recommendations such as segregation of infected patients from other
219 patients (58).

220 The mode of transmission of pathogenic NTM to humans is still poorly understood, with many studies
221 seeking evidence of human to human transmission using molecular techniques such as WGS. A study
222 undertaken in 2001 sought to address this question; a retrospective analysis of 1062 respiratory
223 specimens taken from 214 patients with CF revealed 5 patients with *M. abs* lung infection. These 5
224 patients each had isolates with a unique genotype that was not shared with any of the other patients,
225 which led the authors to conclude that patient to patient transmission of *M. abs* was not occurring
226 within their cohort (59).

227 In 2014, a small-scale study was performed on 27 *M. abs* isolates from 20 paediatric CF patients (60).
228 The authors used a combination of epidemiology, variable number tandem repeat (VNTR) profiling
229 and WGS to find evidence of cross-infection between paediatric CF patients. They hypothesized that
230 patients with strains that had identical VNTR profiles would have had intense exposure to each other
231 compared with patients with strains that had different VNTR profiles. They found little evidence of
232 transmission between patients, except for 2 patients who were siblings and therefore had higher

233 intensity of exposure. They concluded that cross-infection was uncommon in their cohort, and that
234 transmission is most likely to be from a common environmental source (60).

235 The biggest shift in our understanding of transmission came in 2013 when a major study was published
236 in which WGS was used to identify transmission of *M. abs* between patients at an adult CF centre in
237 the UK between 2007 and 2011 (20). The authors found a high level of relatedness between isolates
238 of *M. abscessus* subsp. *abscessus*, but clusters were clearly segregated from one another, indicating
239 that patients have independently acquired either genetically diverse strains or a dominant circulating
240 clone. In the case of *M. abscessus* subsp. *massiliense*, however, the authors found isolates from
241 different individuals with almost identical genomic sequences, strongly indicating transmission
242 between patients. Analysis of the environment revealed no NTM species isolated from the water
243 supply to the clinic, showerheads, dish washers, bronchoscopes or the local River Cam or Papworth
244 Hospital Pond. Further investigation into possible transmission routes revealed patients with isolates
245 from the same genetic relatedness clusters were present in the clinic at the same time as each other,
246 further supporting their hypothesis that *M. abscessus* subsp. *massiliense* is likely transmitted from
247 patient to patient rather than independently from the environment. This finding represents a major
248 clinical advance which may require patients infected with *M. abs* to be segregated from *M. abs*-naïve
249 patients to prevent onward transmission.

250 Following on from the localised retrospective study published in 2013 (20), a global WGS initiative was
251 launched on 1080 isolates from 517 patients from the UK, USA, Republic of Ireland, mainland Europe
252 and Australia (53). This study found that the majority of isolates were from densely clustered
253 genotypes that were not diverse, suggesting a high level of human-human transmission. Phylogenetic
254 analysis also revealed that there are 3 dominant circulating clones globally, and these clones are
255 associated with higher virulence and poor clinical outcomes. Human-human transmission appears to
256 have facilitated the evolution of *M. abs* from an environmental pathogen to a transmissible human
257 pathogen.

258

259 **Diagnosis and treatment:**

260 As *M. abs* and other NTM species are ubiquitous in the environment, including drinking water supplies,
261 the presence of culture-positive respiratory tract sample for NTM does not always indicate NTM-
262 pulmonary disease (NTM-PD). Therefore, patients must also have characteristic symptoms,
263 compatible radiology, and two or more positive sputum samples for the same NTM species, as well as
264 the exclusion of other potential causes of pulmonary disease (61).

265 For clinical laboratory identification of NTM species, the British Thoracic Society (BTS) recommends
266 that isolates be obtained from sputum samples, and if this is not possible (for example in children),
267 bronchoalveolar lavage or transbronchial biopsy samples should be taken when NTM pulmonary
268 disease is suspected (61). NTM infection can be validated in the laboratory, with the use of auramine-
269 phenol staining and microscopy, as well as culture on solid and liquid media.

270 All clinical isolates of *M. abs* undergo susceptibility testing for clarithromycin, ceftazidime and amikacin.
271 They also recommend that other antibiotics such as tigecycline, imipenem, minocycline, moxifloxacin
272 and clofazimine are tested in this manner (61).

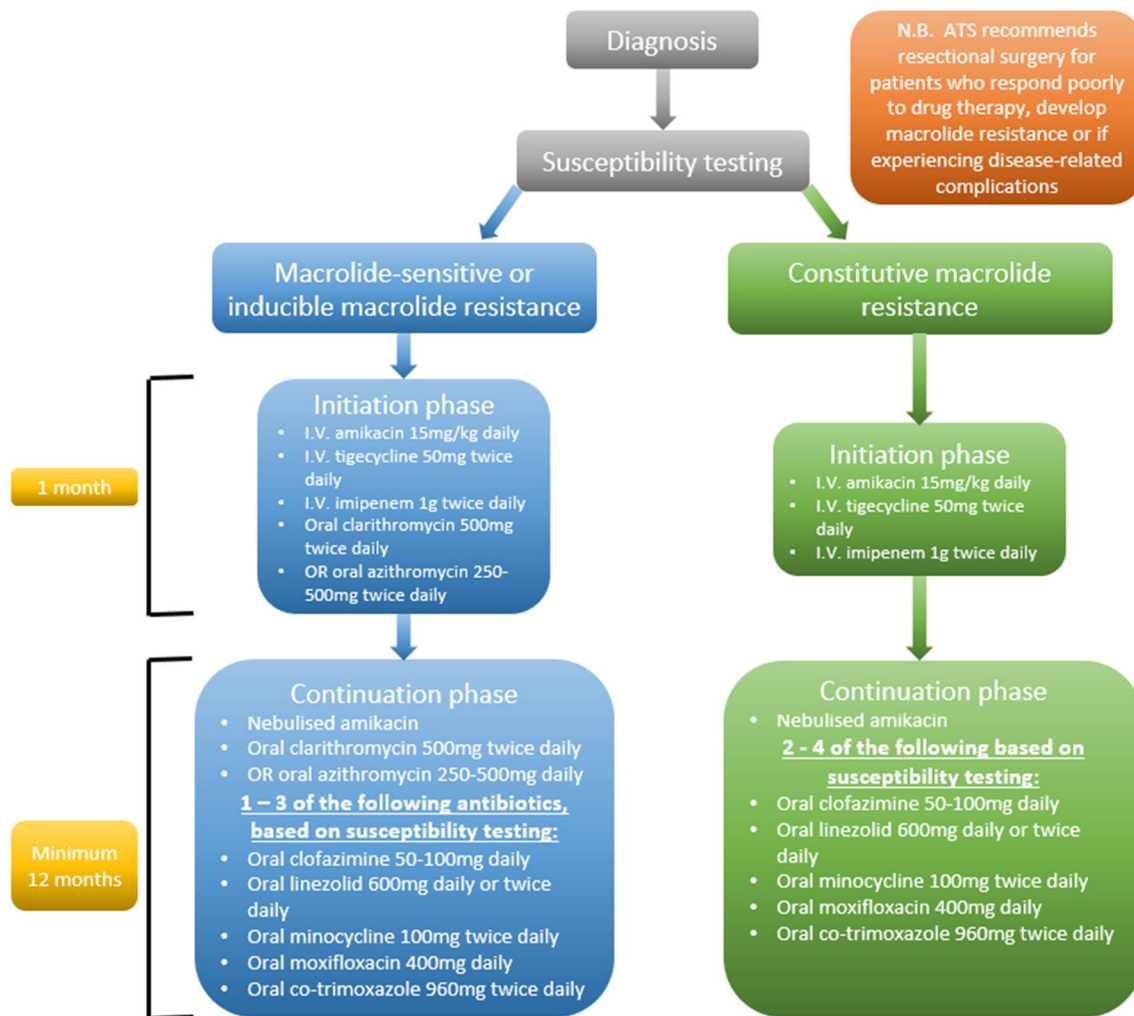
273 **Treatment**

274 When *M. abs* was first isolated in 1952, it was thought the patient was initially infected with the
275 pathogen at the age of 14 years old. The patient's condition resolved without intervention and so for
276 some time, treatment wasn't considered a priority in *M. abs* infections (6).

277 Of course, today it is well known that treatment for *M. abs* pulmonary infection is essential to give the
278 patient the best chance of survival. Unfortunately, antimicrobial chemotherapy for *M. abs* infection is
279 particularly difficult due to its intrinsic and acquired resistance to most of the commonly used
280 antibiotic classes. Further complications in the treatment of *M. abs* infection is the lack of evidence
281 that *in vitro* susceptibility of antibiotics corresponds to *in vivo* efficacy in treating pulmonary disease

282 (62). Because chemotherapy-based treatment of *M. abs* infection is often unsuccessful, the American
283 Thoracic Society advises that certain patients may have the best chance of disease regression with
284 resectional surgery, especially if the patient exhibits a poor response to drug therapy, if macrolide-
285 resistance develops, or if the patient is experiencing disease-related complications such as
286 haemoptysis (62).

287 Current treatment guidelines from the BTS (61) recommend that treatment for *M. abs* pulmonary
288 disease should consist of an initial phase antibiotic regimen that includes intravenous (I.V.) and oral
289 antibiotics, followed by a continuation phase comprising of oral and inhaled antibiotics (Figure 2).
290 Further genetic analysis of clinical isolates can provide information on the *erm*(41) (inducible
291 macrolide resistance) and/or presence of 23S rRNA point mutation (constitutive macrolide resistance)
292 in clinical isolates of *M. abs*, which can then be used to inform patient-specific treatment regimens.



293

294 **Figure 2:** Flow chart showing treatment regimen for *M. abs*-pulmonary disease based on laboratory
 295 susceptibility testing results as recommended by the British Thoracic Society. Treatment will differ
 296 based on the whether the isolate displays macrolide sensitivity/inducible macrolide resistance or
 297 constitutive macrolide resistance. The initial phase of treatment involves three intravenous (I.V.)
 298 antibiotics, and for macrolide sensitive/inducible macrolide resistance 1 of 2 oral macrolides, and this
 299 phase lasts one month. The continuation phase also depends on laboratory susceptibility testing
 300 results and clinicians will typically administer 1-4 oral antibiotics over a period of at least 12 months.
 301 It is also important to note that the American Thoracic Society recommends surgical resection of
 302 infected area if the patient is not responding to therapy, if macrolide resistance develops, and/or if
 303 the patient develops disease-related complications such as haemoptysis.

304 Side effects of *M. abs* treatment are common and can be severe. A retrospective analysis of 65
305 patients undergoing treatment for *M. abs* lung disease in South Korea (63) revealed frequent adverse
306 reactions to cefoxitin; 51% of patients developed leukopenia, 6% of patients developed
307 thrombocytopenia, and 15% of patients experienced drug-induced hepatotoxicity. As a result,
308 cefoxitin was discontinued in 60% of patients and side effects resolved. Another common side effect
309 observed was gastrointestinal problems (nausea, anorexia, or diarrhoea), which affected 22% of
310 patients and caused 4 patients (6%) to completely stop antibiotic treatment. A clinical
311 recommendation was made to consider imipenem as an alternative to cefoxitin, however prolonged
312 treatment with imipenem can cause neutropenia.

313 Another study that analysed treatment outcomes in 65 patients with *M. abs* in North America also
314 found a high prevalence of side effects. IV amikacin (65% of patients) and azithromycin (71% of
315 patients) were the most commonly used antimicrobials in this cohort. They found 74 different side
316 effects reported in 62% of patients, most commonly nausea/vomiting (31%) and skin changes (20%).
317 They attributed many of these side effects to amikacin or tigecycline, and as a result, of those received
318 amikacin or tigecycline therapy, 51% and 36% of patients, respectively, had to adjust or stop
319 medication due to severe side effects such as ototoxicity. Similar to the South Korean study, 4 patients
320 had to totally stop treatment because of their side effects (64).

321 Clarithromycin is one of the most commonly used antibiotics to treat *M. abs* (35). However,
322 clarithromycin has been associated with hearing loss, with one study citing a 7% hearing loss rate in
323 their patients. This side effect did resolve in all but one patient, but the authors state that the patient
324 had a pre-existing condition that hindered their ability to attribute this hearing loss solely to
325 clarithromycin (65). A case study on an 81-year-old woman, who was being treated with
326 clarithromycin for infective exacerbation of chronic pulmonary obstructive disease (COPD) showed
327 another example of clarithromycin-related permanent hearing loss, despite evidence that
328 clarithromycin is relatively well tolerated (66) (67). The major issue with using clarithromycin to treat

329 *M. abs* is the presence of a functional inducible *erm(41)* gene that confers macrolide resistance in both
 330 *M. abscessus* subsp. *abscessus* and *M. abscessus* subsp. *bolletii* but not *M. abscessus* subsp.
 331 *massiliense*.

332

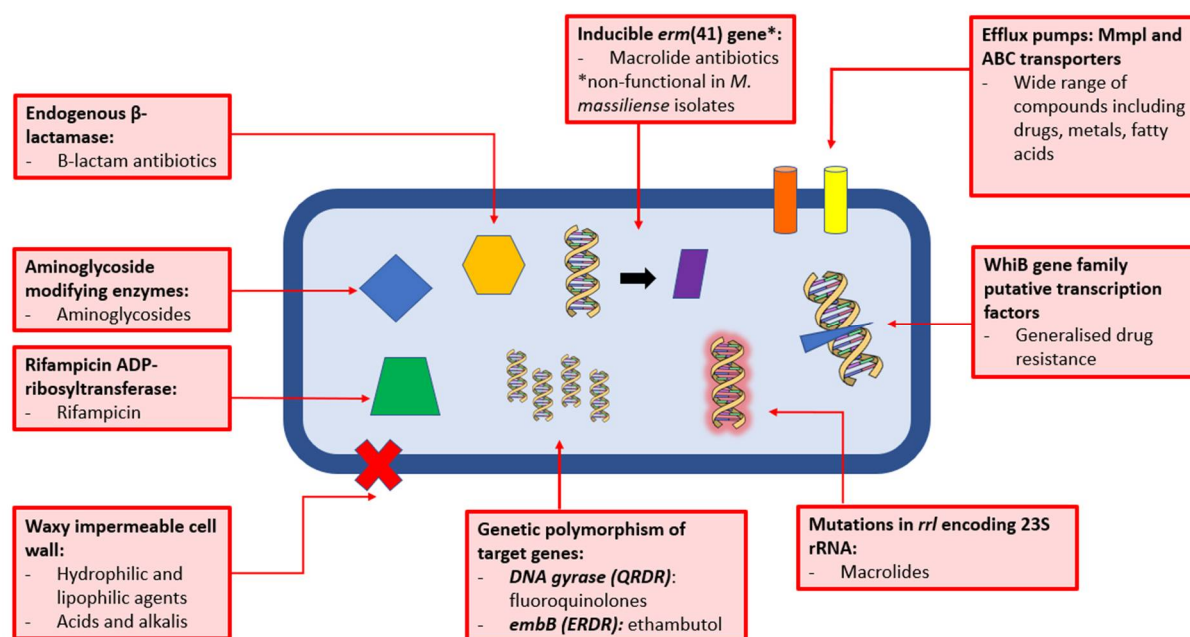
333 Future perspectives for *M. abscessus*

334

335 The resistance problem: why the drugs don't work:

336 *M. abs* is known for its intrinsic resistance to most chemotherapeutic agents, including all the anti-
 337 tuberculous drugs used to treat *M. tuberculosis* infection (68) (69). Furthermore, *in vitro* drug
 338 susceptibility testing on *M. abs* often proves unhelpful in guiding treatment regimens (70). There are
 339 a number of natural resistance mechanisms displayed by *M. abs* (along with other mycobacteria),
 340 including a waxy and impermeable cell wall, drug export systems, antibiotic modifying/inactivating
 341 enzymes, and genetic polymorphism of target genes (12).

342



343

344

345 **Figure 3:** Graphical summary of the resistance mechanisms exhibited by *Mycobacterium abscessus*
346 (*M. abs*). There are several mechanisms involving different physiological, enzymatic and genomic
347 processes that contribute to the notoriously drug-resistant profile of *M. abs*. It is likely that these
348 processes work in synergy to produce a highly resistant pathogen, such as efflux pumps and drug
349 resistance genes.

350

351 The greatest contributing factor to the lack of *M. abs* sensitivity to many major classes of antibiotic is
352 the mycobacterial cell wall, the role of which has long been studied. The high lipid content and unusual
353 thickness of the mycobacterial cell wall provides an effective barrier for hydrophilic and lipophilic
354 agents (71). In 1990 it was shown that the lack of permeability of the *M. chelonae* (then grouped
355 together with *M. abs*) cell wall plays a vital role in making the pathogen resistant to antibiotics (52).
356 The cell wall barrier is also responsible for *M. abs*' intrinsic resistance to acids and alkalis (72). The cell
357 wall of mycobacteria also contains porins, it was shown in 1990 that *M. chelonae* possesses a 59 kDa
358 cell wall protein that allows for the diffusion of small, hydrophilic solutes. This porin, however, is
359 minor, unlike that of *E. coli* where they are the most abundant cell wall protein, explaining the low
360 permeability to hydrophilic solutes (11). The cell wall cannot explain all of the intrinsic drug resistance
361 seen in *M. abs*, in fact it is known that the cell wall, particularly the porins, act synergistically with
362 internal systems that are activated by the presence of intracellular antibiotics, and that the low
363 permeability of the mycobacterial cell wall means that the bacteria has time to induce the expression
364 of drug resistance genes (73).

365 As a constituent of the mycobacterial cell wall, active efflux pumps can be described as one of the
366 main causative factors of drug resistance in mycobacteria (12) (74) (75). They primarily act to protect
367 bacteria against toxic compounds and bacterial homeostasis by transporting toxins or metabolites to
368 the extracellular environment (75). *M. abs* encodes protein members of the major facilitator family
369 ABC transporters as well as mycobacterial membrane protein large (MmpL) families (76). ABC

370 transporters are found in all forms of life and make use of adenosine triphosphate (ATP) to transport
371 molecules across membranes. The MmpL transporter family is a subclass of a large family of multidrug
372 resistance pumps known as Resistance-Nodulation-Cell-Division (RNCD) permeases. MmpLs export
373 lipid components across the cell envelope of mycobacteria (77). The role of MmpLs in *M. abs* drug
374 resistance is yet to be fully understood, however there is evidence that MmpL7 in *M. tuberculosis*
375 confers resistance to isoniazid (78), suggesting that MmpLs may play a major role.

376 Macrolides are one of the mainstays of *M. abs* treatment (35), yet despite this, *M. abs* infections tend
377 to respond poorly to macrolide therapy, even when they appear sensitive to clarithromycin *in vitro*
378 (79). A study performed in 2009 revealed the presence of an inducible *erm(41)* gene in 7 out of 10 *M.*
379 *abs* clinical isolates that confers resistance to macrolides with a minimum inhibitory concentration
380 (MIC) of ≥ 32 $\mu\text{g}/\text{mL}$. The 3 remaining susceptible isolates had *erm(41)* gene, however it appeared to
381 be non-functional (79). The *erm(41)* gene produces a functional 23S rRNA methylase, contributing to
382 macrolide resistance along with point mutations in the *rrl* encoding 23S rRNA gene (80). Following on
383 from this, it was shown that macrolides may be useful in treating approximately 20% of *M. abs*
384 infections in the U.S., and that sequencing of the *erm(41)* gene is a potentially useful tool in predicting
385 macrolide susceptibility (81). It is also noteworthy that *M. abscessus* subsp. *massiliense* contains a
386 large 97 base pair deletion in *erm(41)*, rendering it useless and therefore meaning *M. abscessus* subsp.
387 *massiliense* retains susceptibility to macrolides, except in the case of *rrl* mutants (82) (79) (83) (84). *M.*
388 *abs* isolates possessing an *rrl* mutant display constitutive resistance to macrolide antibiotics. This
389 phenomenon is known to be mediated by a mutation in *rrl* encoding the bacterial 23S rRNA gene,
390 particularly at positions 2058 and 2059, i.e. the drug binding pocket of the gene (85).

391 If macrolide therapy is not advised due to evidence of constitutive resistance, there are of course other
392 chemotherapeutic options available. However, in many of the conserved genes in *M. abs* that can
393 potentially act as drug targets there is the presence of genetic polymorphisms, which can often confer
394 drug resistance (12).

395 A 1998 study showed revealed an amino acid substitution at position 83 (Ser83Ala) in the quinolone-
396 resistance-determining-region (QRDR) in fluoroquinolone-resistant isolates of *M. abs* (86). This
397 substitution occurs in the region of DNA gyrase subunit *GyrA* that binds DNA, and as fluoroquinolones
398 bind strongly to the gyrase-DNA complex, and weakly to protein or DNA alone, this mutation results
399 in fluoroquinolone resistance (87). Genetic polymorphisms also occur within the *emb* operon that
400 codes for several homologous arabinosyl transferases. These are enzymes involved in the
401 polymerisation of arabinogalactan, an essential component of the mycobacterial cell wall and can be
402 inhibited by the tuberculosis drug ethambutol. A 1997 study showed that polymorphisms at position
403 306 in a highly conserved *embB* gene conferred natural resistance across many species of
404 mycobacteria, including *M. abs* (88). *M. abs* has high natural levels of resistance to ethambutol (MIC
405 >64mg/L), and the same study transferred the *M. abs emb* region to ethambutol-susceptible *M.*
406 *smegmatis* resulted in a 500-fold increase in the MIC to ethambutol (88).

407 *M. abs* also produces a number of target-modifying enzymes. Rifampicin ADP-ribosyl transferase,
408 *Arr_Mab* inactivates rifamycins such as rifampicin. Aminoglycoside 2'-N-acetyltransferase and
409 aminoglycoside phosphotransferases mediate the susceptibility to aminoglycoside antibiotics. *M. abs*
410 has also been shown to produce an endogenous β -lactamase (*Bla_{Mab}*), that efficiently hydrolyses the
411 β -lactam ring of β -lactam antibiotics, rendering them ineffective (89).

412 Aside from antibiotic-specific internal drug resistance mechanisms, a family of transcriptional
413 regulators, the *WhiB* family, is exclusive to actinomycetes and may be involved in conferring drug
414 resistance in *M. abs*. Members of this family have been shown to regulate systems of drug resistance
415 in *M. tuberculosis*, including antibiotic export and activation (90). *M. abs* has been shown to possess
416 a homologue of the *M. tuberculosis WhiB7*. When *M. abs WhiB7* is deleted, the result is increased
417 sensitivity to clinically relevant antibiotics that target the ribosome, such as clarithromycin, amikacin
418 and tetracycline (91).

419

420 **Future treatments:**

421 It perhaps goes without saying that there is an urgent, unmet need for safe and effective treatments
422 against *M. abs* pulmonary disease. There have been instances of successful treatment of *M. abs* with
423 already available antibiotics. One such case was reported in 2002, where a 63-year-old patient whose
424 infection had not responded to the traditional regimen was prescribed a course of faropenem, a new
425 member of the β -lactam antibiotic class. Treatment was successful and produced no adverse side
426 effects (92). It is not just antimicrobials that have potential in enhancing *M. abs* treatment. In 2012,
427 Okazaki *et. al.* reported that the use of clarithromycin, amikacin and imipenem/cilastatin to treat a
428 case of *M. abs* pulmonary was greatly enhanced with the addition of corticosteroids. The authors
429 recommend that the presence of organising pneumonia (a non-specific inflammatory pulmonary
430 process) or an allergic reaction may have helped to explain the poor response to antibiotic treatment
431 alone in some patients, and that this possibility should be considered when applicable to improve
432 treatment outcomes (93).

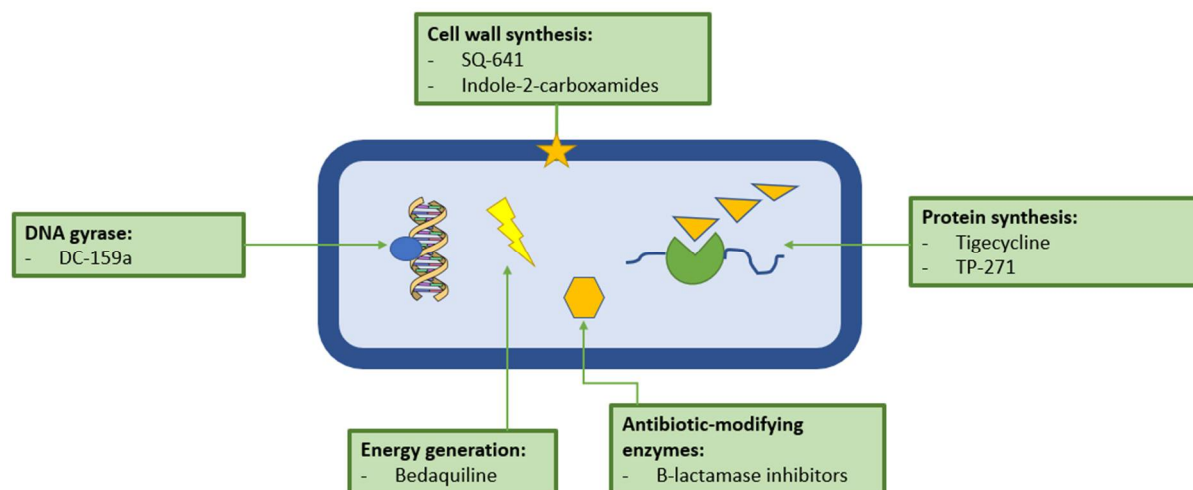
433 One of the enzymatic resistance mechanisms employed by *M. abs* is the production of an endogenous
434 β -lactamase, Bla_{Mab} (Figure 3). Cefoxitin and imipenem, both β -lactam antibiotics, are commonly used
435 to treat *M. abs*. In order to improve the efficacy of these antibiotics, a β -lactamase inhibitor may be
436 administered in conjunction during therapeutic treatment. A 2015 study revealed that avibactam, a
437 β -lactamase inhibitor is able to efficiently inhibit Bla_{Mab} (94), and a subsequent 2017 study showed
438 that avibactam improves the efficacy of imipenem against *M. abs* both *in vitro* and in macrophage,
439 and zebrafish models of infection (95).

440 Aside from these examples, very few case studies have reported successful treatment with
441 repurposed antibiotics. Therefore, novel drug targets in *M. abs* must be discovered and elucidated,
442 and novel compounds that safely and effectively inhibit these targets discovered.

443 There are potentially a wide variety of viable drug targets in *M. abs* (Figure 4) Many of the most
444 promising leads against *M. abs* have come about as a result of concerted effort to find novel drugs for

445 *M. tuberculosis*, which a handful of researchers have applied to *M. abs* and other NTM species.
 446 Unfortunately, only a small percentage of the novel drugs which are active against *M. tuberculosis*,
 447 are also active against *M. abscessus*, further highlighting just how resistant and dangerous this
 448 pathogen is proving to be.

449



450

451 **Figure 4:** Graphical summary of the exploitable drug targets in *Mycobacterium abscessus* (*M. abs*).

452 There are several potential target areas in *M. abs* including physiological, genomic, enzymatic and
 453 metabolic processes. Many of the drugs with potential to be used as part of *M. abs* treatment are
 454 old classes of antibiotics that have been repurposed, such as β -lactamase inhibitors, or have been
 455 discovered as part of the anti-tuberculous drug discovery pipelines, such as bedaquiline.

456

457 One potential target in *M. abs* is DNA gyrase, despite the fact that *M. abs* is naturally resistant to
 458 quinolones (96), a novel fluoroquinolone, DC-159a was developed in 2010 as part of the Working
 459 Group on TB Drugs, and was found to be active against *M. abs* with an MIC of 16 $\mu\text{g}/\text{mL}$, which was 4
 460 to 8-fold lower than the other already available quinolones tested (97). The authors stressed the
 461 importance of *in vivo* testing of DC-159a, however, no publications attesting to the *in vitro* activity of
 462 DC-159a against *M. abs* have been released to date.

463 The mycobacterial cell wall, in all its complexity, can offer an attractive range of potential antibiotic
464 targets. The three distinct layers of the mycobacterial cell wall: core peptidoglycan, arabinogalactan
465 and mycolic acids are each essential to the pathogen and involve a number of exploitable processes
466 (98). A 2010 study subjected several species of NTM to a capuramycin analogue SQ641 (99).
467 Capuramycins are a novel class of nucleoside antibiotics that work by targeting phosphor-*N*-
468 acetylmuramyl-pentapeptide-translocase (translocase-1 or TL-1) which is essential for peptidoglycan
469 synthesis. They found that the drug had an MIC of 0.25-1 µg/mL, as well as finding synergy between
470 SQ641 and rifabutin and streptomycin. This drug has great potential as it is fast-acting and displays a
471 long post-antibiotic effect (100). In 2017 a study was published in which several members of the newly
472 synthesized MmpL3 inhibitors, indole-2-carboxamides, have shown potent activity against *M. abs*.
473 These inhibitors have been shown to work by inhibiting the transfer of mycolic acids to their cell
474 envelope acceptors in *M. abs* strains (101). Further work has been done on this class of inhibitors; in
475 2019, Pandya *et. al.* reported that oral administration of the inhibitors shows a statistically significant
476 reduction in bacterial load in the lungs and spleens of *M. abs*-infected mice (102).

477 It has been demonstrated that *M. abs* displays high levels of intrinsic resistance to the tetracycline
478 class of antibiotics via the monooxygenase, MabTetX, a *WhiB7*-independent pathway (103). This is not
479 the end of the road for this class of antibiotics. Tigecycline, the first developed glycylcycline, a new
480 class of tetracycline antibiotics originally developed for SSTIs, was shown in 2014 to be highly effective
481 *in vivo* against *M. abs* pulmonary disease (104). Further work in 2018 revealed that tigecycline is a
482 poor substrate of MabTetX and is incapable of inducing its expression, explaining its high efficacy in
483 comparison with other tetracycline antibiotics (103). Tigecycline is now of the recommended
484 treatment options for *M. abs* pulmonary disease, and is arguably one of the most effective, with one
485 study citing clinical improvement in >60% patients with *M. abs* pulmonary disease when tigecycline is
486 employed as part of the multi-drug regimen against *M. abs* (104). Tigecycline is not the only
487 tetracycline showing activity against *M. abs*. A 2012 study tested the *in vitro* activity of a novel
488 fluorocycline antibiotic, TP-271 (a tetracycline-related antibiotic) against 22 isolates of *M. abs*. They

489 found all the isolates to have an MIC of ≤ 1 $\mu\text{g}/\text{mL}$ with an average of 0.5 $\mu\text{g}/\text{mL}$, which is decidedly
490 superior than that of the other orally available tetracycline antibiotics such as moxifloxacin and
491 tetracycline (105).

492 Bedaquilin, the latest drug indicated for the treatment of multi-drug resistant TB (MDR-TB) was
493 approved by the FDA in 2011, and it works by targeting the ATP synthase of mycobacteria. Obregon
494 *et. al.* (106) demonstrated an MICs of 1.0 $\mu\text{g}/\text{mL}$ against *M. abs* reference strain and then in 2017,
495 Vesenbeckh and colleagues pointed to bedaquiline as a potential antimicrobial against *M. abs* after
496 the drug exhibited MICs of ≤ 1 $\mu\text{g}/\text{mL}$ against 20 *M. abs* clinical isolates *in vivo*. (107)

497

498 **Summary:**

499 *M. abs* is increasingly being recognised as an important pathogen responsible for a wide range of
500 infections and implicated in severe, and often untreatable pulmonary infections in people with CF and
501 other structural lung disorders. Almost all of the currently available antibiotics are useless against the
502 pathogen, with even official guideline treatment regimens having little to no evidence of *in vivo*
503 efficacy. With such high treatment failure rates, clinicians are often forced to administer last-resort
504 antibiotics in the hope of a cure. Coupled with increasing prevalence and its already extensively drug
505 resistant profile, it is glaringly obvious that novel, effective and safe treatments are needed. Many of
506 the novel drugs mentioned above are in various phases of clinical trial against *M. tuberculosis* and
507 there is a significant paucity of data regarding their efficacy against *M. abs* and other NTM species.
508 Furthermore, there is a startling lack of *in vivo* efficacy data for any of these drugs, which is particularly
509 worrying considering the inconsistencies between *in vitro* and *in vivo* anti-*M. abs* activity. Whilst TB
510 has many dedicated drug-discovery programmes, NTM has none. A dedicated NTM drug discovery
511 pipeline is essential to ensure the disease burden of NTM does not become overwhelming.

512

513 **Author Contributions:**

514 R. C. L., J. H., M. D. and J. A. G. C. reviewed the literature, intellectually conceived and wrote the
515 manuscript.

516

517 **Funding:** This research was funded by Birmingham Women's and Children's Hospital Charity Research
518 Foundation (BWCHCRF) (R. C. L. 50% PhD Studentship, match funded by Aston University Prize
519 Scheme) and the Academy of Medical Sciences and Global Challenges Research Fund with a
520 Springboard Grant (SBF003\1088:).

521

522 **Acknowledgements:** J. A. G. C. is grateful to the Academy of Medical Sciences, Global Challenges
523 Research Fund and Birmingham Women's and Children's Hospital Charity Research Foundation
524 (BWCHCRF) for their continued support of the Mycobacterial Research Group at Aston University.

525

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