## 1 Thiol Stress Fuels Pyrazinamide Action Against *Mycobacterium* 2 *tuberculosis*

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14 Author Contributions: Lev Ostrer, Taylor A. Crooks and Anthony D. Baughn conceived of the 15 study. Lev Ostrer and Taylor A. Crooks designed and performed microbiology-related experiments, 16 data analysis and interpretation. Sang Vo synthesized and performed quality control for the study 17 compound. Michael D. Howe and Taylor A. Crooks developed Tn-seq and RNA-seq pipelines and 18 conducted computational analyses. Pooja Hegde performed mass spectrometry data acquisition 19 and analysis. Ziyi Jia constructed and validated targeted mutations in mycobacterial strains and 20 conducted growth inhibition studies with these strains. Courtney C. Aldrich provided critical 21 resources and expertise for synthesis and analysis of the study compound and its products. Lev 22 Ostrer and Taylor A. Crooks prepared all display images. All authors discussed the results and 23 contributed to writing and editing of the final manuscript, with major contributions by Anthony D. 24 Baughn, Lev Ostrer and Taylor A. Crooks.

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### 28 Abstract

29 Pyrazinamide (PZA) is a cornerstone of first-line antitubercular drug therapy and is unique in its 30 ability to kill nongrowing populations of Mycobacterium tuberculosis through disruption of 31 coenzyme A synthesis. Unlike other drugs, PZA action is conditional and requires potentiation by 32 host-relevant environmental stressors, such as low pH and nutrient limitation. Despite its pivotal 33 role in tuberculosis therapy, the mechanistic basis for PZA potentiation remains unknown and the 34 durability of this crucial drug is challenged by the emergent spread of drug resistance. To advance 35 our understanding of PZA action and facilitate discovery efforts, we characterized the activity of a 36 more potent PZA analog, morphazinamide (MZA). Here, we demonstrate that like PZA, MZA acts 37 in part through impairment of coenzyme A synthesis. Unexpectedly, we find that, in contrast to 38 PZA, MZA does not require potentiation due to aldehyde-mediated disruption of thiol metabolism 39 and maintains bactericidal activity against PZA-resistant strains. Our findings reveal a novel dual action mechanism of MZA that synergistically disrupts coenzyme A synthesis resulting in a faster 40 41 rate of killing and a higher barrier to resistance relative to PZA. Together, these observations 42 resolve the mechanistic basis for potentiation of a key first-line antitubercular drug and provide new 43 insights for discovery of improved therapeutic approaches for tuberculosis.

## 44 Significance Statement

45 Pyrazinamide is the only antitubercular drug of its kind, capable of targeting persistent 46 Mycobacterium tuberculosis through disruption of the coenzyme A biosynthetic pathway. A peculiar 47 feature of this drug is that its activity is conditional, requiring low pH for its action. Despite decades 48 of investigation, the precise basis for this conditional susceptibility has remained elusive which has 49 been a barrier to discovery of more effective next-generation analogs. Using approaches in 50 chemical biology, functional genomics and bacterial physiology we demonstrate that activation of 51 thiol stress is the basis for pyrazinamide potentiation. These findings resolve a long-standing question regarding the mechanistic basis for conditional PZA susceptibility of *M. tuberculosis* and 52 53 reveal novel avenues for antimicrobial drug discovery efforts.

## 54

# 55 Main Text

### 57 Introduction

58 59 For several decades, pyrazinamide (PZA) has been an integral part of the first-line 60 standard short course therapy for tuberculosis (TB)<sup>1</sup>. A unique feature of PZA is its ability to kill 61 non-replicating populations of the causative agent Mycobacterium tuberculosis<sup>2</sup>. Use of PZA has 62 played a pivotal role in reducing TB relapse rates and shortening the duration of treatment from 9 63 to 6 months<sup>1</sup>. In spite of this success, wide use of this treatment regimen has resulted in the 64 emergent spread of PZA resistance. Current rates of PZA resistance in M. tuberculosis clinical 65 isolates range from 16 to 42% depending upon patient cohort<sup>3</sup>. The primary mechanism of clinical 66 PZA resistance occurs through spontaneous loss-of-function mutations in the M. tuberculosis pncA 67 gene which encodes an amidase that is required for activation of PZA to pyrazinoic acid (POA)<sup>4,5</sup>. 68 Identification of potent next-generation analogs of PZA that circumvent resistance will be critical to 69 maintain use of this important therapeutic tool.

70 Despite its crucial role in TB therapy the antitubercular mechanism of PZA action is not 71 fully defined. Recent evidence demonstrates that PZA acts through disruption of coenzyme A (CoA) 72 biosynthesis, in part, through destabilization of L-aspartate decarboxylase (PanD) by POA<sup>6,7</sup>. 73 Intriguingly, while CoA plays an indispensable role in M. tuberculosis metabolism, PZA 74 susceptibility is conditional and requires activation of the cell envelope stress response through 75 exposure of bacilli to low pH or a variety of other stressors<sup>8</sup>. How disruption of CoA biosynthesis 76 connects with conditional susceptibility of *M. tuberculosis* to PZA has not been resolved<sup>9</sup>. Yet, it 77 has recently been shown that exposure of *M. tuberculosis* to low pH results in thiol stress<sup>10</sup> which 78 likely interferes with CoA synthesis through limitation of cysteine, an essential pathway substrate.

Of note, consistent with an essential role for environmentally driven potentiation of PZA action *in vivo*, PZA shows no detectable antitubercular activity in mice that lack T cells and are thereby impaired for macrophage activation which is required for full phagosomal acidification<sup>11</sup>. Identification of means to bypass the need for host-mediated potentiation of PZA action would advance antitubercular therapy through promoting drug potency in the context of compromised immunity which is causally associated with progression of TB<sup>9</sup>.

85 To address the unmet needs outlined above, we set out to characterize the mechanism of 86 action of an intriguing PZA analog, morphazinamide (MZA, Figure 1A). Early clinical studies 87 demonstrated efficacy of MZA against *M. tuberculosis* in humans with equivalent potency and 88 safety profiles relative to PZA<sup>12,13</sup>. Further, in vitro assays indicated that the antitubercular activities 89 of MZA and PZA were comparable, with similar levels of growth inhibition in broth culture<sup>14</sup>. 90 However, it was noted that MZA and other aminomethylene analogs did not require low pH to show 91 inhibitory activity and retained activity against PZA resistant isolates, suggesting that they operate 92 by a distinct mechism<sup>14</sup>. In this study we applied approaches in chemical biology, functional 93 genomics and bacterial physiology to characterize the mechanism of action of MZA. Similar to its 94 sustained activity against pncA null strains of M. tuberculosis, we find that MZA also retains activity 95 against *M. tuberculosis* strains with newly described PZA resistance mechanisms<sup>8</sup>. By using 96 transcriptional profiling and transposon sequencing (Tn-seq) we find many parallels between 97 responses of *M. tuberculosis* to PZA, POA and MZA that highlight the impact of these drugs on 98 CoA metabolism. Importantly, we uncover an association between thiol metabolism and 99 susceptibility to these drugs and find that MZA susceptibility in particular is strongly influenced by 100 the mycothiol reductase MscR due to release of aldehyde. Together, our findings reveal a central 101 role for thiol stress as a principal driver of PZA action which resolves the basis for conditional 102 susceptibility and illuminates new opportunities to improve potency and raise the resistance barrier 103 to PZA analogs.

## 104

## 105 **Results** 106

107 Impact of MZA on PZA susceptible and resistant *M. tuberculosis*. To assess the antitubercular 108 activity of MZA, we performed minimum inhibitory concentration assays (MICs) using a panel of 109 strains with differing mechanisms of PZA resistance and under various conditions that influence 110 PZA susceptibility. We began by comparing activity between PZA and MZA using standard growth 111 conditions at circumneutral pH with both H37Ra and H37Rv backgrounds. Consistent with the 112 known conditional susceptibility of M. tuberculosis to PZA, cultures grew unimpaired at concentrations as high as 4.5 mM in medium at circumneutral pH (Figure 1B; Table S1). In striking 113 114 contrast, under these same conditions we observed >90% inhibition of growth in the presence of 115 280 µM MZA (Figure 1B; Table S1). Next, we assessed whether MZA activity could be enhanced 116 by conditions that potentiate PZA and diminished under conditions that antagonize PZA. To 117 evaluate potentiation. MICs were determined under acidic (pH 5.8) conditions (Figure 1C.D). To 118 determine if MZA activity could be antagonized similarly to PZA, the effect of the CoA intermediate 119 pantothenate on drug action was evaluated. Consistent with previous work, PZA was fully inhibitory 120 at 500 µM under acidic conditions (Figure 1C,D; Table S1), while the addition of 230 µM 121 pantothenate fully antagonized PZA action<sup>15</sup> (Figure 1E; Table S1). Acidic conditions were able to 122 enhance MZA activity by two-fold, lowering its MIC to 140 µM (Figure 1C,D; Table S1), whereas 123 addition of pantothenate caused a slightly antagonistic effect on MZA activity at both acidic and 124 circumneutral pH (Figure 1E; Table S1). These observations indicate that MZA is mildly impacted 125 by conditions that strongly influence PZA action.

We next sought to determine whether MZA retained activity against several previously described PZA resistant isolates. These strains included mutants deleted for *pncA* (required for activation of PZA to POA) and *sigE* (required for potentiation of PZA susceptibility at low pH), as well as a *clpC1* promoter mutant and a strain with truncation of the 3' end of *panD*<sup>8</sup>, both of which are thought to interfere with POA action through stabilization of PanD<sup>7</sup>. Despite PZA showing diminished activity against this panel of strains, MZA was highly active with MICs ranging from 140 to 280  $\mu$ M (Figure 1F,G; Table S1). It is worth noting that the  $\Delta pncA$  strain showed a two-fold

increase in its MZA MIC relative to the parental strain (Figure 1G; Table S1) indicating a minorcontribution of PncA to MZA action.

135 One of the key features of PZA is its unique ability to elicit a bactericidal effect on non-136 growing *M. tuberculosis*<sup>16</sup>. Thus, we sought to determine whether MZA also has bactericidal activity 137 against actively growing and metabolically quiescent drug tolerant bacteria by performing kill curves 138 under nutrient rich and nutrient limiting conditions. When mid-exponential cultures under either 139 acidic or circumneutral pH growth conditions were treated with 0.9 mM MZA, greater than 99.999% 140 killing was observed after 4 days (Figure 1H,I). After 14 days of incubation following a single 141 treatment, no surviving CFU were observed. We next tested the bactericidal activity of PZA and 142 MZA under nutrient limiting conditions by exposing starved bacteria to either 0.9 mM PZA or MZA. 143 After 56 days of stasis, the majority of untreated bacilli survived, whereas there was more than 144 99.9% loss in viability of bacilli exposed to 0.9 mM PZA (Figure 1J). In contrast to PZA, MZA 145 bactericidal activity was transient, with 80% killing in the first 2 days followed by a plateau (Figure 1J). To determine whether the plateau in killing was due to drug inactivation or due to enrichment 146 147 of a drug tolerant population, starved cultures were washed and treated with fresh MZA every 3 148 days. As before, we observed 80% killing after 3 days for both H37Rv (Figure 1K) and H37Rv 149  $\Delta pncA$  (Figure 1L). Following the second treatment over 99.5% of bacteria were no longer viable 150 for both strains (Figure 1K,L). The third treatment resulted in sterilization of bacteria with no 151 detectible CFU for either strain (Figure 1K,L). These data indicate that MZA has a potent PncA-152 independent bactericidal activity against both actively growing and metabolically quiescent M. 153 tuberculosis.

To assess the antibacterial spectrum of MZA action, we evaluated its inhibitory activity against a panel of human-associated ESKAPE pathogens<sup>17</sup>. All ESKAPE pathogens that were tested showed MICs greater than 10-fold higher than those for *M. tuberculosis* (Figure 1M,N), suggesting that like PZA, MZA activity is highly selective for mycobacterial species.

158 Transcriptional responses of M. tuberculosis to PZA, POA and MZA exposure. To gain a better understanding of the impact of PZA, POA and MZA on *M. tuberculosis* physiology and to 159 160 determine similarities and differences in global responses of bacilli to these drugs, we performed 161 genome-wide transcriptional profiling using RNA-seq. Since PZA is only active at low pH, all 162 treatments were performed using M. tuberculosis grown in acidic (pH 5.8) media. Cultures of 163 exponentially growing H37Rv were exposed to 200 µM PZA, POA, MZA or vehicle only (DMSO) 164 for 24 hours. The resulting transcriptional profiles revealed strikingly similar results (Figure 2A-C; 165 Figure S1A). Specifically, we observed a significant overexpression of early genes in the CoA 166 biosynthesis pathway, including panB, panG, panC, panD and coaX (Figure 2A-C). This 167 observation is consistent with the current understanding that POA interferes with CoA biosynthesis through disruption of PanD activity<sup>18,19</sup>. Unexpectedly, expression of most other genes of the CoA 168 pathway were not greatly altered by exposure to PZA, POA and MZA with the exception of the final 169 170 gene of the pathway coaE which was down-regulated by 1.5 to 2-fold relative to the no drug control. 171 Due to the central role of CoA in lipid metabolism, we also observed differential expression of 172 numerous genes involved in fatty acid synthesis (FAS) and utilization. The majority of genes 173 associated with the FAS-I and FAS-II systems that are critical for long-chain and mycolic acid 174 biosynthesis, respectively, were overexpressed under all three treatment conditions (Figure 2A-175 C,F). Interestingly, we also observed down regulation of the large gene cluster for synthesis and 176 export of phthiocerol dimycocerosate (PDIM) virulence lipids, as well as differential expression of 177 several other less well characterized genes potentially involved in polyketide and fatty acid 178 synthesis (Figure 2F,G). Beyond lipid synthesis and metabolism, there were also indications of 179 altered respiratory activity and oxidative stress with induction of genes for fumarate reductase, the 180 microaerophilic-type cytochrome bd oxidase and catalase peroxidase katG (Figure 2F).

181 These similarities in responses of *M. tuberculosis* to MZA, PZA and POA exposure 182 suggests mechanistic conservation in their antimicrobial action. Our observation that deletion of 183 *pncA* conferred a slight increase in the MIC for MZA led us to speculate that POA may be released 184 from MZA in a PncA-dependent manner. To understand the role of PncA in response of *M.* 185 *tuberculosis* to MZA, we performed RNA-seq using the H37Rv  $\Delta pncA$  strain exposed to PZA or 186 MZA (Figure 2D,E). As anticipated, the transcriptional profile of H37Rv  $\Delta pncA$  showed no

significant differences between the PZA-treated and untreated control (Figure 2D; Figure S1B). In contrast, the transcriptional profile of H37Rv  $\Delta pncA$  exposed to MZA was distinct from the untreated control, suggesting that MZA exerts additional stress onto bacilli beyond PZA release (Figure 2E; Figure S1B). Despite this distinction, there were only 12 transcripts that met a threshold cutoff for significant differential abundance (2-fold change, adj *p*-value ≤0.05), none of which were immediately informative regarding the PncA-independent action of MZA.

193 Release of PZA and POA from MZA. Based on the distinctions in antitubercular activity of PZA 194 and MZA, and on the profoundly different transcriptional responses of H37Rv and H37Rv  $\Delta pncA$ 195 to MZA exposure, we hypothesized that MZA likely acts through two interacting mechanisms, one 196 that is PncA-dependent and another that is PncA-independent. To parse these mechanisms, we 197 assessed MZA metabolism in strains with and without PncA activity. We began by characterizing 198 the rate of PZA release and conversion to POA. Exponentially growing H37Ra (PncA-proficient) 199 cultures were amended with 250 µM MZA and abundance of MZA, PZA and POA was measured 200 in cell extracts and media over an 8-hour time course via LC-MS. In the first 4 hours, cell-associated 201 MZA rapidly accumulated then steadily declined, while PZA levels rose rapidly from 17 µM to 138 202  $\mu$ M (Figure 3A). With continued incubation, levels of MZA and PZA declined to 34  $\mu$ M and 121  $\mu$ M, 203 respectively. Meanwhile, limited POA accumulation was observed in cell lysates during the time 204 course. The concentration of MZA present in the supernatant steadily declined from 250 to 100 µM 205 over the eight-hour exposure, during which we also observed a steady increase of PZA from 4 to 206 67  $\mu$ M and POA from 1 to 100  $\mu$ M, respectively (Figure 3B).

To determine whether PncA is essential for formation of POA from MZA, a similar experiment was performed with overnight exposure using the vaccine strain *M. bovis* BCG which lacks pyrazinamidase activity due to a loss-of-function mutation in *pncA*. Following overnight exposure, cell-associated MZA was near the limit of detection for both BCG and the control strain H37Ra. Both strains showed accumulation of high levels of PZA, whereas only H37Ra showed an abundant level of POA (Figure 3C). Together, these data demonstrate a rapid conversion of MZA to PZA, followed by PncA-dependent conversion of PZA to POA.

**MZA promotes thiol stress.** To further characterize the PncA-independent mechanism of MZA action, we selected for spontaneously resistant mutants of H37Rv, H37Rv  $\Delta pncA$  and BCG. Selections were performed using solid medium containing a geometric series of concentrations from 0.28 to 0.9 mM MZA. No resistant isolates were recovered from the H37Rv background at any of the tested concentrations. In contrast, H37Rv  $\Delta pncA$  and BCG strains yielded mutants resistant to 0.41 mM MZA at a frequency of 10<sup>-5.9</sup>, and to 0.61 mM MZA at a frequency of 10<sup>-7</sup>. No resistant isolates were observed for either H37Rv  $\Delta pncA$  or BCG at 0.9 mM MZA.

221 After confirming resistance phenotypes, six independently selected isolates of H37Rv 222 ApprcA and four of BCG were analyzed using whole genome sequencing. All ten isolates had single 223 nucleotide mutations within the promoter region of mscR (Figure 3D) encoding mycothiol reductase 224 which is critical for formaldehyde tolerance of mycobacteria<sup>20</sup>. To confirm the protective nature of 225 mutations in the mscR promoter region against aldehyde stress, we exposed wild-type and MZA 226 resistant mutants to methenamine, a bactericidal antibiotic that acts through the spontaneous 227 release of formaldehyde and is commonly used to treat urinary tract infections<sup>21</sup>. While the MIC of 228 methenamine for the wild-type BCG strain was 0.9 mM, the MZA resistant strains showed 229 resistance to 7.1 mM methenamine (Figure 3E,F).

230 To confirm whether the identified promoter mutations conferred MZA resistance through 231 increased MscR expression, we constructed GFP reporter fusions to the mscR promoters from 232 wild-type and C-15T resistant variant in the integrative reporter plasmid pYUB1227. Reporter 233 constructs were integrated at the L5 attachment site in BCG and confirmed strains were grown to 234 exponential phase. After fluorescence was normalized to the OD600 a 1.7 fold increase in 235 fluorescence was observed for the mscR promoter variant relative to the wild-type mscR promoter 236 (Figure 3G) confirming that resistance was due to increased expression of mycothiol reductase. To 237 validate the role of mscR in intrinsic resistance of M. tuberculosis to MZA we constructed a H37Rv 238 ΔmscR strain and assessed MZA susceptibility. This strain showed an eight-fold enhanced level of 239 susceptibility relative to the parental control (Figure 3H).

240 Given the role of MscR in MZA susceptibility and resistance, we reasoned that cysteine 241 supplementation would antagonize MZA action through alleviating thiol stress. To test this 242 hypothesis, BCG wild-type and *mscR* promoter variants were exposed to inhibitory concentrations 243 of 0.9 mM methenamine or MZA in the presence of varying concentrations of cystine as a source 244 of cysteine (Figure 3I,J). For wild-type BCG, 4.1 mM cysteine supplementation was highly 245 antagonistic for both MZA and methenamine. For the mscR promoter mutant, cysteine 246 supplementation was highly antagonistic for MZA at an eight-fold lower concentration than was 247 required for the wild-type strain. Together, these data demonstrate a central role for thiol stress in 248 MZA action and establish increased MscR expression as a mechanism for MZA and methenamine 249 resistance.

250 Genome-wide fitness profiling and chemical biology reveal the unique self-potentiating 251 mechanism of MZA. To better understand pathways that modulate resistance and susceptibility 252 of M. tuberculosis to PZA, POA and MZA, we conducted fitness profiling by using Tn-seq. A 253 saturated library of ~70,000 transposon mutagenized H37Ra was prepared using the himar1 254 mariner transposon<sup>22</sup>. Colonies were harvested and homogenized to form a pooled library and 255 aliquots were precultured in the absence of drug. Exponentially growing libraries were then 256 subcultured and grown for 5 generations in the absence of drug or in the presence of a 257 concentration of PZA (233 μM), POA (105 μM) or MZA (112 μM) that resulted in a two-fold 258 extension of the generation time. These concentrations and number of generations were chosen 259 to allow for the highest resolution of gain- and loss-of-fitness conferred by transposon insertion. 260 Following growth, cells were harvested, genomic DNA was extracted and deep-sequencing of 261 transposon junctions was performed to determine relative abundance of each insertion<sup>8</sup>. Intra-262 population fitness was determined by comparing abundance of individual insertion sites before and 263 after enrichment using a previously established methodology as described in Opijnen et al. 2009<sup>23</sup>. 264 To determine the change in fitness of specific Tn-insertion mutants, the relative fitness of each 265 insertion site was evaluated by calculating the change in fitness with drug compared to the absence 266 of drug. Insertions with relative fitness scores > 1 were considered gain-of-fitness mutants, while 267 relative fitness scores < 1 were considered loss-of-fitness mutants. Consistent with our previous 268 findings on PZA and POA susceptibility<sup>8</sup>, insertions in genes involved in phosphate utilization and 269 transport (pstA1, pstC2, pstS3, phoT), cell wall stress response regulator (sigE), and acyl-CoA 270 synthetase (fadD31) demonstrated substantially greater fitness in the presence of all 3 drugs 271 (Figure 4A-C). We also observed a consistent gain-of-fitness for mutants with insertions in the 272 PDIM locus which correlates with our RNA-seg data and previous indications that PDIM mutants 273 are moderately resistant to PZA<sup>19</sup>. In addition, pncA insertions were found to have a gain-of-fitness 274 in PZA and MZA treated samples, but not POA treated samples, supporting our observation that 275 PncA plays a supporting role in MZA activity (Figure 4A-C). Further, insertions in the mscR locus 276 harbored substantial fitness defects in MZA treated samples relative to PZA and POA treated 277 samples (Figure 4C). This finding is consistent with our observation that MZA susceptibility is 278 modulated by MscR activity in mycobacteria. Interestingly, strains with insertions in sseA, recently 279 shown to be critical for maintaining thiol homeostasis in *M. tuberculosis*<sup>24</sup>, showed a substantial 280 loss-of-fitness in the presence of PZA, POA and MZA, consistent with a direct connection between 281 thiol stress and activity of these drugs. Collectively these data indicate MZA, PZA and POA show 282 conserved interactions within M. tuberculosis metabolism, yet, that MZA action is distinct due to its 283 unique ability to exacerbate thiol stress.

284 Since MZA action shares many features with PZA but does not require exposure to low pH 285 for this aspect of its activity, we next evaluated whether aldehyde can potentiate PZA action in lieu 286 of exposure to low pH. To do so, we conducted a DiaMOND assay<sup>25</sup> testing for synergy between 287 methenamine and PZA at circumneutral pH (Figure 4E). While the individual MIC of PZA at neutral pH surpassed 8 mM and MIC of methenamine was 0.44 mM, the combination of the two produced 288 a Fractional Inhibitory Concentration Index (based on the IC<sub>90</sub> of individual drugs) (FICI) below 0.5 289 290 indicative of synergy<sup>25</sup>, indicating that aldehyde stress sufficiently potentiates PZA activity. To 291 further characterize the synergistic dual action of MZA we assessed whether the common point for synergy between aldehyde stress and PZA action was the depletion of CoA pools<sup>26</sup>. To test this 292 293 hypothesis, we measured intracellular CoA abundance using H37Ra treated with formaldehyde,

PZA, MZA, or a combination of PZA and formaldehyde. Reduction of CoA in PZA treated cells was
reproducible and consistent with previous reports<sup>27</sup> (~24%), but did not meet statistical significance
in our assay. CoA reduction in cells treated with MZA (21%), formaldehyde (34%) and the
combination of formaldehyde and PZA (30%) was highly robust and significant (Figure 4F).
Together, these data demonstrate that the synergistic dual action of MZA is driven through
collateral impairment of CoA metabolism.

300 MZA shows superior potency relative to PZA against intracellular M. tuberculosis. Given the 301 critical role of cell-mediated responses in PZA efficacy and the growing problem of drug resistant 302 *M. tuberculosis*, we assessed the ability of MZA to kill intracellular *M. tuberculosis* in resting or IFN-303 y activated macrophages. RAW 264.7 macrophages were exposed to M. tuberculosis H37Rv or 304 H37Rv *ApricA* at a multiplicity of infection of 1:1 and extracellular bacteria were removed by 305 washing adhered macrophages. For assessing the impact of macrophage activation on MZA efficacy, 5 ng/ml IFN-y was added one day prior to *M. tuberculosis* infection and replenished every 306 307 other day. MZA and PZA were used at a concentration of 0.9 mM and compared to treatment with 308 no drug (DMSO). At the concentration tested, PZA showed limited activity against H37Rv and no 309 activity against H37Rv  $\Delta pncA$  in resting and activated macrophages (Figure 5A-D). In contrast, 310 MZA caused a greater than 3 log<sub>10</sub> decrease in H37Ry viability by day four of treatment in both 311 resting and activated macrophages, and after a week, bacterial loads fell below the detection 312 threshold (Figure 5A.B). Comparable findings were also observed when macrophages were 313 infected with H37Rv  $\Delta pncA$  (Figure 5C,D). Taken together these data demonstrate that the robust 314 bactericidal activity of MZA is both independent of macrophage potentiation and highly effective 315 against intracellular PZA resistant isolates.

316

## 317 Discussion

318

319 In this study we employed a combination of approaches in functional genomics, bacterial 320 physiology and chemical biology to characterize the synergistic dual-action *M. tuberculosis*-specific 321 mechanism of the PZA analog MZA. Our analyses revealed a striking overlap in a central 322 component of the mechanisms of action of PZA and MZA while also resolving the underlying basis 323 for the peculiar difference in their antitubercular activity. These findings led us to identify a self-324 potentiating mode of action specific to MZA that is both unique from and more potently bactericidal 325 than its PZA counterpart. The unique features of MZA, which include rapid killing, high target 326 specificity, low rates of resistance and independence from environmentally-driven potentiation, can 327 be attributed to the drug-specific release of aldehyde resulting in disruption of thiol homeostasis. 328 The rate of aldehyde release, as determined by mass spectrometry, takes place over the course of 329 hours, which allows for elimination of bacilli from infected macrophages within a matter of days. 330 The superior feature of MZA is orchestrated by the synergistic actions of its two components, 331 aldehyde release followed by PZA collateral susceptibility (Figure 6). The rapid engagement of aldehyde elicits bactericidal activity against both actively growing and dormant cells alike while 332 simultaneously sensitizing any survivors to the activity of PZA. This potentiation event ensues via 333 334 depletion of biologically active thiols, resulting in bacterial growth arrest and increased demand for 335 CoA with consequent disruption of fatty acid biosynthesis, a pathway known to be perturbed by 336 PZA<sup>15,28</sup>. The mechanism proposed here is further confirmed by synergistic activity of methenamine 337 with PZA and the antagonistic activity of cysteine against MZA.

338 The robust activity of MZA against non-growing *M. tuberculosis* suggests that entry into the 339 cytoplasm is likely a passive event. Hence, MZA is effective at killing *M. tuberculosis* independent 340 of metabolic activity or growth state of the cells. Passive entry of MZA into the cytoplasm also 341 supports a low likelihood for emergence of drug resistant strains via mutations that alter drug 342 transport. While low-level resistance to aldehyde release can be achieved via activation mutations 343 in the mscR promoter, achieving full resistance to MZA will likely be a challenge due to the 344 promiscuous nature of aldehyde-mediated damage. Moreover, we find that PZA resistant strains 345 of *M. tuberculosis* and *M. bovis* BCG with mscR promoter-up mutations still show measurable 346 susceptibility to MZA, suggesting either that the rate of internal aldehyde release is greater than

the MscR turnover rate or that other aldehydes that cannot be effectively neutralized by MscR are
released by MZA. In either case, the bacilli will be faced with a greater adaptive burden to develop
full resistance to MZA (Figure 6).

350 Given the pivotal role of PZA in TB treatment regimens and the limited activity of this drug 351 in models of compromised immunity<sup>11</sup>, it will be critical to resolve the importance of aldehyde-driven 352 thiol stress in its antitubercular efficacy. In the present study, aldehyde release from MZA was able 353 to reduce bacterial loads in resting macrophages to levels below the limit of detection in under a 354 week. Based on loss of PZA efficacy in the athymic mouse model<sup>11</sup>, it has been speculated that 355 antitubercular activity of PZA may be diminished in the context of T cell deficiency or dysfunction 356 due to insufficient macrophage activation, resulting in reduced treatment efficacy and higher rates 357 of drug resistance<sup>9</sup>. Circumnavigating the need for host-driven potentiation of PZA may prove to be 358 a valuable approach in the treatment of *M. tuberculosis* in the context of compromised immunity.

359 Interestingly, it has recently been recognized that production of host-derived aldehydes are 360 modulated via IFN-y activation and play a role as antimicrobial effectors against intracellular pathogens, such as M. tuberculosis and Francisella tularensis 29,30. Whether host-derived 361 362 aldehydes participate in the bactericidal activity of PZA in vivo has yet to be determined, but, may represent an opportunity for host-directed therapy for bolstering the action of this critical 363 antitubercular agent. Further, microbe-derived aldehyde production may also represent an 364 365 opportunity that can be explored for novel therapeutic discovery. For example, selective disruption 366 of cytokinin metabolism has been shown to result in accumulation of toxic aldehyde species in M. 367 tuberculosis and further sensitize the bacilli to other host stressors<sup>31</sup>. Further, as noted by Darwin 368 and Stanley<sup>29</sup>, mycobacterial glycolysis may represent an exploitable target for driving toxic aldehyde accumulation in *M. tuberculosis*<sup>32</sup>. Along these lines, it is curious to note that strains 369 bearing loss-of-function mutations in *glpK* (encoding glycerol kinase) show a selective advantage 370 both in humans and in mice treated with antitubercular agents<sup>33,34</sup>. Since disruption of glycerol 371 phosphate synthesis would reduce the level of endogenous metabolic aldehydes through limitation 372 373 of glyceraldehyde phosphate and dihydroxyacetone phosphate synthesis, glycerol kinase likely 374 represents a key mediator in endogenous mycobacterial aldehyde production<sup>29</sup>. In strong support 375 of this concept, Dick and colleagues previously determined that mutations in *glpK* are associated 376 with PZA resistance in vitro<sup>19</sup>. Our findings, coupled with the emerging role of host- and microbe-377 derived aldehydes in compromising the fitness of *M. tuberculosis*, advocate for further evaluation 378 of the role of aldehyde-mediated potentiation of antimicrobial agents such as PZA for improved 379 therapeutic approaches.

380

# 381 Materials and Methods382

383 Standard growth conditions. Middlebrook 7H9 broth (Difco) supplemented with 10% Middlebrook 384 OADC (Difco), 0.02% (V/V) glycerol and 0.05% (V/V) tyloxapol were used for all experiments 385 involving liquid *M. tuberculosis* and *M. bovis* cultures. Depending on experimental requirements, 386 the media were adjusted to pH 7.2 or pH 5.8. Cultures were incubated shaking at 100 RPM 37°C. All solid media used in experiments involving M. tuberculosis and M. bovis were based on 387 388 Middlebrook 7H10 (Difco) supplemented with 10% Middlebrook OADC, 0.02% (V/V) glycerol. For 389 *M. smegmatis* the same media were used with the exclusion of OADC and replacing glycerol with 390 0.2% (W/V) dextrose. ESKAPE pathogens and *E. coli* (for plasmid amplification) were propagated 391 in Luria-Bertani broth (LB, Difco) under standard growth conditions (37°C shaking at 250 RPM).

392 Strain construction. The knockout strains (H37Rv  $\Delta pncA$  and H37Rv  $\Delta mscR$ ) and were 393 constructed using the ORBIT system<sup>35</sup>. H37Rv was electroporated with pKM461. After selection on 394 50 µg/mL kanamycin, H37Rv pKM461 was induced with anhydrotetracycline and electroporated 395 with pKM464 the position specific oligonucleotide 5'and pncA TACCTCGGCGCCACGGCGGCGGACCCGGCCCGCGCCCGGTGGCTCCT 396 397 GCACTTCGGCATGGTGGGCCGCAGGTTTGTACCGTACACCACTGAGACCGCGGTGGTTGA

398 CCAGACAAACCCCTCGACTCGCTTCCGACAGCACCTCGAAGACCGCTTCGGGTGCGTGAG

399 CACGCTGGGCGGTTCGCAGTG-3' or mscR 5'-GCGCGCATGGTCAGCGACGCTACACCGT 400 AGGTTGGACACCATGAGTCAGACGGTGCGCGGTGTGATCGCAGGTTTGTCTGGTCAACCAC 401 CGCGGTCTCAGTGGTGTACGGTACAAACCAAGGTATTGCGTTCGGTGGTGATGTTGTGATG 402 GCCGCCATCGAGCGCGTCATCACCCACGGCACCTTCGA-3'. Transformed bacteria were then 403 plated on supplemented 7H10 medium containing 50 µg/mL hygromycin B and 10% sucrose to 404 select for recombinants and counter select against pKM461. Individual colonies were then re-405 streaked, confirmed via PCR, and full genome sequenced to validate desired modification and no 406 additional mutations.

407 Fluorescent reporter mscR strains were constructed using a modified integrative reporter 408 plasmid pYUB1227. Restriction enzymes Pvul and Mfel (NEB) were used to prepare the backbone 409 and the prompter inserts, followed by a T4 ligation. Primers used to validate promoter changes were mscRF 5'-TTGCAACGCATCCCTGATCT-3' and mscRR 5'-AGGGCAGATTGTGTGGACAG-410 411 3'. Additional validation was performed via Nanopore sequencing. Plasmid amplification was 412 performed in *E. coli* DH5α. Plasmids were purified using a QIAprep Spin Miniprep Kit (cat.# 27104). 413 BCG Pasteur 1173P2 was then transformed using previously prepared plasmids via standard 414 electroporation protocol. Successful transformants were selected for on standard media amended 415 with 50 µg/mL hygromycin B and validated via genomic DNA extraction and PCR using validation 416 primers described above.

417 Determination of minimum inhibitory concentrations. Bacteria were grown in 30 mL PETG 418 square media bottles (Nalgene) containing 5 mL of 7H9 media under standard growth conditions 419 (37°C shaking at 100 RPM) to OD<sub>600</sub> of 0.15-0.25. Bacteria were then diluted to OD<sub>600</sub> of 0.01 and 420 allowed to incubate for 2 weeks in presence of the compound(s) being tested. Concentrations of 421 compounds were determined as a function of a geometric series (0.05, 0.1, 0.2, 0.4, 0.81, 1.62, 422 3.25 and 6.5 mM) for PZA, (0.06, 0.11, 0.22, 0.45 and 0.9 mM) for MZA and (0.06, 0.11, 0.22, 0.44, 423 0.88, 1.75 and 3.5 mM) for methenamine. MIC values were then calculated by plotting inhibition 424 curve and calculating based on the appropriate slope intercept.

Antagonism assays. Bacteria were prepared in the same way as for MIC assays and were then inoculated into 7H9 media containing either 0.9mM methenamine, 0.9mM MZA or DMSO (vehicle control) and antagonist (pantothenate or cystine) at indicated concentrations. Pantothenate was used at a single concentration of 230  $\mu$ M, cystine concentrations ranged from 15  $\mu$ g/mL to 500  $\mu$ g/mL in 2-fold increments. The media pH was adjusted to 7.2 and cells were incubated in 96-well plated for 8 days at 37°C with no shaking. Measurements were performed using a BioTek Synergy H1 plate reader (Agilent). All experiments were performed in triplicate.

Bacterial kill curves. To assess the bactericidal activity of MZA and methenamine, against
exponentially growing cells, bacteria were grown in 30 mL bottles containing 5 mL of 7H9 media
under standard growth conditions (37°C shaking at 100 RPM) to OD<sub>600</sub> of 0.15-0.25. Bacteria were
then diluted to OD<sub>600</sub> of 0.03 (approximately 5X10<sup>6</sup> bacilli) and plated on 7H10 media to determine
the starting population of bacteria. After addition of (0, 0.28, 0.41, 0.61 or 0.9mM of MZA) and (0.06,
0.11, 0.22, 0.44, 0.88, 1.75 and 3.5 mM) µg/mL Methenamine, CFU counts were determined at 0,
2-, 4-, 7- and 14-day time intervals.

439 To assess bactericidal activity of PZA and MZA against nutrient limited *M. tuberculosis*, 440 bacteria were grown in 30 mL bottles containing 5 mL of 7H9 media under standard growth 441 conditions (37° C shaking at 100 RPM) to OD<sub>600</sub> of 0.15-0.25. Bacteria were then washed three 442 times with 1xPBS saline to remove any residual media and resuspended in 1xPBS saline amended 443 with 0.05% tyloxapol. Prior to addition of the drugs CFU/mL were determined by serial dilution 444 plating on 7H10 media, 900 µM PZA, MZA or DMSO were then added to the buffer and CFU 445 measurements were taken at predetermined intervals 0, 2-, 3-, 4-, 7- and every 7 days afterwards 446 until 62-day timepoint was reached. For MBC measurements under starvation conditions with 447 multiple treatments, bacteria were prepared in the same way as for a single treatment. However, 448 at every timepoint culture was washed three times to remove residual antibiotics and split into two 449 cultures. One of the two cultures were treated with DMSO while the other was amended with fresh 450 MZA for a final concentration of 900 µM.

451 RNA-seq. For bulk RNA-seq library preparation and sequencing, exponentially growing H37Rv and 452 H37Rv  $\Delta pncA$  were grown to the OD<sub>600</sub> of 0.15 and inoculated with 200  $\mu$ M MZA, PZA or equivalent 453 volume of DMSO (as a vehicle control). Cells were then incubated under standard condition for 24 454 hours prior to harvesting. Following incubation, TRIzol RNA purification was performed. Cells were 455 chilled, pelleted and resuspended in a 500 µL of TriReagent (Invitrogen) with 1% polyacryl carrier 456 (Molecular Research Center) and lysed using 250 µL of 0.1 mm zirconia beads (BioSpec). Samples 457 were then centrifuged and supernatant was transferred to a clean 2 mL microcentrifuge tube and 458 combined with 50 µL of 5-bromo-3-chloro-propane. Following a 10-minute incubation samples were 459 centrifuged aqueous phase was removed and transferred to a clean 1.5 mL Eppendorf tube where 460 samples were treated with 250 µL of isopropanol. Following a 10-minute incubation samples were 461 centrifuged again, isopropanol was removed and RNA coating the tube was washed with 300 uL 462 of 75% EtOH. After the final wash, RNA was dried and eventually resuspended in 50 uL of DEPCtreated water (Invitrogen). A DNA-free (<sup>™</sup>) DNAse treatment (Invitrogen) was used to remove 463 464 residual DNA from the samples prior to sequencing. Samples were treated with Ribo-Zero Plus kit 465 and Libraries were prepared using an Illumina Stranded Total RNA Prep Ligation. Sequencing was performed at the University of Minnesota Genomic Center using a 50 paired-end NovaSeq S-prime 466 467 platform.

468 For bulk RNA-seq analysis, the pipeline used for preprocessing raw fastq files can be found 469 at https://github.com/MDHowe4/RNAseg-Pipeline. Quality control of RNA sequencing read quality 470 was assessed with FastQC. Read length thresholding and t-overhang trimming of forward and 471 reverse reads was completed with Cutadapt with minimum read length cutoff of 30 bp. Reads were 472 mapped to the M. tuberculosis H37Rv reference genome (NC 000962.3) using the STAR aligner 473 without spliced alignment detection (--alignIntronMax 1)<sup>36</sup>. Total reads per gene were counted using featureCounts<sup>37</sup>. Genes with <10 reads mapped occurrences across wild-type and mutant 474 475 experiments were not included in further analysis. Differentially expressed genes (DEGS) were 476 found using the negative binomial generalized linear model of DESeg2<sup>38</sup>. Genes were considered 477 differentially expressed if they displayed a log<sub>2</sub> fold-change  $\geq$  1.5 or  $\leq$  -1.5 and adjusted *P*-value  $\leq$ 478 1x10<sup>-6</sup>. Volcano plots were generated using the EnhancedVolcano package.

479 Mass spectrometry. For mass spectrometry analyte preparation, for each sample, a total of 40 480 mL of bacteria were grown under standard growth conditions at pH 5.8. Upon reaching OD<sub>600</sub> of 481 0.4, bacteria were inoculated with either 250 µM MZA or PZA. After three days of treatment the 482 samples were pelleted. Cell extract samples were prepared for mass spectrometry analysis by 483 resuspending pellets in 1 mL of extraction buffer (40% methanol, 40% acetonitrile and 20% water) 484 and bead beaten using 250 µL of 0.1 mm zirconia beads (BioSpec). Samples were then centrifuged 485 to remove the solids. Liquid fractions were then filtered using 22 µm filter. A portion of the filtrate 486 was then used to determine protein content using a Pierce<sup>™</sup> BCA Protein Assay Kit (Thermo Scientific). The remaining samples were further purified using a 3 kDa column (PALL Life 487 488 Sciences). Supernatant sample preparation was performed in the identical way as cell extract, 489 except for the omission of the first bead beating step. Bacterial cell extracts were diluted 10-fold 490 (10 µL in 100 µL) in DI water. 100 µL of diluted solution was treated with 100 µL methanol: water 491 solution containing para-aminosalicylic acid (PAS, internal standard), mixed well by vortexing, 492 centrifuged (2 min,  $15000 \times q$ ), and supernatant ( $150 \mu L$ ) was analyzed by LC-MS/MS.

493 For LC-MS/MS analysis, reverse-phase LC was performed on a Zorbax Eclipse XDB- C8 494 column (150 mm, 4.6 mm, 5 µm; Agilent) using Shimadzu UFLC XR instrument. The elution 495 gradient was carried out with binary solvent system consisting of 0.1% formic acid in H<sub>2</sub>O (solvent 496 A) and 0.1% formic acid in MeCN (solvent B). A linear gradient profile with the following proportions 497 (v/v) of solvent B was applied (t (min), %B): (0, 5), (1, 5), (4, 95), (6, 95), (7, 5), (8, 5) with 2 min for 498 re-equilibration to provide a total run time of 10 min. The flow rate was 0.5 mL/min and the column 499 oven was maintained at 40°C. The injection volume was 10 µL. The retention times for MZA, PZA, 500 POA and PAS (internal standard) were 3.5, 2.6, 3.8 and 6.8 min, respectively, MS/MS analysis 501 were carried out using triple quadrupole/linear ion trap instrument (AB SCIEX QTRAP 5500). MZA, 502 PZA and POA peak areas were calculated using MultiQuant, version 2.0.2. MZA, PZA and POA 503 areas were normalized to p-amino salicylate areas and the MZA, PZA and POA concentration in 504 samples were determined using the standard curves of MZA, PZA and POA. For the preparation

of MZA, PZA and POA standard curve, average area of standard 0 was subtracted from MZA, PZA
 and POA of all higher standards prior to area normalization. Sample concentrations were back
 calculated to account for dilutions. Each sample was performed in triplicate.

Transposon mutagenesis and transposon sequencing. M. tuberculosis H37Ra was 508 509 mutagenized with the mariner himar1 transposon using the phAE180 temperature-sensitive 510 mycobacteriophage<sup>39</sup>. Approximately 70,000 independent mutants were selected on 7H10 media amended with 50 µg/mL kanamycin. Transposon mutants were pooled, homogenized, then 511 aliquoted into 25% (V/V) glycerol stocks and stored at -80°C. Growth curves to establish 50% 512 inhibition of growth rate using a geometric series of drug concentrations were carried out using 7H9 513 514 medium at a pH of 6.0. Cultures were seeded at OD<sub>600</sub> of 0.01 in 80 mL of medium using transposon 515 mutants pre-cultured in a shaking incubator for 48 hours at 37°C. Enrichment was carried out in 516 the presence or absence of drugs for 5 generations as determined by  $OD_{600}$ . After 5 generations, 517 cultures were flash frozen in liquid nitrogen for genomic DNA (gDNA) extraction. gDNA extraction 518 was carried out using a previously established protocol<sup>40</sup>. Library prep and sequencing was 519 performed at the University of Minnesota Genomics Center with read mapping performed using a 520 previously fragmentation established protocol<sup>8</sup>. DNA and Illumina P7 521 (CAAGCAGAAGACGGCATACGAGAT) ligation was performed. Amplification of transposon site 522 junction was carried out using a P7 (CAAGCAGAAGACGGCATACGAGAT) and mariner-specific 523 Mariner 1R TnSeg noMm primer

524 (TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGCCGGGGACTTATCAGCCAACC).

Amplification of himar1-enriched samples was performed using a P5 indexing primer 525 526 (AATGATACGGCGACCACCGAGATCTACAC[i5]TCGTCGGCAGCGTC: [i5] barcode sequence) and a P7 primer HotStarTag master mix kit (Qiagen) after a 1:50 dilution. Sequencing was 527 528 performed on an Illumina NovaSeq. The 5' ends of reads were trimmed to remove adapter and 529 transposon sequence using Cutadapt to leave the TA-gDNA junction intact<sup>41</sup>. Trimmed reads with less than 18 bp were discarded and reads were mapped to the M. tuberculosis H37Ra genome 530 531 (GenBank Accession: CP000611) with 1 bp permitted mismatch using Bowtie 2<sup>42</sup> (PMID: 532 22388286). Mapped reads were printed in SAM format and counted per TA dinucleotide sight using 533 SAMreader\_Ta script<sup>43</sup>. Scripts used for read processing may be found on GitHub 534 (https://github.com/MDHowe4/Himar1-TnSeq-Pipeline).

The relative abundance of insertions at TA sites was calculated for each sample. TA sites lacking mapped insertions at the start of enrichment were omitted. TA sites without mapped insertions at the end of enrichment that were present at the start of the enrichment were set to the limit of detection without contributing to the total number of insertions. Relative fitness of sitespecific mutants were calculated with a previously characterized function using the population expansion factor, as well as the relative number of insertions at the beginning and end of enrichment<sup>23</sup>.

542 Isolation of spontaneous mutants. To isolate spontaneously MZA resistant mutants, H37Rv 543  $\Delta pncA$  and BCG cells were grown up to mid-log phase for selection. 1 mL of culture diluted to OD<sub>600</sub> 544 of 0.3 was plated onto standard 7H10 agar supplemented with 409  $\mu$ M MZA for H37Rv  $\Delta pncA$ , and 545 607 μM for BCG. Plates were then incubated at 37°C in stationary incubator for 14-21 days. 546 Resistant mutants were then picked and streaked again for isogenic backgrounds, then grown 547 again for 14-21 days at 37°C in a stationary incubator. Isogenic colonies were picked, then grown 548 in standard 7H9 broth for 14 days followed by genomic DNA extraction using a the previously 549 described DNA extraction methodology. Illumina sequencing was performed at SeqCenter. Read 550 alignment and variant calling to the H37Rv reference genome (NC 000962.3) and BCG reference 551 genome (BCG Pasteur 1173P2) was performed using Breseq version 0.36.0.

**Coenzyme A detection assays.** Bacteria were prepared for extraction in the same way as described in mass-spectrometry methods. 80 mL per treatment sample of H37Ra was grown to an OD<sub>600</sub> of 0.13 in 7H9 media in the shaking incubator at 100 RPM and 37°C. All of the samples were then treated with 200  $\mu$ M of either PZA, MZA, formaldehyde or formaldehyde + PZA. After an eighthour exposure cells were harvested and resuspended in 400  $\mu$ L of ice cold 1xPBS and lysed using

557 zirconia beads. Samples were then centrifuged and filtered using 0.22-µm filter. 50 µl aliquots were 558 then taken to perform a BCA assay (Pierce™ BCA Protein Assay Kit, ThermoScientific). The 559 remaining extracts were then filtered one more time using a using a 3 kDa column (PALL Life 560 Sciences). Twice filtered lysate was then used to perform fluorometric assay using the Free 561 Coenzyme A Assay kit (Sigma-Aldrich). The readout was then normalized to the protein 562 abundance. All assays were performed in triplicate.

563 Macrophage Infections. RAW 264.7 macrophages were prepared by seeding density of 1.0-2.0 x 564 10<sup>5</sup> cells per well in DMEM/F-12 medium containing 10% FBS without antibiotics (DMEM/F-12 565 complete) in 12 well (1 mL/well) plates. In order to allow for cells to adhere, macrophages were 566 incubated overnight in a humidified 5% CO<sub>2</sub> chamber. The following day cells were rinsed with 567 Hank's buffer three times and replenished with fresh DMEM/F-12 media. A 5 ng/mL of IFN-y was 568 added to the well containing macrophages meant to be activated. Following a 14-16h incubation, 569 macrophages were washed again in Hank's buffer and replenished with fresh, drug and interferon 570 free DMEM media. Upon transfer of macrophages into BSL-3 space, media was removed, cells 571 were washed with Hank's buffer three times and fresh DMEM media containing M. tuberculosis 572 H37Rv. An MOI of 1:1 was used for the initial infection. To prepare bacteria, H37Rv was grown 573 under standard growth conditions. Upon reaching OD<sub>600</sub> of 0.25, culture was washed three times. 574 resuspended, and then diluted in DMEM media to the 1.0-2.0 x 10<sup>5</sup> CFU/mL. Following a 2h 575 infection, infected macrophages were washed three times with Hank's buffer and resuspended in 576 complete DMEM media amended with 900 µM MZA, PZA or DMSO (as a vehicle control) as well 577 as 5ng/ml of IFN-y for activated macrophages. Media was changed daily for the duration of the 578 experiment with re-induction via IFN-y taking place on every other day. During every media change, 579 cells were washed three times with Hank's buffer and adherence and appearance of macrophages 580 was monitored via a stereoscope. On plating days, cells were washed as described above, and 581 macrophages were lysed via TritonX-100. The lysate was then serially diluted in 0.05% Tyloxapol 582 1xPBS pH 7.2 buffer using 10-fold dilution in a 96well plate format and plated on 7H10 media. 583 Plates were incubated for two weeks and colonies counted for CFU determination. All experiments 584 were performed in triplicate.

585 **MZA synthesis and NMR analysis**. Formaldehyde 37% in water (89 µL, 1.9 mmol) was prepared 586 with PZA (200 mg, 1.6 mmol) and morpholine (1.1 mL, 12.9 mmol) in 100% ethanol solvent with 587 one drop of concentrated HCI. The reaction was refluxed for 6 hours and neutralized with one drop 588 of 1 N NaOH. The crude mixture was evaporated and separated via silica gel flash chromatography 589 with methanol/dichloromethane solvent system. The pure compound (>95%) was a white solid. The 590 calculated yield was 63.7%. Observed (M+H) + of 223.0 m/z with mass spectrometry analysis.

591 <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>-d, ppm):  $\delta$  = 9.35 (s, 1), 8.71 (d, 1), 8.48 (d,1), 8.12 (s, 1), 4.29 592 (d, 2), 3.65 (t, 4), and 2.58 (t, 4). <sup>13</sup>C NMR (100 MHz, DMSO-d<sub>6</sub> ppm):  $\delta$  = 164.2, 148.0, 145.0, 593 144.1, 143.7, 66.4, 60.8, and 50.5. Purity: >99%.

Quantification and statistical analyses. Where applicable, 95% confidence intervals were used
 to determine statistical significance. Numbers of biological replicates and statistical tests that were
 employed are described in respective figure legends.

597 **Materials availability.** Materials will be made available upon reasonable request and may require 598 completion of a Material Transfer Agreement and payment for related expenses.

599 **Data and code availability.** All data are available in the main manuscript or supplemental 600 information. Primary sequencing data from RNA-seq, Tn-seq, and whole-genome sequencing 601 studies are publicly available through the National Center for Biotechnology Information via SRA 602 link <u>https://www.ncbi.nlm.nih.gov/sra/PRJNA1104292</u>.

603 Code for bulk RNA-seq analysis pipeline used for preprocessing raw fastq files can be 604 found on GitHub at <u>https://github.com/MDHowe4/RNAseq-Pipeline</u>. Code used for read processing 605 can be found on GitHub https://github.com/MDHowe4/Himar1-TnSeq-Pipeline.

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## 608

## 609 Acknowledgments

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- 617 for determination of MZA purity.
- 618

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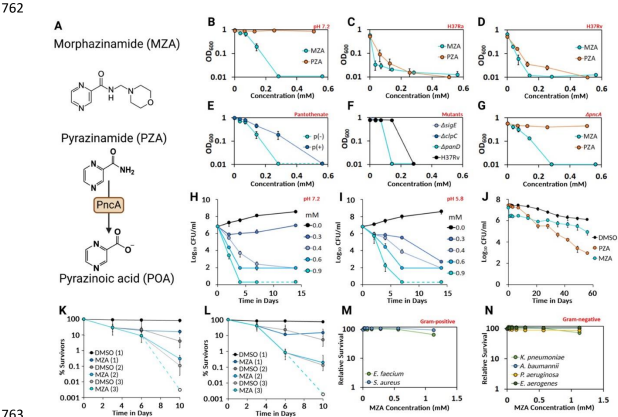
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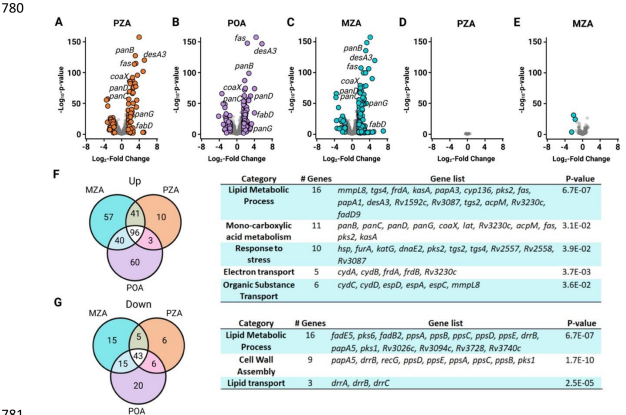


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764 Figure 1. Impact of MZA on PZA susceptible and resistant M. tuberculosis. A, Structure of 765 MZA and activation of PZA to POA by PncA<sub>Mtb</sub>. Dose response curves comparing MZA and PZA susceptibility for *M. tuberculosis* strains **B.** H37Ra at pH 7.2 and **C.** H37Ra at pH5.8 and **D.** H37Rv 766 at pH 5.8. E, Assessment of pantothenate antagonism of MZA action on H37Ra at pH 7.2. 767 Determination of MZA susceptibility of PZA resistant strains **F**, H37Rv  $\Delta sigE$ , H37Rv  $P_{clpC1}$ ::himar1, 768 769 H37Rv panD::himar1 and G, H37Rv ApncA compared to parental H37Rv. Dose dependent kill curves of H37Rv exposed to MZA at H, pH of 7.2 and I, pH 5.8 over a 14-day period. J, Kill curves 770 771 performed with H37Rv under starvation conditions following a one-time treatment with PZA, MZA 772 or DMSO over a 60-day period at pH of 7.2. Kill curves performed with K, H37Rv and L, H37Rv ApprcA (PZA resistant) incubated at neutral pH under starvation conditions with MZA replenished 773 every 3 days. Evaluation of MZA activity against M, Gram-positive and N, Gram-negative ESKAPE 774 775 pathogens. All assays were performed in biological triplicate with mean displayed and error-bars 776 indicative of standard deviation.

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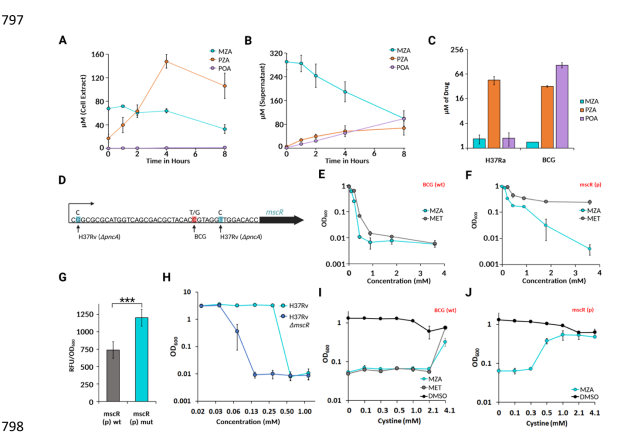
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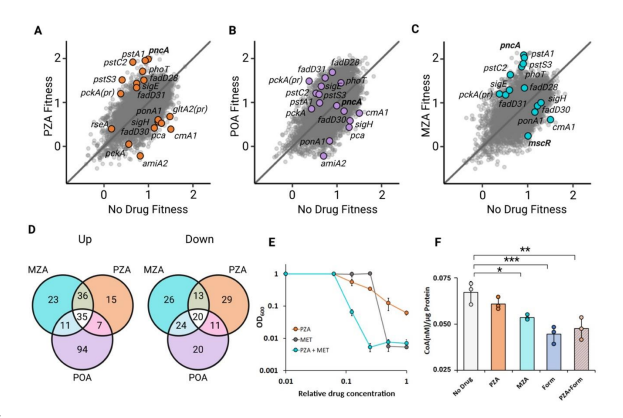
782 Figure 2. Transcriptional profiling of *M. tuberculosis* exposed to MZA, PZA and POA. Volcano 783 plots showing significantly differentially expressed genes in presence of A, PZA, B, POA and C, 784 MZA. Cells were treated with 200 µM MZA, POA or PZA for 24h prior to RNA purification and 785 sequencing. The MZA, POA and PZA transcriptional profiles share many key features, including upregulation in coenzyme A and lipid metabolism pathways. Transcriptional changes associated 786 787 with exposure of a H37Rv ΔpncA to **D**, PZA and **E**, MZA. Cells were treated with 200 μM MZA or 788 PZA for 24h prior to RNA purification and sequencing. F, Venn diagram showing upregulated genes 789 in MZA, POA and PZA treated cultures, with the corresponding GO term analysis of common genes 790 to the right. G, Venn diagram showing downregulated genes in MZA, POA and PZA treated 791 cultures, with the corresponding GO term analysis of common genes to the right. RNA-seq was 792 performed in biological triplicate.

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799 Figure 3. Metabolism of MZA results in release of PZA and drives thiol stress. Time course 800 monitoring A, intracellular and B, extracellular abundance of MZA to PZA and POA from H37Ra 801 exposed to 250 µM MZA. C, Intracellular abundance of MZA, PZA and POA following a 24-hr treatment of H37Ra (PZA susceptible) and *M. bovis* BCG (PZA resistant) with MZA. D, mscR 802 803 promoter region highlighting MZA resistance mutations in H37Rv  $\Delta pncA$  (blue) and BCG (red) 804 backgrounds, respectively. Inhibition curve of an E, wild-type BCG and a F, mscR promoter mutant 805 exposed to MZA and methenamine (MET). G, Comparison of GFP expression between wild-type mscR promoter and mscR promoter variant GFP reporter constructs in BCG. All samples were 806 807 normalized to the OD<sub>600</sub>, significance was determined using a 2-tailed Student's t-test using n=6. 808 H, Inhibition curve comparing susceptibility of H37Rv and H37Rv *AmscR* to MZA. I, BCG wild-type 809 and J, BCG mscR promoter mutant exposed to 0.9 mM MZA or MET in the absence or presence 810 of cystine. All assays were performed in biological triplicate unless indicated otherwise with mean 811 displayed and error-bars indicative of standard deviation.

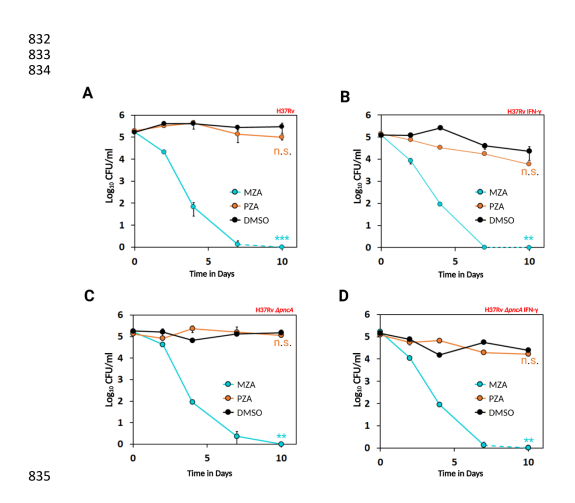
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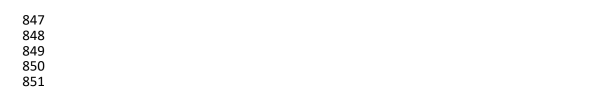
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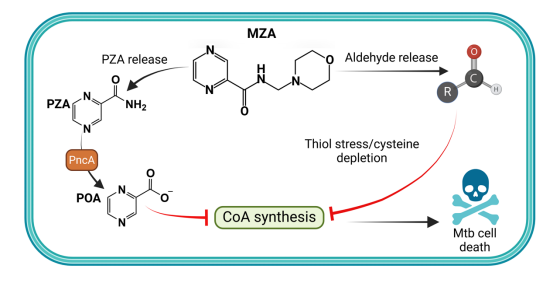
Figure 4. Genome-wide fitness profiling of *M. tuberculosis* transposon insertion mutants 816 and chemical biology reveal the unique self-potentiating mechanism of MZA. Saturated 817 library of Tn-insertion mutants treated with A, 233 µM PZA B, 105 µM POA and C, MZA 112 µM 818 819 MZA at pH 6 to achieve a 50% reduction in growth rate. Each point corresponds to a specific TA 820 site harboring a transposon insertion within the genome. The plot shows every transposon insertion 821 identified. Highly depleted or enriched TA sites are highlighted with genes of interest noted. D, 822 Venn diagram showing genes enriched for (left) and depleted (right) shared by all three treatments. 823 E, DiaMOND assay showing exposure of *M. tuberculosis* H37Ra to a geometric series of PZA and 824 methenamine (MET) alone and in combination. F, Abundance of CoA from *M. tuberculosis* H37Ra 825 treated with DMSO (No Drug), MZA, PZA, formaldehyde (Form) and a combination of PZA and formaldehyde (PZA+form). Values were normalized to total protein. Significance was determined 826 827 using a one-tailed Dunnett's test with Bonferroni correction. All assays were performed in biological 828 triplicate with mean displayed and error-bars indicative of standard deviation.

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836 Figure 5. MZA efficacy against PZA susceptible and PZA resistant *M. tuberculosis* strains in resting and activated macrophages. A, CFU comparison between RAW 264.7 macrophages 837 infected with H37Rv and treated with either MZA, PZA or DMSO. B, H37Rv infected macrophages 838 839 were activated using IFN-y. C, Same experimental design as in A but using the PZA resistant 840 H37Rv ApricA strain to test efficacy of MZA against PZA resistant strains. D, RAW 264.7 macrophages activated with IFN-y and infected with PZA resistant H37Rv ApricA. Cells were 841 842 treated with 0.9 mM MZA or PZA with daily media exchange. All assays were performed in biological triplicate with mean displayed and error-bars indicative of standard deviation. 843 844 Significance was determined via Kruskal-Wallis test combined with post hoc Dunn test with p<0.01 845 (\*\*), *p*<0.001 (\*\*\*).

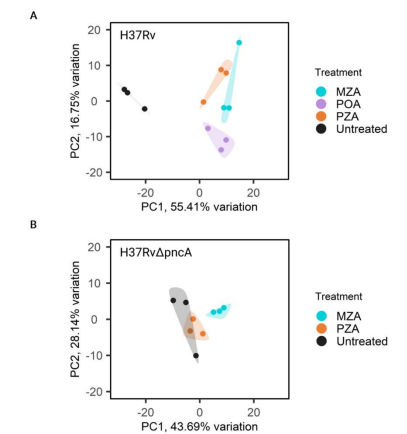




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Figure 6. Thiol stress mediated potentiation of PZA action. Exposure to low pH drives thiol
stress in *M. tuberculosis* resulting in potentiation of PZA activity. MZA releases aldehyde and PZA
in *M. tuberculosis* regardless of pH. PZA is converted to POA which interferes with CoA synthesis.
Aldehyde release drives thiol stress resulting in synergistic disruption of CoA metabolism with PZA
resulting in rapid bacterial cell death.

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Figure S1. PCA for RNA-seq treatment groups. A, PCA plots from data from wildtype M. tuberculosis H37Rv cells treated with DMSO, PZA, POA or MZA. B, PCA plots from the M. 863 tuberculosis H37Rv ApricA cells treated with DMSO, PZA or MZA. Each point represents an 864

- 865 individual biological replicate.
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## Table S1. PZA and MZA MIC values under potentiating and antagonizing conditions.

Strain		MIC in 7H9 media	
	Condition	PZA (mM)	MZA (mM)
H37Ra	pH 5.8	0.5	0.14
H37Ra	pH 5.8, pantothenate	>4.5	0.28
H37Ra	pH 7.2	>4.5	0.28
H37Ra	pH 7.2, pantothenate	>4.5	0.56
H37Rv	pH 5.8	0.5	0.14
H37Rv	pH 7.2	>4.5	0.28
H37Rv ΔpncA	pH 5.8	>4.5	0.28
H37Rv ∆ <i>sigE</i>	pH 7.2	>4.5	0.14
H37Rv P <sub>clpC1</sub> ::himar1	pH 7.2	>4.5	0.14

Experiments were performed using *M. tuberculosis* strains H37Ra and H37Rv and their derivatives. For potentiation of PZA action, pH of 5.8 was used. For antagonism of PZA action, pantothenate was used at a concentration of 0.23 mM. Strains H37Rv  $\Delta pncA$ ,  $\Delta sigE$  and  $P_{clpC1}$ ::*himar1* were previously described<sup>8</sup> for their differing levels of PZA resistance. All assays were performed in biological triplicate with mean displayed.