Accepted Manuscript

Clofazimine improves clinical outcomes in multidrug-resistant tuberculosis: a randomized controlled trial

Hongfei Duan, Xiaohong Chen, Zhihui Li, Yu Pang, Wei Jing, Ping Liu, Tingting Wu, Cui Cai, Junwei Shi, Zhihua Qin, Hongyun Yin, Chao Qiu, Chunxiang Li, Ying Xia, Wei Chen, Zhizhong Ye, Zhiyue Li, Gang Chen, Sheng Wang, Yufeng Liu, Lixiang Chu, Min Zhu, Tao Xu, Qingfeng Wang, Jing Wang, Yadong Du, Jun Wang, Naihui Chu, Shaofa Xu

PII: S1198-743X(18)30530-5

DOI: 10.1016/j.cmi.2018.07.012

Reference: CMI 1382

To appear in: Clinical Microbiology and Infection

Received Date: 29 January 2018

Revised Date: 5 July 2018

Accepted Date: 10 July 2018

Please cite this article as: Duan H, Chen X, Li Z, Pang Y, Jing W, Liu P, Wu T, Cai C, Shi J, Qin Z, Yin H, Qiu C, Li C, Xia Y, Chen W, Ye Z, Li Z, Chen G, Wang S, Liu Y, Chu L, Zhu M, Xu T, Wang Q, Wang J, Du Y, Wang J, Chu N, Xu S, Clofazimine improves clinical outcomes in multidrug-resistant tuberculosis: a randomized controlled trial, *Clinical Microbiology and Infection* (2018), doi: 10.1016/j.cmi.2018.07.012.

This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.



1	Clofazimine improves clinical outcomes in multidrug-resistant
2	tuberculosis: a randomized controlled trial
3	
4	Running Title: Treatment of MDR-TB with CFZ in China
5	
6	Hongfei Duan ^{1#} , Xiaohong Chen ^{2#} , Zhihui Li ^{3#} , Yu Pang ^{4#} , Wei Jing ^{1#} , Ping Liu ⁵ ,
7	Tingting Wu ⁶ , Cui Cai ⁷ , Junwei Shi ⁸ , Zhihua Qin ⁸ , Hongyun Yin ⁹ , Chao Qiu ¹⁰ ,
8	Chunxiang Li ¹¹ , Ying Xia ¹¹ , Wei Chen ¹² , Zhizhong Ye ¹³ , Zhiyue Li ¹⁴ , Gang Chen ¹⁵ ,
9	Sheng Wang ¹⁶ , Yufeng Liu ¹⁷ , Lixiang Chu ¹⁸ , Min Zhu ¹⁹ , Tao Xu ²⁰ , Qingfeng Wang ¹ ,
10	Jing Wang ¹ , Yadong Du ¹ , Jun Wang ¹ , Naihui Chu ^{1*} , Shaofa Xu ^{21*}
11	
12	¹ Department of Tuberculosis, Beijing Chest Hospital affiliated to Capital Medical
13	University, Beijing Tuberculosis & Thoracic Tumor Research Institute, Beijing,
14	101149, P.R.China
15	² Department of Tuberculosis, Pulmonary Hospital of Fuzhou of Fujian Province,
16	Fuzhou, 350008, P.R.China
17	³ Department of Tuberculosis, Hebei Chest Hospital, Shijiazhuang, 050048,
18	P.R.China
19	⁴ National Clinical Laboratory on Tuberculosis, Beijing Chest Hospital affiliated to
20	Capital Medical University, Beijing Tuberculosis & Thoracic Tumor Research
21	Institute, Beijing, 101149, P.R.China
22	⁵ Depatment of Respiratory Medicine, The People's Hospital of Changshou Distirct,
23	Chongqing, 401220, P.R.China

- ⁶Department of Respiratory Medicine, The Second Affiliated Hospital of Chengdu.
- 25 Chongqing Medical University, 610031, P.R.China
- ⁷Department of Tuberculosis, Guiyang Pulmonary Hospital, Guiyang, 550006,
- 27 P.R.China
- ⁸Department of Tuberculosis, The Sixth People's Hospital of Nantong City Jiangsu
- 29 Province, Nantong, 226011, P.R.China
- ⁹Department of Tuberculosis, Shanghai Pulmonary Hospital Affiliated to Tongji
- 31 University, Shanghai, 200433, P.R.China
- 32 ¹⁰Department of Drug-resistant tuberculosis, Jiamusi Tuberculosis Control Hospital,
- 33 Jiamusi, 154007, P.R.China
- ¹¹The Third Ward of Pulmonary Hospital, Changsha Central Hospital, Changsha,
- 35 410000, P.R.China
- ¹²Department of Tuberculosis, Shenyang Chest Hospital, Liaoning Province
- 37 Shenyang, 110044, P.R.China
- ¹³Department of Pulmonary, The First Affiliated Hospital of Xiamen university,
- 39 Xiamen, 361022, P.R.China
- 40 ¹⁴Department of Tuberculosis, Liaoyang Tuberculosis Hospital of Liaoning Province,
- 41 Liaoyang, 111000, P.R. China
- 42 ¹⁵Department of Tuberculosis, Chongqing Pulmonary Hospital, Chongqing, 404100,
- 43 P.R. China
- ¹⁶Department of Thoracic Surgery, Chest Hospital of Xinjiang Uyghur Autonomous
- 45 Region of the PRC, Wulumuqi, 830049, P.R. China
- 46 ¹⁷Department of Chest, Qingdao Chest Hospital, Qingdao, 266043, P.R.China
- 47 ¹⁸Department of Tuberculosis, The Seventh People's Hosipital of Mudanjiang,
- 48 Mudanjiang, 157011, P.R.China

- ⁴⁹ ¹⁹Tuberculosis Treatment Center, Hangzhou Red Cross Hospital, Hangzhou, 31003,
- 50 P.R.China
- ²⁰Department of Epidemiology and Biostatistics, Institute of Basic Medical Sciences,
- 52 Chinese Academy of Medical Sciences & School of Basic Medicine, Peking Union
- 53 Medical College, Beijing, 100005, P.R.China
- ⁵⁴ ²¹Department of Thoracic Surgery, Beijing Chest Hospital affiliated to Capital
- 55 Medical University, Beijing Tuberculosis & Thoracic Tumor Research Institute,
- 56 Beijing, 101149, P.R.China
- 57
- [#]These authors contributed equally to this work.
- 59
- 60 ^{*}Correspondence:
- 61 Prof. Naihui Chu
- 62 chunaihui1994@sina.com
- 63 Tel: +86-10-8950 9301
- 64 Fax: +86-10-8950 9301
- 65 Postal Address: No 9, Beiguan Street, Tongzhou District, Beijing 101149, P.R.China
- 66
- 67 Prof. Shaofa Xu
- 68 xushaofa@263.net
- 69 Tel: +86-10-8950 9188
- 70 Fax: +86-10-8950 9188
- 71 Postal Address: No 9, Beiguan Street, Tongzhou District, Beijing 101149, P.R.China
- 72
- 73

	ACCEPTED MANUSCRIPT
74	
75	
76	Abstract
77	Objectives: We carried out a randomised multicenter study in China to investigate
78	whether the clofazimine (CFZ) would improve the efficacy of the standardised
79	regimen in multidrug-resistant tuberculosis (MDR-TB) patients.
80	
81	Methods: MDR-TB patients managed in 17 TB specialised hospitals in China
82	between September 2009 and September 2011 were randomly assigned to the
83	treatment groups at enrolment. In the intervention group 100 mg CFZ per day was
84	added to the standardised regimen. The primary outcome was the proportion of
85	patients with successful outcomes.
86	
87	<i>Results:</i> From the 156 patients that were screened, 74 were assigned to the control
88	group and 66 to the CFZ group. Of the 66 cases analysed for clinical outcome in the
89	CFZ group, 36 patients were cured, and 7 completed treatment, yielding a favourable
90	outcome rate of 65.1%. The proportion of patients with favourable outcomes among
91	control regimen was 47.3% (35/74), which was significantly lower than that in the
92	CFZ group (<i>P</i> =0.034, RR=0.661, 95%CI: 0.243-0.949).
93	
94	<i>Conclusions</i> : The addition of clofazimine to the standard regimen improved the
95	treatment of MDR-TB.

- 97 Keywords: multidrug-resistant tuberculosis; clofazimine; treatment; China; adverse
 98 events
- 99

100 Introduction

- 101 Multidrug-resistant tuberculosis (MDR-TB), defined as resistance to at least
- 102 rifampicin (RIF) and isoniazid (INH), is a major public health threat that jeopardizes
- 103 the progress in TB control worldwide [1, 2]. According to an estimation by the World
- 104 Health Organization (WHO), in 2016 490,000 MDR-TB cases emerged globally, and
- 105 of these 240,000 died as a result of MDR/RIF-resistant (RR)-TB [1]. Among new and
- 106 previously treated TB cases, the proportions of MDR/RR-TB cases were 4.1% (95%
- 107 confidence interval [CI]: 2.8%-5.3%) and 19% (95% CI: 9.8%-27%), respectively [1].

108

109 China has the third largest number of TB patients worldwide [1]. Despite the steady 110 decline in the overall TB notification rate [3], the MDR-TB epidemic emerges as the 111 greatest challenge facing TB control in this country [4], with estimated rates of 7.1% 112 and 24% among new and previously treated TB cases, respectively [1]. More 113 importantly, only a small proportion of affected individuals are actually diagnosed 114 and can access proper treatment in China[5], contributing to increasing treatment 115 failures and ongoing transmission within communities.

116

Treatment of patients with MDR-TB is more complicated than those with drugsusceptible TB due to the limited efficacy of second-line drugs, an increased number
of adverse events associated with the drugs, and the long duration of therapy [6,
7].The treatment outcome of MDR-TB is generally poor, and only 48% of MDR-TB
cases worldwide achieve a favourable outcome [8]. We need novel TB drugs that are

122	active against drug-resistant bacteria [7]. Given the costly and lengthy process of new
123	drug discovery, repurposing existing drugs has emerged as an alternative strategy to
124	provide accessible anti-TB drugs for patients infected with MDR-TB [7]. Among
125	these candidate drugs, clofazimine (CFZ), a member of riminophenazine antibiotic
126	class, probably improves outcomes of MDR-TB and is classified by WHO as a group
127	C drug [9]. In 2010, a clinical trial conducted in Bangladesh revealed that a 9-month
128	treatment regimen including CFZ could cure nearly 90% of patients with MDR-TB
129	[10], indicating the potential role of CFZ for improving the treatment outcome of this
130	serious form of TB. The finding was subsequently confirmed by several observational
131	studies from other researchers [11, 12].
132	
133	To provide further evidence on the use of CFZ in the treatment of MDR-TB cases, we
134	carried out a randomised multicentre study in China focused on the potential of
135	adding CFZ to the standardised regimen. The adverse events associated with CFZ
136	were analysed to evaluate its safety in Chinese population.
137	

- 138 Methods
- 139 **Ethic statement**

The study was approved by the Medical Research Ethics Committee, Beijing Chest
Hospital, Capital Medical University (2009-28). Eligible participants infected with
MDR-TB were required to provide written informed consent. Patients could withdraw
from the trial at their own request. This study was registered after its completion
with the Chinese Clinical Trial Registry (ChiCTR, www.chictr.org.cn) under
identifier ChiCTR1800014800.

147 Study design

148 A multicentre, randomized trial was conducted among MDR-TB patients who

registered in 17 TB specialized hospital between September 2009 and September

150 2011. The study consisted of 3 phases: (1) screening; (2) treatment of intensive phase

151 (6 months); (3) treatment of consolidation phase (18 months).

152 Participants were randomized (1:1) to control or experimental group at enrolment.

153 Randomization was conducted by using a computer-generated random-number table,

154 statistical staff generated the random allocation sequence. Clinical doctors enrolled

155 participants. All participants and clinicians involving in this study were unblinded to

156 the treatment allocation. Patients in the control group received amikacin

157 (capreomycin), levofloxacin, pyrazinamide, ethambutol, para-aminosalicylic acid

158 (protionamide), and amoxicillin/clavulanate for 6 months; and then were subsequently

administered a baseline regimen of levofloxacin, pyrazinamide, ethambutol, para-

160 aminosalicylic acid (protionamide), and amoxicillin/clavulanate for 18 months. The

161 dose of drugs was listed in Table S1. Patients in the CFZ group received 100 mg of

162 CFZ per day in addition to the baseline regimen within the whole 24-month treatment

163 period. Patients and clinicians were unblinded to the treatment received throughout

164 the trial. At enrolment, data were collected on demographic and clinical

165 characteristics, including age, sex, body mass index (BMI), anti-TB treatment

- 166 duration, and co-morbidity. No changes were made to study methods after
- 167 commencement of the trial.

168

169 Participants

170 Patients were recruited from 17 hospitals in China (Table S2). Eligible patients were

171 at least 18 years of age, not pregnant, had sputum smear-positive pulmonary TB, and

172	had MDR-TB confirmed by conventional drug susceptibility testing. Reasons for
173	exclusion included: (i) XDR-TB (MDR-TB strains with additional resistance to any
174	fluoroquinolone and one injectable second-line drug); (ii) patients infected with non-
175	tuberculous mycobacteria; (iii) severe comorbidity (Table S3); (iv) previous anti-
176	tuberculosis treatment with clofazimine.
177	
178	Assessment
179	Sputum smears and solid culture were performed monthly during 2-year study period.
180	Drug susceptibility testing (DST) for four first-line anti-TB drugs (rifampicin,
181	isoniazid, ethambutol, and streptomycin) and 6 second-line drugs (amikacin,
182	capreomycin, ethionamide, para-aminosalicylic acid, ofloxacin, and levofloxacin) was
183	performed using the proportional agar method on Löwenstein–Jensen (L-J) medium
184	[14]. In addition, routine blood counts, biochemical tests, and urinalysis were assessed
185	monthly to monitor the occurrence of adverse events. Skin discoloration was defined
186	as the visible presence of reddish discoloration/pigmentation and ichthyotic changes
187	of the skin. Hepatic damage was defined as the elevation of serum transaminases to at
188	least three times the normal levels in the presence of gastrointestinal symptoms, or
189	serum transaminases to at least five times the normal levels without symptoms. Renal
190	damage was defined as the elevation of creatinine to at least 1.3 times the normal
191	levels. Adverse events were graded according to an adaptation of the AIDS Clinical
192	Trials Group Table for Grading Adverse Experiences [15]. Study regimen was
193	temporarily discontinued for all patients with grade 3 or 4 adverse events, defined as
194	serious adverse event.
195	The primary outcome was the proportion of patients with successful outcomes. The

196 clinical outcomes were assessed by the local investigator without blinding. The

197 following treatment outcome definitions were adapted from WHO guidelines. Cure 198 was defined by at least 3 consecutive negative cultures and no positive culture during 199 the last 18 months of treatment. Treatment completion was defined by bacteriological 200 conversion through the end of treatment but fewer than three consecutive negative 201 culture. Death was defined as death for any reason during the course of MDR-TB 202 treatment. Default was defined as treatment interruption for 2 or more consecutive months for any reason without medical approval. Treatment failure was defined as 203 204 persistence of two or more positive cultures of the five cultures recorded in the final 205 12 months, persistence of one or more positive cultures of the final three months, or 206 early treatment termination because of poor clinical or radiological response or 207 adverse events. Successful outcome included cure and treatment completion, while adverse outcome included any death, default, and treatment failure [16]. There were 208 209 no changes to trial outcomes after the trial commenced.

210

211 Sample size calculation

By reviewing previous studies [10, 13], we estimated that the rates of patients with 212 213 favourable outcomes at the end of treatment were 50% for the control group and 80% 214 for the CFZ group. The sample size calculation determined that 51 subjects per 215 treatment arm would provide a power of 80% to show the difference of the CFZ 216 intervention to the control regimen, assuming a one-side type I error of 0.05. In 217 addition, we estimated that 10% of the MDR patients in each study group would have 218 XDR-TB and that 20% would be loss of follow-up or default. Hence, a sample of 65 219 subjects per arm was recruited during the study period.

220

221 Data analysis

222	The original data of treatment records were entered into a computer by a double data
223	entry method using Epidata-Entry (http://www.epidata.dk/). We used SPSS 20.0 for
224	analysis. We used Chi-square analysis to investigate the clinical outcomes, occurrence
225	of adverse events of patients randomly assigned in the control and experimental
226	groups. Student's t-test were conducted for continuous variables. In addition,
227	univariate analysis and multivariate analysis were conducted to assess the potential
228	risk factors associated with a poor clinical outcome, respectively. The Kaplan-Meier
229	curve was generated to describe and compare the overall rate of bacteriological
230	conversion over a two-year period. The difference was declared as significant if the P
231	value was less than 0.05.
232	
233	Results
234	Participants
234 235	Participants Between September 2009 and September 2011, a total of 156 patients were screened,
235	Between September 2009 and September 2011, a total of 156 patients were screened,
235 236	Between September 2009 and September 2011, a total of 156 patients were screened, and 140 underwent randomisation in this study, where 74 and 66 patients were
235 236 237	Between September 2009 and September 2011, a total of 156 patients were screened, and 140 underwent randomisation in this study, where 74 and 66 patients were assigned to the control and CFZ groups, respectively. All recruited patients had a
235 236 237 238	Between September 2009 and September 2011, a total of 156 patients were screened, and 140 underwent randomisation in this study, where 74 and 66 patients were assigned to the control and CFZ groups, respectively. All recruited patients had a negative test result for the human immunodeficiency virus (HIV). The trial ended on
 235 236 237 238 239 	Between September 2009 and September 2011, a total of 156 patients were screened, and 140 underwent randomisation in this study, where 74 and 66 patients were assigned to the control and CFZ groups, respectively. All recruited patients had a negative test result for the human immunodeficiency virus (HIV). The trial ended on the date of the final follow-up of the patient who was last randomised. During the
 235 236 237 238 239 240 	Between September 2009 and September 2011, a total of 156 patients were screened, and 140 underwent randomisation in this study, where 74 and 66 patients were assigned to the control and CFZ groups, respectively. All recruited patients had a negative test result for the human immunodeficiency virus (HIV). The trial ended on the date of the final follow-up of the patient who was last randomised. During the study period, 39 patients discontinued their treatment (Fig. 1). The principal reason
 235 236 237 238 239 240 241 	Between September 2009 and September 2011, a total of 156 patients were screened, and 140 underwent randomisation in this study, where 74 and 66 patients were assigned to the control and CFZ groups, respectively. All recruited patients had a negative test result for the human immunodeficiency virus (HIV). The trial ended on the date of the final follow-up of the patient who was last randomised. During the study period, 39 patients discontinued their treatment (Fig. 1). The principal reason for discontinued treatment was a failure to follow-up (<i>n</i> =19), followed by treatment
 235 236 237 238 239 240 241 242 	Between September 2009 and September 2011, a total of 156 patients were screened, and 140 underwent randomisation in this study, where 74 and 66 patients were assigned to the control and CFZ groups, respectively. All recruited patients had a negative test result for the human immunodeficiency virus (HIV). The trial ended on the date of the final follow-up of the patient who was last randomised. During the study period, 39 patients discontinued their treatment (Fig. 1). The principal reason for discontinued treatment was a failure to follow-up (n =19), followed by treatment modification due to self-reported intolerable adverse events (n =8) and early treatment

- 246 previous treatment duration of 18 months and 24 months for control and CFZ group,
- 247 respectively. One tenth of patients had comorbidity (Table 1).
- 248

249 Treatment efficacy

- 250 Of the 66 cases analysed for clinical outcome in the CFZ group, 36 patients were
- cured, and 7 achieved treatment completion who had documented bacteriological
- conversion through the end of treatment but fewer than three consecutive negative
- culture, yielding a favourable outcome rate of 65.1%. Out of 23 patients meeting the
- criteria of adverse outcome, 4 died, 10 defaulted, and 9 failed the treatment in the
- 255 CFZ group. The proportion of patients with favourable outcomes among those
- receiving the control regimen was 47.3% (35/74, 26 cured and 9 treatment
- completion), which was significantly lower than that in the CFZ group (P=0.034,
- 258 RR=0.661, 95%CI: 0.243-0.949) (Table 2).
- 259 Of the 140 study patients, 101 with culture results were included in Kaplan–Meier
- analyses. As shown in Fig. 2, MDR patients in the CFZ group had conversion to
- 261 culture-negative status sooner than those in the control group by using mycobacterial
- 262 culture with L-J medium (P=0.031) (Fig. 2).
- 263

Adverse events

A total of 44 adverse events occurred in 44 patients in this study, including 14 in the control group and 30 in the CFZ group. There was a significant difference in the incidence of adverse events between the two groups (P=0.001). Data on the adverse events is detailed in Table 3.

270	Nine patients (9/44, 20.5%) had serious adverse events, including 3 in the control and
271	6 in the CFZ groups, respectively (Table 3 & Table S4). Anti-TB treatment in the
272	control grouped was stopped and not restarted due to a gastrointestinal reaction and an
273	occurrence of anaemia. Also, the adverse effect of the patient suffering
274	gastrointestinal reaction was resolved by stopping treatment, and the initial regimen
275	was reused after one-month of interruption. In the CFZ group, 6 different reactions
276	(two of hepatic damage, two of gastrointestinal reaction, one of renal damage, and
277	one of leukocytopenia) caused serious adverse events; thus, treatments were
278	discontinued and not restarted.
279	
280	Discussion
281	In this study involving 140 MDR-TB patients, we found that the addition of
282	clofazimine to the treatment regimen significantly improved outcomes among MDR-
283	TB patients. Similar results were observed in the prospective cohort studies from
284	Norway (86.9%) [17] and Bangladesh (87.1%) [10] and were higher than those from
285	Brazil (65.2%) [18], Shanghai (62.9%) [12] and Peru (59.9%) [19]. The discrepancy
286	across various reports may be related to the study's population and treatment regimen
287	[18]. The individuals enrolled in this study had MDR-TB instead of XDR-TB, which
288	may explain the greater treatment success rate in our study. This difference may also
289	be due to longer treatment duration of clofazimine during the whole 24-month
290	treatment period. There is evidence that an extended duration of treatment is
291	associated with favourable outcomes [20, 21]. In addition to the significant benefit
292	effect on the clinical outcomes, the cost of CFZ is more affordable compared with
293	other second-line drugs. This further highlights its use as an important candidate drug
294	against MDR-TB, especially in low-resource settings.

270	
296	Despite exhibiting promising efficacy against MDR-TB, several major concerns
297	regarding the application of CFZ should be taken into consideration in clinical
298	practice. For instance, cross-resistance to bedaquiline and clofazimine has been noted
299	by some researchers [24, 25], where prior exposure to clofazimine could cause
300	resistance to both drugs due to sharing the same efflux pump system [25]. The abuse
301	of clofazimine may facilitate the emergence of bedaquiline resistance, thereby
302	resulting in the rapid loss of this new drug. Therefore, the evaluation of in vitro CFZ
303	resistance is essential before its clinical application. Furthermore, the critical
304	concentration of CFZ has not yet to be established by WHO [14]; thus, there is an
305	urgent need to develop the accurate and reproducible DST method for CFZ.
306	
307	The beneficial effect of clofazimine was tempered, as expected, by the high rates of
308	drug-related adverse events. While skin discoloration is the most common adverse
309	event associated with the administration of CFZ [12], previous studies demonstrated
310	that lowering the dose of clofazimine to 100 mg every other day could help manage
311	the side effects of skin discoloration [12]. However, the effect of decreased exposure
312	to CFZ on clinical outcomes remains unknown. Moreover, hepatic damage (according
313	to our definition) was observed more often patients assigned to the CFZ group
314	compared with patients in the control group, though the difference did not reach
315	statistical significance due to the small sample size. Our findings indicate that routine
316	determination of hepatic enzyme levels should be performed in patients administered
317	the CFZ-containing regimen to avoid the occurrence of severe hepatic injury.
318	

319	Our study has several limitations. First, we limited our analysis to the primary
320	outcome of treatment success rate at the end of the treatment course, while the long-
321	term effect of CFZ on relapse among this cohort of MDR-TB cases was not evaluated.
322	Second, due to lack of a reliable DST method for the detection of CFZ resistance, we
323	could not assess the correlation of in vitro DST results of CFZ with clinical response
324	to treatment. Likewise, the acquired resistance following exposure to CFZ was not
325	collected in this clinical trial. Third, all patients enrolled in this study had MDR-TB,
326	which means that it was not possible to determine whether CFZ exhibits promising
327	efficacy for patients with XDR-TB. Fourth, although great efforts were focused on
328	patient follow-up, 19 out of 140 study patients failed to show for follow-up visits,
329	which increases the risk of statistical bias. Despite these limitations, our findings echo
330	the increasing evidence that the addition of CFZ is more effective in achieving
331	favourable outcomes for individuals infected with MDR-TB.
332	
333	In conclusion, our data demonstrate that the addition of clofazimine to the routine

treatment regimen exhibits promising efficacy for the treatment of MDR-TB. The
high incidences of CFZ-related skin discoloration and hepatic dysfunction highlight
the need to conduct routine examination to avoid the occurrence of serious adverse
events.

338

339 Conflicts of interest None declared.

340

341 Acknowledgments We express our thanks to Yongai Luo, Xichen Huang (The

342 First Affiliated Hospital of Chongqing Medical University), Yan Ruan, Wei Lin,

343 Sufang Fang, Guolan Wu (Fuzhou Pulmonary Hospital of Fujian), Qing Zhang

344	(Shanghai Pulmonary Hospital affiliated to Tongji University), Lisi Wang, Liming
345	Zhang (Chongqing People's Hospital), Kesar Ughur, Junlian Li (Xinjiang Uygur
346	Autonomous Region Chest Hospital) for their time and effort in data collection and
347	patients' follow-up.
348	
349	Author contributions NC and SX designed the study. HD, XC, ZL, YP, WJ, TX
350	participated in data analysis. HD, XC, ZL, PY and WJ wrote the manuscript. PL, TW,
351	CC, JS, ZQ, HY, CQ, CL, YX, WC, ZY, ZL, GC, SW, YL, LC, MZ, QW, JW, YD
352	and JW participated in data collection and patients' follow-up. All authors approved
353	the final version of the paper.
354	
355	Access to to the full data: Tao Xu
356	
357	Funding
358	This work was supported by the Ministry of Science and Technology of the People's
359	Republic of China [2017ZX09304009] and Key Clinical Specialty Discipline
360	Construction Program of Fuzhou, Fujian, P.R.C(201510302).
361	
362	References
363	[1] World Health Organization. Global Tuberculosis Report 2017. 2017.
364	[2] Gandhi NR, Nunn P, Dheda K, Schaaf HS, Zignol M, van Soolingen D, et al.

- 365 Multidrug-resistant and extensively drug-resistant tuberculosis: a threat to global
- 366 control of tuberculosis. Lancet. 2010;375:1830-43.

- 367 [3] Wang L, Zhang H, Ruan Y, Chin DP, Xia Y, Cheng S, et al. Tuberculosis
- 368 prevalence in China, 1990-2010; a longitudinal analysis of national survey data.
- 369 Lancet. 2014;383:2057-64.
- 370 [4] Zhao Y, Xu S, Wang L, Chin DP, Wang S, Jiang G, et al. National survey of drug-
- 371 resistant tuberculosis in China. N Engl J Med. 2012;366:2161-70.
- 372 [5] Li Y, Ehiri J, Tang S, Li D, Bian Y, Lin H, et al. Factors associated with patient,
- and diagnostic delays in Chinese TB patients: a systematic review and meta-analysis.
- 374 BMC Med. 2013;11:156.
- [6] Caminero JA, Sotgiu G, Zumla A, Migliori GB. Best drug treatment for
- 376 multidrug-resistant and extensively drug-resistant tuberculosis. Lancet Infect Dis.
- 377 2010;10:621-9.
- 378 [7] Zumla A, Nahid P, Cole ST. Advances in the development of new tuberculosis
- drugs and treatment regimens. Nat Rev Drug Discov. 2013;12:388-404.
- [8] Kwon YS, Kim YH, Suh GY, Chung MP, Kim H, Kwon OJ, et al. Treatment
- 381 outcomes for HIV-uninfected patients with multidrug-resistant and extensively drug-
- 382 resistant tuberculosis. Clin Infect Dis. 2008;47:496-502.
- 383 [9] World Health Organization. WHO treatment guidelines for drug-resistant
- tuberculosis (2016 update). 2016.
- 385 [[10] Van Deun A, Maug AK, Salim MA, Das PK, Sarker MR, Daru P, et al. Short,
- 386 highly effective, and inexpensive standardized treatment of multidrug-resistant
- tuberculosis. Am J Respir Crit Care Med. 2010;182:684-92.
- 388 [11] Tang S, Yao L, Hao X, Liu Y, Zeng L, Liu G, et al. Clofazimine for the
- 389 treatment of multidrug-resistant tuberculosis: prospective, multicenter, randomized
- 390 controlled study in China. Clin Infect Dis. 2015;60:1361-7.

- 391 [12] Xu HB, Jiang RH, Xiao HP. Clofazimine in the treatment of multidrug-resistant
- tuberculosis. Clin Microbiol Infect. 2012;18:1104-10.
- 393 [13] P.G. Suarez, K. Floyd, J. Portocarrero, E. Alarcon, E. Rapiti, G. Ramos, C. et al.
- 394 Feasibility and cost-effectiveness of standardised second-line drug treatment for
- 395 chronic tuberculosis patients: a national cohort study in Peru. Lancet.
- 396 2002;359(9322):1980-9.
- 397 [14] World Health Organization. Guidelines for surveillance of drug resistance in
- tuberculosis 5th edition. 2009.
- 399 [15] DAIDS. Division of AIDS Table for Grading the Severity of Adult and Pediatric
- 400 Adverse Events. Bethesda, MD, USA: DAIDS.2004.
- 401 [16] Laserson KF, Thorpe LE, Leimane V, Weyer K, Mitnick CD, Riekstina V, et al.
- 402 Speaking the same language: treatment outcome definitions for multidrug-resistant
- 403 tuberculosis. Int J Tuberc Lung Dis. 2005;9:640-5.
- 404 [17] von der Lippe B, Sandven P, Brubakk O. Efficacy and safety of linezolid in
- 405 multidrug resistant tuberculosis (MDR-TB)--a report of ten cases. J Infect.
- 406 2006;52:92-6.
- 407 [18] Dey T, Brigden G, Cox H, Shubber Z, Cooke G, Ford N. Outcomes of
- 408 clofazimine for the treatment of drug-resistant tuberculosis: a systematic review and
- 409 meta-analysis. J Antimicrob Chemother. 2013;68:284-93.
- 410 [19] Mitnick CD, Shin SS, Seung KJ, Rich ML, Atwood SS, Furin JJ, et al.
- 411 Comprehensive treatment of extensively drug-resistant tuberculosis. N Engl J Med.
- 412 2008;359:563-74.
- 413 [20] Van Deun A, Salim MA, Das AP, Bastian I, Portaels F. Results of a standardised
- 414 regimen for multidrug-resistant tuberculosis in Bangladesh. Int J Tuberc Lung Dis.
- 415 2004;8:560-7.

- 416 [21] Masjedi MR, Tabarsi P, Chitsaz E, Baghaei P, Mirsaeidi M, Amiri MV, et al.
- 417 Outcome of treatment of MDR-TB patients with standardised regimens, Iran, 2002-
- 418 2006. Int J Tuberc Lung Dis. 2008;12:750-5.
- 419 [22] Leimane V, Riekstina V, Holtz TH, Zarovska E, Skripconoka V, Thorpe LE, et
- 420 al. Clinical outcome of individualised treatment of multidrug-resistant tuberculosis in
- 421 Latvia: a retrospective cohort study. Lancet. 2005;365:318-26.
- 422 [23] Leimane V, Dravniece G, Riekstina V, Sture I, Kammerer S, Chen MP, et al.
- 423 Treatment outcome of multidrug/extensively drug-resistant tuberculosis in Latvia,
- 424 2000-2004. Eur Respir J. 2010;36:584-93.
- 425 [24] Hartkoorn RC, Uplekar S, Cole ST. Cross-resistance between clofazimine and
- 426 bedaquiline through upregulation of MmpL5 in Mycobacterium tuberculosis.
- 427 Antimicrob Agents Chemother. 2014;58:2979-81.
- 428 [25] Almeida D, Ioerger T, Tyagi S, Li SY, Mdluli K, Andries K, et al. Mutations in
- 429 pepQ Confer Low-Level Resistance to Bedaquiline and Clofazimine in
- 430 Mycobacterium tuberculosis. Antimicrob Agents Chemother. 2016;60:4590-9.
- 431
- 432
- 433
- 434 Figure Legends
- 435 **Figure 1 Enrolment and follow-up of the study patients.**
- 436 **Figure 2 Time to sputum-culture conversation in the control and experimental**
- 437 groups

	Experimental group	Control group
Characteristic	(N=66)	(N= 74)
Ageyears		
Mean	36.8	36.4
Range	19~65	18~61
Male sexno.(%)	44(66.7)	44(59.5)
Body mass indexkg/m ²		
Mean	19.9	19.8
Range	15.0~27.3	14.0~25.7
Treatment history	(
New casesno.(%)	3 (4.5)	5 (6.8)
Previously treatedno.(%)	63 (95.5)	69 (93.2)
Treatment duration of previously		
treated patientsmonths		
Mean	29.9	23.0
Range	1~140	1~120
Co-morbidityno.(%)		
Diabetes	2 (3.0)	2 (2.7)
COPD ^a	2 (3.0)	2 (2.7)
Cardiopathy	1 (1.5)	1 (1.4)

Table 1 Demographic and clinical characteristics of MDR-TB patients enrolled in this study

^aCOPD, chronic obstructive pulmonary disease.

Live pulmon.

Treatment outcome	No. of patien	of patients (%)	
	Experimental group	Control group	-
	(N=66)	(N= 74)	
Favorable outcome			0.034
Cure	36 (54.5)	26 (35.1)	
Treatment completion	7 (10.6)	9 (12.2)	
Adverse outcome			
Treatment failure	9 (13.6)	24 (32.4)	
Death	4 (6.1)	2 (2.7)	
<i>Default^a</i>	10 (15.2)	13 (17.6)	

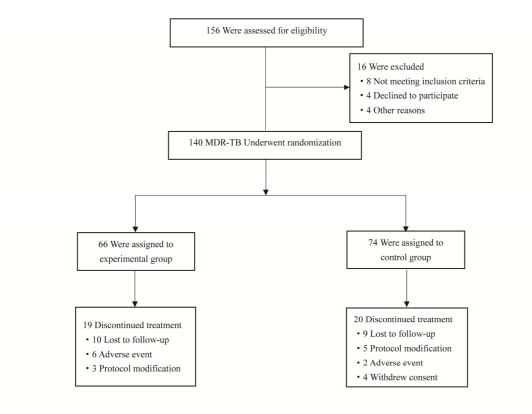
Table 2 Treatment outcomes of patients with multidrug-resistant tuberculosis

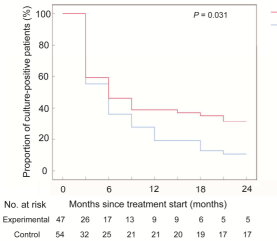
^aFour patients withdrawing consent due to in control group are classified into default category.

Adverse event	No. of patients (%)		P value
	Experimental group	Control group	_
	(N=66)	(N= 74)	
Skin discoloration	8 (12.1)	0 (0.0)	0.002
Hepatic damage	8 (12.1)	2 (2.7)	0.046
Hyperuricemia	3 (4.5)	2 (2.7)	0.667
Gastrointestinal reaction	3 (4.5)	5 (6.8)	0.722
Others ^a	8(12.1)	5 (6.8)	0.275

 Table 3 Adverse events during 24-month treatment among patients enrolled in this study

^aOther adverse events include renal damage, rash, leukocytopenia, anemia, arthralgia and hearing loss.





Control group Experimental group