

***In Vitro* efficacy comparison of linezolid, tedizolid, sutezolid and delpazolid against rapid growing Mycobacteria isolated in Beijing, China**

3

4 Shuan Wen^{1§}, Xiaopan Gao^{2§}, Weijie Zhao^{3§}, Fengmin Huo¹, Guanglu Jiang¹, Lingling Dong¹,
5 Liping Zhao¹, Fen Wang¹, Xia Yu^{1*}, Hairong Huang^{1*}.

6 1 National Clinical Laboratory on Tuberculosis, Beijing Key laboratory for Drug-resistant
7 Tuberculosis Research, Beijing Chest Hospital, Capital Medical University, Beijing, China;
8 2MOH Key Laboratory of Systems Biology of Pathogens, Institute of Pathogen Biology, Chinese
9 Academy of Medical Sciences & Peking Union Medical College, Beijing, China.

10 3 The administration office of clinical trial, Beijing Chest Hospital, Capital Medical University,
11 Beijing, China

12

13

14 §These authors contributed equally to this study. Author order was determined by working load.

15

16

17

18 *Correspondence should be addressed to Dr. Huang at huanghairong@tb123.org or Dr. Yu at
19 yuxiasmart@163.com, BeiguanSt, No. 9, Tongzhou Qu, Beijing 101149, China.

20

21

22 **Running title:** The efficacy of oxazolidinones against RGM

23

ABSTRACT

The natural resistance of rapid growth Mycobacterium (RGM) against multiple antibiotics renders the treatment of caused infections less successful and time consuming. Therefore, new effective agents are urgently needed. The aim of this study was to evaluate the *in vitro* susceptibility of 115 isolates, constituting different RGM species, against four oxazolidinones i.e.delpazolid,sutezolid,tedizolid and linezolid.Additionally,32 reference strains of different RGM species were also tested.The four oxazolidinones exhibited potent *in vitro* activity against the recruited RGM reference strains,24 out of 32 RGM species had MICs $\leq$ 8 μ g/mL against all four oxazolidinones whereas tedizolid and delpazolid generally presented lower MICs than linezolid or sutezolid. Tedizolid showed the strongest activity against clinical isolates of *M.abscessus* with MIC₅₀=1 μ g/mL and MIC₉₀=2 μ g/mL.MIC values for tedizolid were usually 4- to 8-fold less than the MICs of linezolid for *M.abscessus* subsp. *abscessus*.The MIC distributions of sutezolid and linezolid were similar, while delpazolid showed 2-fold lower MIC as compared with linezolid.Linezolid was not active against most of the tested *M.fortuitum* isolates, 22 out of the 25 *M.fortuitum* were resistant against linezolid. However,delpazolid exhibited better antimicrobial activity against these isolates with 4-fold lower MIC values,in contrast with linezolid.In addition,the protein alignment of RplC and RplD and structure based analysis showed that there may be no correlation between oxazolidinones resistance and mutations in *rplC*,*rplD* and *23srRNA* genes in tested RGM.This study showed tedizolid harbors the strongest inhibitory activity against *M.abscessus in vitro*,while delpazolid presented the best activity against *M.fortuitum*,which provided important insights on the potential clinical application of oxazolidinones to treat RGM infections.

KEYWORDS: Rapidgrowing Mycobacteria, delpazolid, sutezolid, tedizolid, linezolid, antimicrobial activity

INTRODUCTION

Non-tuberculous mycobacteria (NTM) are recognized as important opportunistic pathogens of humans that can cause pulmonary infection, lymphadenitis, skin abscesses, disseminated infection and systematic infection. The prevalence of NTM infections has increased globally and even surpassed tuberculosis (TB) in certain countries (1-5). According to their speed of growth (i.e. appearance of visible colonies within or after 7 day cultivation on solid medium), NTM can be categorized as rapid growing Mycobacteria (RGM) or slow growing mycobacteria (SGM). Compared with SGM, RGM are more resistant to conventional anti-TB agents and other general antibiotics, therefore, increasing the chances of treatment failure(6). *M.abscessus* and *M.fortuitum* are among the most frequently isolated and pathogenic RGMs(1-5). *M.abscessus* often cause severe pulmonary infections with poor clinical outcomes and have been frequently reported to cause soft tissue infections(7). *M.fortuitum* can cause soft tissue infection during trauma and surgery, while lung disease caused by them is rare(8). The limited efficacies and availability of only fewer choices of medications highlight the requirement of identifying new and more potent antimicrobials against RGMs.

Oxazolidinones have demonstrated promising efficacies against *M.tuberculosis* (TB) *in vitro* and *in vivo*. Due to their distinct mechanism of action (binding to the 23S ribosome, thereby blocking microbial protein synthesis) without cross-resistance to the existing TB drugs. Oxazolidinones are proposed to be used for the treatment of multiple drug resistant TB. Linezolid (LZD), licensed in 2000, is an oxazolidinone which exhibited excellent antibacterial activity against drug resistant tuberculosis (DR-TB) and NTM infection(9-12). However, serious hematologic and neurologic toxicities can be caused by LZD during long term therapy due to its inhibition of mitochondrial protein synthesis which often requires dose reduction or discontinuation(13). Thus, new oxazolidinones drugs with superior efficacy and reduced toxicity are continuously sought.

Recently, three new next-generation oxazolidinones have been developed for potential use against DR-TB. Tedizolid (TZD) phosphate is a novel, potent

oxazolidinone pro-drug that has been approved by the American FDA (2014) and the European Medicine Agency (2015) for treatment of acute bacterial skin and soft tissue infections(7). The pharmacokinetic/pharmacodynamics properties of TZD allow it to be administered orally once daily, facilitating its usage in prolonged treatment course. Sutezolid (SZD) (PNU-100480) is a thiomorpholinyl analog of LZD with preliminary evidence of superior efficacy against *M. tuberculosis*(14). SZD was found to be generally safe, well tolerated in TB patients, and with readily detectable bactericidal activity in sputum and blood. Delpazolid (LCB01-0371)(DZD) is a thiomorpholinyl analog of LZD, which showed superior efficacy against *M.tuberculosis* in the hollow-fiber, mouse model, and whole-blood model (2–4). DZD was well tolerated and showed bacteriostatic and bactericidal activity comparable to LZD against *S. aureus*, *E. faecalis* and methicillin-resistant *Staphylococcus aureus* in a recently completed phase I clinical trial(15).

To better understand the efficacies of these three new-generation oxazolidinones against different RGM species, we detected the MICs of 32 RGM reference strains and 115 RGM clinical isolates collected in Beijing, China. Furthermore, we investigated the three reported LZD-resistance genes (including *rplC*, *rplD* and *23srRNA*) from different RGM species to identify their potential relationships with oxazolidinone resistance.

RESULTS

MICs of SZD,TZD,DZD and LZD against RGM reference strains

The MICs of the 32 reference strains against SZD, TZD, DZD and LZD are presented in Table 1. All four oxazolidinones exhibited antimicrobial activities *in vitro* against the recruited RGM reference stains. Majority of the species had MICs equal to or below 8μg/mL for all four drugs. Only *M.fortuitum* and *M.rhodesiae* had MICs greater than 32μg/mL. Generally, a given isolate presented uniform tendency against all four oxazolidinones, the MIC values were either high for or low for the four drugs. For *M.abscessus*, the efficacy of TZD was stronger than LZD.

The MIC distributions of *M.abscessus* and *M.massiliense* against LZD,TZD,SZD and DZD

The MIC distributions of *M.abscessus* and *M.massiliense* against LZD, TZD,

SZD and DZD are shown in Figure 1. MICs for TZD were generally 4- to 8-fold less than the MICs of LZD for the two species. The MIC distribution of SZD was similar to LZD, while DZD values were generally half of LZD. Notably, TZD showed strongest activity against *M.abscessus* with MIC₅₀=1μg/mL and MIC₉₀=2μg/mL. According to the CLSI resistance criteria for LZD(16), the susceptibility rate of *M.abscessus* against LZD, TZD, SZD and DZD was 73.5%(36/49), 100%(49/49), 71.4%(35/49), 87.8%(43/49), respectively. The susceptibility rate of *M.massiliense* against LZD, TZD, SZD and DZD was 65.8%(23/35), 82.9%(29/35), 68.6%(24/35) and 74.3%(26/35), respectively. In general, the MIC distributions of *M.massiliense* had uniform tendency than *M.abscessus*, but with an exception for TZD. The MICs of *M.massiliense* isolates were higher than *M.abscessus*. 6 out of 35 isolates of *M.massiliense* had MICs ≥ 16μg/mL against TZD, the MICs of all the tested *M.abscessus* were ≤ 4μg/mL. In addition, the MIC outcomes for species with less than five isolates are presented in Table 2.

The MIC distributions of *M.fortuitum* against LZD, TZD, SZD and DZD

The MIC distributions of *M.fortuitum* against LZD, TZD, SZD and DZD are shown in Figure 2. In contrast to *M.abscessus* and *M.massiliense*, *M.fortuitum* presented higher percentage of resistance against the four oxazolidinones. The susceptibility profiles of the clinical isolates exhibited much lower MICs than *M.fortuitum* ATCC6481 reference strain. In Total, 88%(22/25) of the clinical isolates were resistant to LZD, including *M.fortuitum* reference strain. The *in vitro* activity of DZD was relatively better than LZD as indicated by its 2- to 4-fold lower MIC. The MIC distributions of TZD was similar to LZD as only 5 out of 25 isolates indicated MIC ≤ 8 μg /mL. According to the cutoff value of LZD, the susceptibility rates of *M.fortuitum* against TZD, SZD and DZD were 20%(5/25), 12%(3/25), 76%(19/25), respectively.

Alternations in the Oxazolidinones target sites

The entire 23SrRNA, *rplC*, and *rplD* genes were sequenced to identify the potential mutations associated with oxazolidinones resistance. The sequences of the tested clinical isolates of *M.abscessus*, *M.massiliense* and *M.fortuitum* were compared

with their corresponding reference strains. For *M.massiliense* isolates, Ala177Pro in *rplD* was detected in 12 isolates with MIC of LZD $\geq 2 \mu\text{g/mL}$. In addition, two types of synonymous SNPs within the coding region of *rplC* were also observed both in LZD resistant and susceptible isolates, including Leu86Leu(CTG $\rightarrow$ CTT) and Ala92Ala(GCG $\rightarrow$ GCT). A2271G in 23S rRNA was found in one isolate with MIC of LZD = $8 \mu\text{g/mL}$. For *M.abscessus* isolates, no non-synonymous mutation in the coding gene of *rplC* and *rplD* was observed, while most frequently observed mutation i.e. T2650C (n=2) was found in 23S rRNA with MIC of LZD $\geq 2 \mu\text{g/mL}$ (Table 3).

Among the tested *M.fortuitum* isolates, all MICs for LZD were above $2 \mu\text{g/mL}$. No non-synonymous mutation was detected in the *rplC* gene. Among 25 tested *M.fortuitum* isolates, A2090T and C1944T in 23S rRNA were detected in two isolates with MIC = $32 \mu\text{g/mL}$ and $8 \mu\text{g/mL}$ for LZD, respectively. In addition, 21 out of 25 clinical *M.fortuitum* isolates simultaneously showed following nine non-synonymous mutations in the coding protein of *rplD* for the both LZD susceptible and resistant isolates: Ala146Gly(GCG $\rightarrow$ GGC), Thr147Ser(ACC $\rightarrow$ AGC), Val156Ile(GTG $\rightarrow$ ATC), Ala161Thr(GCG $\rightarrow$ ACC), Lys167Arg(AAG $\rightarrow$ CGC), Ser207Ala(TCC $\rightarrow$ GCG), Glu212Gly(GAG $\rightarrow$ GGA), Val213Ala(GTG $\rightarrow$ GCG), Ala215Val(GCC $\rightarrow$ GTC) (Table 4).

Structural mapping of clinical mutants

For *M.massiliense* isolates, Ala177Pro in RplD was detected in 12 isolates, both in LZD susceptible and resistant isolates with MIC $\geq 2 \mu\text{g/mL}$. To gain an insight into the functional relevance of RplC and RplD mutation, multiple sequence alignment of RplC and RplD homologues from different mycobacterial species were performed (Figure S1 and S2). The protein sequence of RplC and RplD in different mycobacterial species are highly conserved. In addition, we used *M. tuberculosis* RplD structure as a model to map *M.massiliense* RplD mutation (PDB ID: 5V7Q) (Figure 5B). The structure shows that Ala177 is located in a high variable region between $\beta 3$ and $\eta 2$ and is far from the LZD binding site which indicates that this mutation may not be related to LZD resistance (Figure 3B). Next, we mapped the 23S rRNA functional mutations of *M.abscessus*, *M.massiliense* and *M.fortuitum*. The

results showed that except A2271 in *M.massiliense*, the other mutations including G2582, A2625 and T2650 were far from the catalytic center (Figure 3C).

DISCUSSION

The treatment of RGM infection is often very difficult because of their higher drug resistance rate than SGM and unavailability of highly potent drugs against them *in vitro*. *M.abscessus* complex and *M.fortuitum* are two most prevalent RGM species around the world. Infections due to *M.abscessus* carry a poor prognosis since this RGM is, for all the correct reasons, considered an “antibiotic nightmare”(17). Thus, identifying drugs that could work potently against *M.abscessus* is a priority. *M.massiliense* is a species that originally split from *M.abscessus* but they are located closely in the phylogenetic tree(18). The treatment response rates to clarithromycin-based antibiotic therapy are much higher in patients with *M.massiliense* than patients with *M.abscessus* lung disease(19). *M.fortuitum* is the main RGM responsible for extra-pulmonary disease, especially in cutaneous and plastic surgery-related infections(20). In contrast to *M.abscessus*, *M.fortuitum* infection has better prognosis due to some available effective drugs(21). However, its emerging drug resistance highlights the need for new and effective drugs(21-23). Several studies have verified the efficacy of LZD in MDR-TB or even in XDR-TB treatment (9, 13, 24). A few studies also proved its antibacterial activity against NTM species either *in vitro* or *in vivo*(25, 26). As a novel oxazolidinone prodrug, TZD exhibited greater potency than LZD against *M. tuberculosis*(6, 27) as well as against NTM(28, 29). Limited studies or no study has been performed to evaluate the efficacy of SZD and DZD against NTM species(28), whereas only a few studies provided preliminarily assessment of their potential usage in TB(14, 30, 31). In this study, we evaluated the efficacies of four oxazolidinones against the reference and clinical isolates of RGM to gain insights on their potential use for specific RGM species.

As new drugs, well recognized susceptibility testing methods for TZD, SZD and DTD have not been developed and the breakpoints to define drug resistance for them have never been discussed yet. Therefore, the MIC data of different RGM species against oxazolidinones still remain scarce. In this study, the four oxazolidinones

exhibited promising activities *in vitro* against the recruited RGM reference stains. The absolute majority of species had MICs below 8µg/mL against the four drugs. However, different species presented non-uniform susceptibility patterns. The MIC distributions of *M.massiliense* had similar tendency to the *M.abscessus*, but the MICs of TZD were obviously higher than *M.abscessus*. In comparison with other oxazolidinones, the MIC values for TZD were the lowest for both *M.abscessus* and *M.massiliense*. Previous studies, including 170 isolates of RGM, showed equivalent or lower (1 to 8 fold) MIC₅₀ and MIC₉₀ values for TZD in contrast with LZD(29). Furthermore, TZD harbors several advantages over LZD in terms of tolerability, safety, dosing frequency, and treatment duration(32). Only a few studies have reported the clinical use of TZD for the treatment of NTM infections. Our results indicated that its usage seems reasonable for the treatment of infection caused by *M.abscessus* and *M.massiliense*. Among the 25 tested *M.fortuitum* isolates in our study, 22(22/25) strains had MICs of LZD at ≥16 mg/L. Based on the CLSI criteria, these strains could be categorized as intermediate resistant or resistant strains, 52% (13/25) of them belong to resistant strains. Using the cutoff value of LZD as the tentative breakpoints, the susceptibility rate of *M.fortuitum* against TZD, SZD and DZD were 20%(5/25), 12%(3/25), 76%(19/25), respectively. DZD exhibited the best antimicrobial activity against the *M.fortuitum*. However, whether this *in vitro* outcome reflects the *in vivo* efficacy or not, requires further investigation.

A major limitation of this study was that no recommended breakpoint of different NTM species against TZD, SZD or DZD had been proposed previously. Beside *in vitro* MIC distributions, the breakpoint determination also correlates with clinical treatment response and pharmacokinetic/pharmacodynamics (PK/PD) data including drug dose. The clinical trial on these new oxazolidinones are either unavailable or very limited. A few studies have been performed on the pharmacokinetic analysis of these drugs. Generally, all the drugs were well-tolerated, and the C_{max} were highly dose-dependent. Recently, Choi et al demonstrated that, after multiple doses of TZD up to 1200mg twice daily for 21days, the peak serum concentration was 16.3µg/mL, which is comparable with peak serum concentrations of LZD=12.5µg/mL at the

dosage of 300mg twice daily(15, 33). In another study, a single 800mg dose of DZD under fasting condition acquired C_{max} at 11.74 μ g/mL(34). STD presented superior efficacy than LZD against experimental murine model of tuberculosis. The C_{max} of its major metabolite PNU-101603, which contributes to its activity, was 6.46 μ g/mL at given dose of 1200mg QD (40). However, since the optimal dosage of these next-generation oxazolidinones is still under investigation, the appropriate breakpoints for the susceptibility definition of these drugs remain beyond known.

LZD works by binding to the peptidyl transferase center of the 50S ribosomal subunit, which is composed of 5S and 23S rRNAs and 36 riboproteins (L1 through L36)(35). Recently, the Cryo-EM structure of the large ribosomal subunit from *M.tuberculosis* bound with a potent LZD analog (LZD-114) was determined(36). LZD-114 is similar with LZD in C ring but different in A and B ring that lacks a fluorine group in the B-ring while the original morpholine ring is replaced by a thiazole ring in the A-ring(Figure 3A). The LZD-114 also was bound in the same pocket and in a similar orientation to LZD in other species(37, 38). The structure showed that *rplC* encoded ribosomal protein L3 and *rplD* encoded ribosomal protein L4 bind directly to 23S ribosomal RNA and was placed relatively close to the LZD binding site on the ribosomes, suggesting that the mechanism for reduced susceptibility may include structural perturbation of the LZD binding site (PDB:5V7Q). Furthermore, previous studies demonstrated that mutations in *rplC* and *rplD* could lead to LZD resistance in *M.tuberculosis*(12, 39). However, there is no non-synonymous mutation in *rplC* against the tested RGM. Ala77Pro mutation was detected in *rplD* which is located in variable site and is far away from LZD-binding site, as shown by the sequence alignment. Except A2271G mutation in 23SrRNA in *M.massiliense* that was closer to binding site of LZD, other mutations are far from the LZD-binding site. Our results combining MIC test, gene mutation and structure based analysis showed there is no obvious correlation between riboproteins mutations (*rplC* and *rplD*) and LZD resistance identified in this study in the RGM species. Mutations located in the LZD binding site may cause LZD resistant. Hence, *rplC*, *rplD* and 23srRNA homologues might not be the only target for LZD to explore its

bacteriostatic activity.

In conclusion, this study demonstrated that oxazolidinones have good *in vitro* activities against the overwhelming majority of RGM species. The efficacies of the four oxazolidinones were variable against different species. TZD showed strongest antimicrobial activity against *M.abscessus* and *M.massiliense*, while DZD owned the strongest activity against *M.fortuitum*. The data provided important insights on the possible clinical applications of oxazolidinones to treat RGM infections.

MATERIAL AND METHODS

Ethics statement

As the study only concerned laboratory testing of mycobacteria without the direct involvement of human subjects, ethics approval was not sought.

Reference strains and clinical isolates

The mycobacterial reference strains stored in the Bio-bank in Beijing Chest Hospital (Beijing, China) were tested against LZD, TZD, SZD and DZD *in vitro*, including 32 RGM species. These reference strains were obtained either from the American Type Culture Collection (ATCC) or from the German Collection of Microorganisms (DSM). The species constitution of these reference strains are listed in Table 1. *M.massiliense* reference strain was not included due to its absence in our stock. One-hundred fifteen isolates of RGM were recruited in Beijing chest hospital from 2016 to 2018 that included 49 *M.abscessus*, 35 *M.massiliense*, 25 *M.fortuitum*. The species constitution of the remaining 6 isolates is presented in Table 2.

All of the 115 RGM clinical strains were isolated from tuberculosis suspected patients. The strains were classified as RGM preliminarily with p-nitrobenzoic acid containing medium, and then were identified by gene sequencing as indicated for each species by 16S rRNA, *hsp65*, *rpoB*, 16-23S rRNA internal transcribed spacer sequencing (40). All the isolates were stored at -80°C and sub-cultured on LJ medium before performing drug susceptibility test.

Minimal inhibitory concentration (MIC) testing

TZD phosphate and LZD were purchased from Toronto Research Chemicals and Sigma-aldrich, respectively. SZD and DZD were purchased from Shanghai Biochempartner Co., Ltd (Shanghai, China) and JHK BioPharma, respectively. Oxazolidinones were dissolved in dimethyl sulfoxide (DMSO). Stock solutions were aseptically prepared at concentrations of 2.56 mg/mL. Broth microdilution method was performed according to the guidelines of Clinical and Laboratory Standards Institute (CLSI)(41). Cation-adjusted Mueller-Hinton broth (CAMHB) was used for MIC test. The inoculum was prepared with fresh culture grown on Lowenstein-Jensen medium. The broth microdilution format was set up as 2-fold dilution, the concentrations of all the tested drugs ranged from 0.063 μ g/mL to 32 μ g/mL. Briefly, a bacterial suspension of 0.5 McFarland standard was prepared, and then diluted and inoculated into 96-well microtiter plate to achieve final bacterial load at 10⁵ colony forming unit (CFU) per well. Plates were then incubated at 37°C for 3 days for RGM. 70 μ L solution containing 20 μ L AlamarBlue (Bio-rad) and 50 μ L Tween80 (5%) was added to each well and incubated for 24 h at 37 °C before assessing color development. A change from blue to pink or purple indicated bacterial growth (42). The MIC was defined as the lowest concentration of antibiotic that prevented a color change from blue to pink.

The breakpoint of LZD was adopted from the CLSI document M24-A2 (susceptible: $\leq$ 8 mg/L; intermediate resistant: 16 mg/L; resistant: $\geq$ 32 mg/L) (16). Since no well-recognized breakpoint has been proposed for TZD, SZD or DZD, a preliminary data analysis was performed for them referring the breakpoint of LZD.

Mutations conferring oxazolidinones resistance and protein Alignment

Sequencing of PCR products was performed using the Sanger method with primers designed to be specific for *rplC*, *rplD* and 23S *rRNA*. We used previously described primers for 23S *rRNA*(43), and design new primers for *rplC*, *rplD* sequencing. The primers used in this study are listed in Table S1 in the supplemental material and were synthesized by Tsingke Biotech Co. (Beijing, China). The *rplC* and *rplD* gene of the reference strains of three RGM species plus *M.tuberculosis* were also sequenced, mutation was defined in contrast with the sequences of the reference

strains. The sequences of *M.massiliense* were adopted from website for alignment. The amplification products were sequenced by Tsingke Company (Beijing, China). Multiple sequence alignment of the homologous proteins was performed using the Clustal Omega software. Structure-based multiple sequence alignment was performed with ESPript 3 based on the crystal structure of RplC and RplD protein of *M.tuberculosis* from the following URL:<http://esprict.ibcp.fr/ESPript/ESPript/>.

Quality control. The MIC for quality control strains was determined using each lot of the prepared microtiter plates, and the results for LZD were within the expected range.

ACKNOWLEDGEMENT

This study was supported by research funding from the Infectious Diseases Special Project, Ministry of Health of China (2018ZX10302302-004-005, 2018ZX10201301302-004.) the Natural Science Fund of China (81672065 , 81802057), Beijing Municipal Administration of Hospitals Clinical Medicine Development of Special Funding Support (ZYLX201824) and Beijing Municipal Administration of Hospitals' Ascent Plan (DFL20181602).

TRANSPARENCY DECLARATIONS

None to declare.

1. Cowman S, van Ingen J, Griffith DE, Loebinger MR. 2019. Non-tuberculous mycobacterial pulmonary disease. *Eur Respir J* 54.
2. Santin M, Barrabeig I, Malchair P, Gonzalez-Luquero L, Benitez MA, Sabria J, Palau-Benavent M, Canete C, Lloret-Queralt JA, Grijota-Camino MD, Dorca J, Alcaide F. 2018. Pulmonary Infections with Nontuberculous Mycobacteria, Catalonia, Spain, 1994-2014. *Emerg Infect Dis* 24:1091-1094.
3. Lin C, Russell C, Soll B, Chow D, Bamrah S, Brostrom R, Kim W, Scott J, Bankowski MJ. 2018. Increasing Prevalence of Nontuberculous Mycobacteria in Respiratory Specimens from US-Affiliated Pacific Island Jurisdictions(1). *Emerg Infect Dis* 24:485-491.
4. Brode SK, Marchand-Austin A, Jamieson FB, Marras TK. 2017. Pulmonary versus Nonpulmonary Nontuberculous Mycobacteria, Ontario, Canada. *Emerg Infect Dis* 23:1898-1901.
5. Yu X, Liu P, Liu G, Zhao L, Hu Y, Wei G, Luo J, Huang H. 2016. The prevalence of non-tuberculous mycobacterial infections in mainland China: Systematic review and

meta-analysis. *J Infect* 73:558-567.

6. Ruiz P, Causse M, Vaquero M, Casal M. 2019. In Vitro Activity of Tedizolid against *Mycobacterium tuberculosis*. *Antimicrob Agents Chemother* 63.

7. Compain F, Soroka D, Heym B, Gaillard JL, Herrmann JL, Dorcène D, Arthur M, Dubee V. 2018. In vitro activity of tedizolid against the *Mycobacterium abscessus* complex. *Diagn Microbiol Infect Dis* 90:186-189.

8. Celdran A, Esteban J, Manas J, Granizo JJ. 2007. Wound infections due to *Mycobacterium fortuitum* after polypropylene mesh inguinal hernia repair. *J Hosp Infect* 66:374-7.

9. Singh B, Cocker D, Ryan H, Sloan DJ. 2019. Linezolid for drug-resistant pulmonary tuberculosis. *Cochrane Database Syst Rev* 3:CD012836.

10. Bilgin H, Tukenmez-Tigen E. 2019. Linezolid for drug-susceptible tuberculosis. *Lancet Infect Dis* 19:357.

11. Bolhuis MS, Akkerman OW, Sturkenboom MGG, Ghimire S, Srivastava S, Gumbo T, Alffenaar JC. 2018. Linezolid-based Regimens for Multidrug-resistant Tuberculosis (TB): A Systematic Review to Establish or Revise the Current Recommended Dose for TB Treatment. *Clin Infect Dis* 67:S327-S335.

12. Zong Z, Jing W, Shi J, Wen S, Zhang T, Huo F, Shang Y, Liang Q, Huang H, Pang Y. 2018. Comparison of In Vitro Activity and MIC Distributions between the Novel Oxazolidinone Delpazolid and Linezolid against Multidrug-Resistant and Extensively Drug-Resistant *Mycobacterium tuberculosis* in China. *Antimicrob Agents Chemother* 62.

13. Lee M, Lee J, Carroll MW, Choi H, Min S, Song T, Via LE, Goldfeder LC, Kang E, Jin B, Park H, Kwak H, Kim H, Jeon HS, Jeong I, Joh JS, Chen RY, Olivier KN, Shaw PA, Follmann D, Song SD, Lee JK, Lee D, Kim CT, Dartois V, Park SK, Cho SN, Barry CE, 3rd. 2012. Linezolid for treatment of chronic extensively drug-resistant tuberculosis. *N Engl J Med* 367:1508-18.

14. Wallis RS, Dawson R, Friedrich SO, Venter A, Paige D, Zhu T, Silvia A, Gobey J, Ellery C, Zhang Y, Eisenach K, Miller P, Diacon AH. 2014. Mycobactericidal activity of sutezolid (PNU-100480) in sputum (EBA) and blood (WBA) of patients with pulmonary tuberculosis. *PLoS One* 9:e94462.

15. Cho YS, Lim HS, Lee SH, Cho YL, Nam HS, Bae KS. 2018. Pharmacokinetics, Pharmacodynamics, and Tolerability of Single-Dose Oral LCB01-0371, a Novel Oxazolidinone with Broad-Spectrum Activity, in Healthy Volunteers. *Antimicrob Agents Chemother* 62.

16. Clinical and Laboratory Standards Institute. 2011. Susceptibility testing of mycobacteria, nocardiae, and other aerobic actinomycetes; approved standard, 2nd ed. CLSI document M24-A2., Wayne, PA.

17. Nessar R, Cambau E, Reytrat JM, Murray A, Gicquel B. 2012. *Mycobacterium abscessus*: a new antibiotic nightmare. *J Antimicrob Chemother* 67:810-8.

18. Adekambi T, Reynaud-Gaubert M, Greub G, Gevaudan MJ, La Scola B, Raoult D, Drancourt M. 2004. Amoebal coculture of "*Mycobacterium massiliense*" sp. nov. from the sputum of a patient with hemoptoic pneumonia. *J Clin Microbiol* 42:5493-501.

19. Koh WJ, Jeon K, Lee NY, Kim BJ, Kook YH, Lee SH, Park YK, Kim CK, Shin SJ, Huitt GA, Daley CL, Kwon OJ. 2011. Clinical significance of differentiation of *Mycobacterium massiliense* from *Mycobacterium abscessus*. *Am J Respir Crit Care Med* 183:405-10.

20. Brown-Elliott BA, Wallace RJ, Jr. 2002. Clinical and taxonomic status of pathogenic nonpigmented or late-pigmenting rapidly growing mycobacteria. *Clin Microbiol Rev* 15:716-46.

- 402 21. Shen Y, Wang X, Jin J, Wu J, Zhang X, Chen J, Zhang W. 2018. In Vitro Susceptibility of
403 Mycobacterium abscessus and Mycobacterium fortuitum Isolates to 30 Antibiotics. Biomed
404 Res Int 2018:4902941.
- 405 22. Aono A, Morimoto K, Chikamatsu K, Yamada H, Igarashi Y, Murase Y, Takaki A, Mitarai S. 2019.
406 Antimicrobial susceptibility testing of Mycobacteroides (Mycobacterium) abscessus complex,
407 Mycolicibacterium (Mycobacterium) fortuitum, and Mycobacteroides (Mycobacterium)
408 chelonae. J Infect Chemother 25:117-123.
- 409 23. Sparks R, Khatami A. 2014. Mycobacterium fortuitum Complex Skin Infection in a Healthy
410 Adolescent. Infect Disord Drug Targets 14:168-71.
- 411 24. Hashemian SMR, Farhadi T, Ganjparvar M. 2018. Linezolid: a review of its properties, function,
412 and use in critical care. Drug Des Devel Ther 12:1759-1767.
- 413 25. Cavusoglu C, Soyler I, Akinci P. 2007. Activities of Linezolid against nontuberculous
414 mycobacteria. New Microbiol 30:411-4.
- 415 26. Brown-Elliott BA, Crist CJ, Mann LB, Wilson RW, Wallace RJ, Jr. 2003. In vitro activity of
416 linezolid against slowly growing nontuberculous Mycobacteria. Antimicrob Agents Chemother
417 47:1736-8.
- 418 27. Prokocimer P, Bien P, Surber J, Mehra P, DeAnda C, Bulitta JB, Corey GR. 2011. Phase 2,
419 randomized, double-blind, dose-ranging study evaluating the safety, tolerability, population
420 pharmacokinetics, and efficacy of oral torezolid phosphate in patients with complicated skin
421 and skin structure infections. Antimicrob Agents Chemother 55:583-92.
- 422 28. Vera-Cabrera L, Brown-Elliott BA, Wallace RJ, Jr., Ocampo-Candiani J, Welsh O, Choi SH,
423 Molina-Torres CA. 2006. In vitro activities of the novel oxazolidinones DA-7867 and DA-7157
424 against rapidly and slowly growing mycobacteria. Antimicrob Agents Chemother 50:4027-9.
- 425 29. Brown-Elliott BA, Wallace RJ, Jr. 2017. In Vitro Susceptibility Testing of Tedizolid against
426 Nontuberculous Mycobacteria. J Clin Microbiol 55:1747-1754.
- 427 30. Zong Z, Jing W, Shi J, Wen S, Zhang T, Huo F, Shang Y, Liang Q, Huang H, Pang Y. 2018.
428 Comparison of in vitro activity and MIC distributions between the novel oxazolidinone
429 delpazolid and linezolid against multidrug-resistant and extensively drug-resistant
430 Mycobacterium tuberculosis in China. Antimicrob Agents Chemother
431 doi:10.1128/aac.00165-18.
- 432 31. McNeil MB, Dennison DD, Shelton CD, Parish T. 2017. In Vitro Isolation and Characterization
433 of Oxazolidinone-Resistant Mycobacterium tuberculosis. Antimicrob Agents Chemother 61.
- 434 32. Flanagan S, Minassian SL, Prokocimer P. 2018. Pharmacokinetics of Tedizolid and
435 Pseudoephedrine Administered Alone or in Combination in Healthy Volunteers. J Clin Med 7.
- 436 33. Choi Y, Lee SW, Kim A, Jang K, Nam H, Cho YL, Yu KS, Jang IJ, Chung JY. 2018. Safety,
437 tolerability and pharmacokinetics of 21 day multiple oral administration of a new
438 oxazolidinone antibiotic, LCB01-0371, in healthy male subjects. J Antimicrob Chemother
439 73:183-190.
- 440 34. Sunwoo J, Kim YK, Choi Y, Yu KS, Nam H, Cho YL, Yoon S, Chung JY. 2018. Effect of food on the
441 pharmacokinetic characteristics of a single oral dose of LCB01-0371, a novel oxazolidinone
442 antibiotic. Drug Des Devel Ther 12:1707-1714.
- 443 35. Dong W, Chochua S, McGee L, Jackson D, Klugman KP, Vidal JE. 2014. Mutations within the
444 rplD Gene of Linezolid-Nonsusceptible Streptococcus pneumoniae Strains Isolated in the
445 United States. Antimicrob Agents Chemother 58:2459-62.

36. Yang K, Chang JY, Cui Z, Li X, Meng R, Duan L, Thongchol J, Jakana J, Huwe CM, Sacchettini JC, Zhang J. 2017. Structural insights into species-specific features of the ribosome from the human pathogen *Mycobacterium tuberculosis*. *Nucleic Acids Res* 45:10884-10894.
37. Eyal Z, Matzov D, Krupkin M, Wekselman I, Paukner S, Zimmerman E, Rozenberg H, Bashan A, Yonath A. 2015. Structural insights into species-specific features of the ribosome from the pathogen *Staphylococcus aureus*. *Proc Natl Acad Sci U S A* 112:E5805-14.
38. Ippolito JA, Kanyo ZF, Wang D, Franceschi FJ, Moore PB, Steitz TA, Duffy EM. 2008. Crystal structure of the oxazolidinone antibiotic linezolid bound to the 50S ribosomal subunit. *J Med Chem* 51:3353-6.
39. Beckert P, Hillemann D, Kohl TA, Kalinowski J, Richter E, Niemann S, Feuerriegel S. 2012. *rplC* T460C identified as a dominant mutation in linezolid-resistant *Mycobacterium tuberculosis* strains. *Antimicrob Agents Chemother* 56:2743-5.
40. Pang YA-Ohoo, Zheng H, Tan Y, Song Y, Zhao Y. 2017. In Vitro Activity of Bedaquiline against Nontuberculous *Mycobacteria* in China. LID - e02627-16 [pii] LID - 10.1128/AAC.02627-16 [doi]. *Antimicrob Agents Chemother* 61.
41. Clinical and Laboratory Standards Institute.2011. Susceptibility testing of mycobacteria, nocardia, and other aerobic actinomycetes; approved standard, 2nd ed; CLSI document M24-A2, Wayne, PA.
42. Coeck N, de Jong BC, Diels M, de Rijk P, Ardizzoni E, Van Deun A, Rigouts L. 2016. Correlation of different phenotypic drug susceptibility testing methods for four fluoroquinolones in *Mycobacterium tuberculosis*. *J Antimicrob Chemother* 71:1233-40.
43. Kim SY, Jhun BW, Moon SM, Jeon K, Kwon OJ, Huh HJ, Lee NY, Shin SJ, Daley CL, Koh WJ. 2019. Genetic mutations in linezolid-resistant *Mycobacterium avium* complex and *Mycobacterium abscessus* clinical isolates. *Diagn Microbiol Infect Dis* 94:38-40.

Table 1.MICs of LZD,TZD,SZD and DZD against the reference strains of 32 RGM species

Table 2. The MICs of LZD,TZD,SZD and DZD against clinically isolated species with less than 5 isolates

Table 3.The MICs of LZD and *rplC*, *rplD* and 23srRNA mutations against *M. abscessus* and *M. massiliense* isolates

Table 4.The MICs of LZD and *rplC*, *rplD* and 23srRNA mutations against *M. fortuitum* isolates

Figure 1. The MIC distributions of *M. abscessus* and *M. massiliense* against LZD,TZD, SZD and DZD.

Figure 2.The MIC distributions of *M. fortuitum* against LZD,TZD, SZD and DZD.

Figure 3.The structure of the ribosomal 23SrRNA and *rplD*.(A)The structure of LZD and its analog LZD-114. (B)The structure of *rplD* and Ala177Pro mutations detected in *M.massiliense* isolated highlighted in red.(C) The structure of ribosomal 23SrRNA and mutations detected in tested RGM clinical isolates were highlighted in red.

484 **Table S1. Primer sets used for target genes in this study**

485 **FigureS1. Sequence alignment *rplC* homologue proteins.** Alignment of the amino acid
 486 sequences of *M. tuberculosis*, *M. abscessus*, *M. massiliense*, *M. chelonae*, *M. fortuitum* and
 487 *M. smegmatis*. The topology of the *rplC* encoded protein of *M. tuberculosis* is shown at
 488 the top. Red boxes with white letters indicate a single, fully conserved residue. Blue frames
 489 indicate highly conserved residues. β -Strands are rendered as arrows.

490 **FigureS2. Sequence alignment *rplD* homologue proteins.** Alignment of the amino acid
 491 sequences of *M. tuberculosis*, *M. abscessus*, *M. massiliense*, *M. chelonae*, *M. fortuitum* and
 492 *M. smegmatis*. The topology of the *rplD* encoded protein of *M. tuberculosis* is shown at
 493 the top. Red boxes with white letters indicate a single, fully conserved residue. Blue frames
 494 indicate highly conserved residues. β -Strands are rendered as arrows.

Table 1. MICs of LZD,TZD,SZD and DZD against reference strains of 32 RGM species

Strain by type	Mycobacterium species (strain)	MIC(µg/ml)			
		LZD	TZD	SZD	DZD
RGM species					
ATCC 19977	<i>Mycobacterium abscessus</i>	16	4	8	8
ATCC 27406	<i>Mycobacterium agri</i>	0.25	0.25	0.25	0.5
ATCC 27280	<i>Mycobacterium aichiense</i>	0.5	0.5	2	1
ATCC 23366	<i>Mycobacterium aurum</i>	0.25	0.125	0.25	0.125
ATCC 33464	<i>Mycobacterium austroafricanum</i>	0.25	0.25	2	0.5
ATCC 14472	<i>Mycobacterium chelonae</i>	8	8	4	8
ATCC 19627	<i>Mycobacterium chitae</i>	1	1	2	1
ATCC 27278	<i>Mycobacterium chubuense</i>	32	2	2	2
DSM 44829	<i>Mycobacterium cosmeticum</i>	4	2	16	1
ATCC 19340	<i>Mycobacterium diernhoferi</i>	1	0.5	2	1
ATCC 43910	<i>Mycobacterium duvalii</i>	1	0.25	0.5	0.5
ATCC 35219	<i>Mycobacterium fallax</i>	4	2	8	1
ATCC 14474	<i>Mycobacterium flavescens</i>	16	2	2	8
ATCC 6841	<i>Mycobacterium fortuitum</i>	32	>32	>32	>32
ATCC 27726	<i>Mycobacterium gadium</i>	0.5	0.125	0.25	0.125
ATCC 43909	<i>Mycobacterium gilvum</i>	0.5	0.25	0.5	0.5
ATCC BAA-955	<i>Mycobacterium goodii</i>	16	4	32	2
DSM 44124	<i>Mycobacterium mucogenicum</i>	1	1	1	1
ATCC 25795	<i>Mycobacterium neoaurum</i>	1	0.5	2	1
ATCC 27023	<i>Mycobacterium obuense</i>	0.5	0.25	0.5	0.5
ATCC 19686	<i>Mycobacterium parafortuitum</i>	1	0.5	2	1
DSM 43271	<i>Mycobacterium peregrinum</i>	2	4	4	1
ATCC 11758	<i>Mycobacterium phlei</i>	2	4	8	16
ATCC 33776	<i>Mycobacterium porcinum</i>	16	8	32	4
ATCC 35154	<i>Mycobacterium pulveris</i>	1	1	1	2
ATCC 27024	<i>Mycobacterium rhodesiae</i>	>32	>32	>32	16
ATCC 700731	<i>Mycobacterium septicum</i>	16	8	8	8
ATCC 19420	<i>Mycobacterium smegmatis</i>	2	2	4	4
ATCC 19527	<i>Mycobacterium thermoresistibile</i>	4	2	2	4
ATCC 27282	<i>Mycobacterium tokaiense</i>	1	2	2	1
ATCC 23292	<i>Mycobacterium triviale</i>	2	2	2	2
ATCC 15483	<i>Mycobacterium vaccae</i>	2	0.5	1	1

Table 2. MICs of LZD,TZD,SZD and DZD against reference strains of 6 RGM species

Mycobacterium species (strain)	Clinical isolates number	MIC(μg/ml)			
		LZD	TZD	SZD	DZD
RGM species					
Mycobacterium chelonae	585	32	8	16	8
Mycobacterium chelonae	752	>32	16	8	16
Mycobacterium chelonae	1354	4	1	2	1
Mycobacterium chelonae	1392	4	1	2	1
Mycobacterium chelonae	1593	4	0.5	2	0.5
Mycobacterium porcinum	29891	4	2	4	1

Table 3. The MICs of LZD and *rplC*, *rplD* and *23srRNA* mutations against *M. abscessus* and *M. massiliense* isolates

MIC of LZD (μg/ml)	Species (NO.)	RplC	RplD	23SrRNA
0.25	<i>M. abscessus</i> (0)	—	—	—
	<i>M. massiliense</i> (1)	Leu86Leu(1)	Gly75Gly(1)	WT
1	<i>M. abscessus</i> (1)	WT	WT	WT
	<i>M. massiliense</i> (0)	—	—	—
2	<i>M. abscessus</i> (4)	WT	WT	G1914A (1) T2650C(1)
	<i>M. massiliense</i> (4)	Leu86Leu(2)	Gly75Gly(2) Ala177Pro(2) Val192Val(2)	WT
4	<i>M. abscessus</i> (14)	WT	Phe23Phe(2)	T2650C(1)
	<i>M. massiliense</i> (8)	Leu86Leu(5)	Gly75Gly(5) Ala177Pro (3) Val192Val(3)	G2582C(1) -2625AC(1)
8	<i>M. abscessus</i> (17)	WT	Phe23Phe(3) Gly111Gly(1)	WT
	<i>M. massiliense</i> (10)	Leu86Leu(4) Ala92Ala(2)	Ile29Ile(1) Gly75Gly(3) Ala177Pro (4) Val192Val(6)	A2271G(1)
16	<i>M. abscessus</i> (13)	WT	Phe23Phe(3)	WT
	<i>M. massiliense</i> (5)	Leu86Leu(2)	Gly75Gly(2) Ala177Pro (3) Val192Val(3)	WT
>16	<i>M. abscessus</i> (0)	—	—	—
	<i>M. massiliense</i> (7)	Leu86Leu(1) Ala92Ala(6)	Gly75Gly(1) Val192Val(5)	WT

Table 4.The MICs of LZD and *rplC*, *rplD* and *23srRNA* mutations against *M. fortuitum* isolates

MIC of LZD (μg/ml)	Species (NO.)	RplC	RplD	23SrRNA
4	1	-	Ala146Gly+ Thr147Ser + Val156Ile+ Ala161Thr+ Lys167Arg+ Ser207Ala+ Glu212Gly+ Val213Ala+ Ala215Val (1)	
8	2		Ala146Gly+ Thr147Ser + Val156Ile+ Ala161Thr+ Lys167Arg+ Ser207Ala+ Glu212Gly+ Val213Ala+ Ala215Val (1)	
			WT	C1944T(1)
16	9		Ala146Gly+ Thr147Ser + Val156Ile+ Ala161Thr+ Lys167Arg+ Ser207Ala+ Glu212Gly+ Val213Ala+ Ala215Val (8)	
32	7		Ala161Thr+ Lys167Arg+ Ser207Ala+ Glu212Gly+ Val213Ala+ Ala215Val(1)	A2090T(1)
			Ala146Gly+ Thr147Ser + Val156Ile+ Ala161Thr+ Lys167Arg+ Ser207Ala+ Glu212Gly+ Val213Ala+ Ala215Val (5)	
>32	6		Ala146Gly+ Thr147Ser + Val156Ile+ Ala161Thr+ Lys167Arg+ Ser207Ala+ Glu212Gly+ Val213Ala+ Ala215Val (6)	

Figure 1

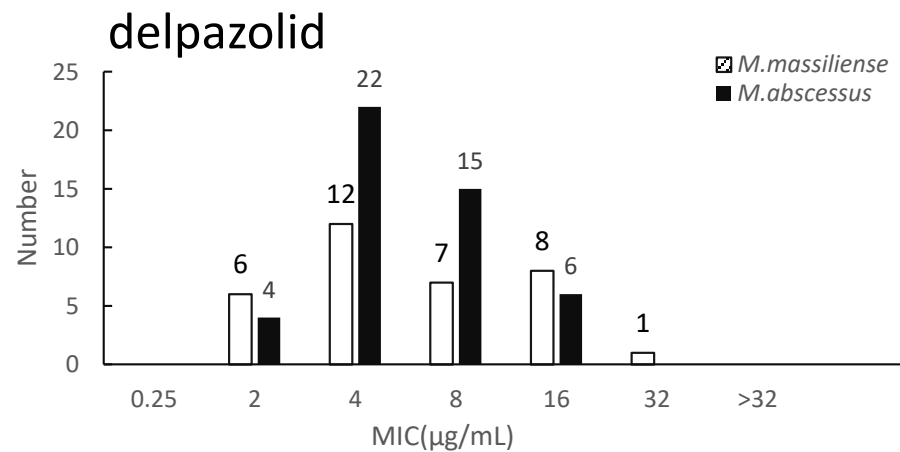
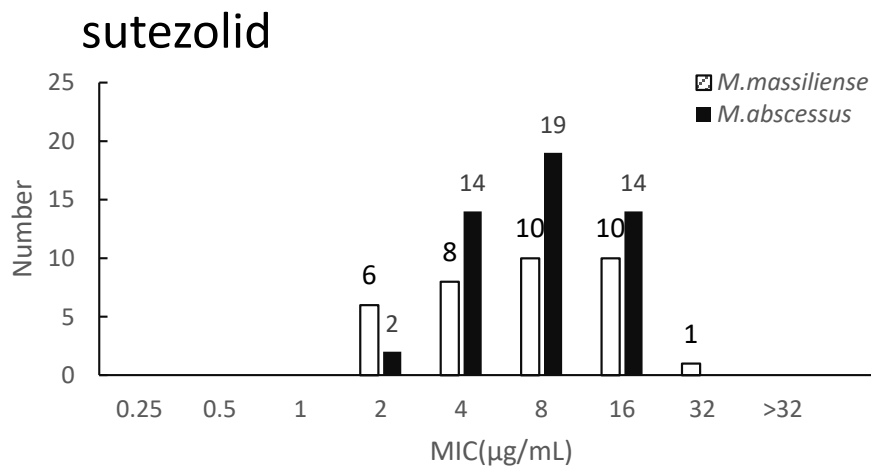
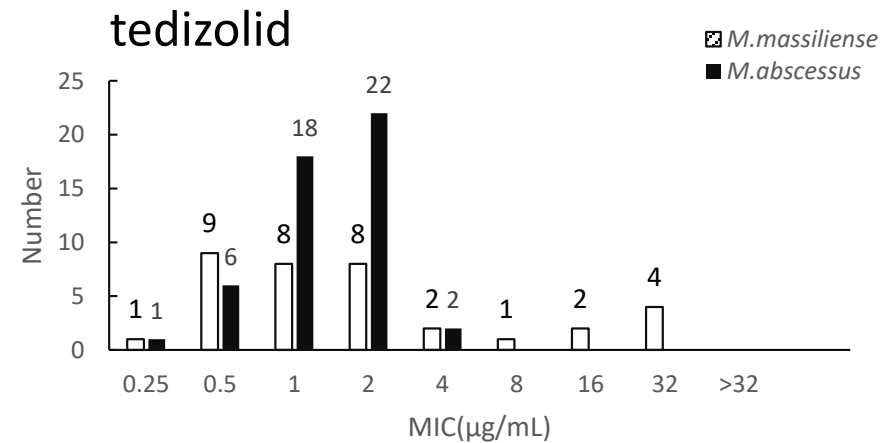
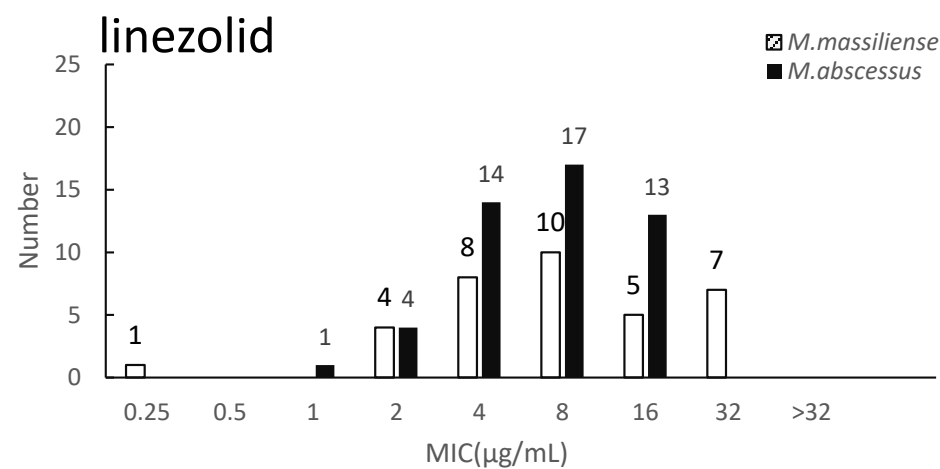


Figure 2

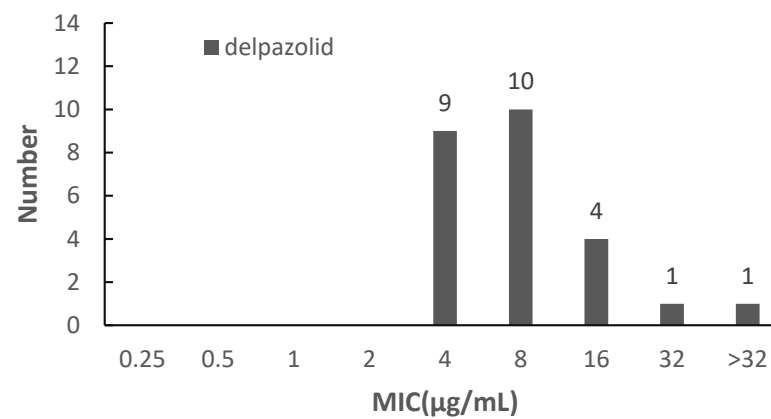
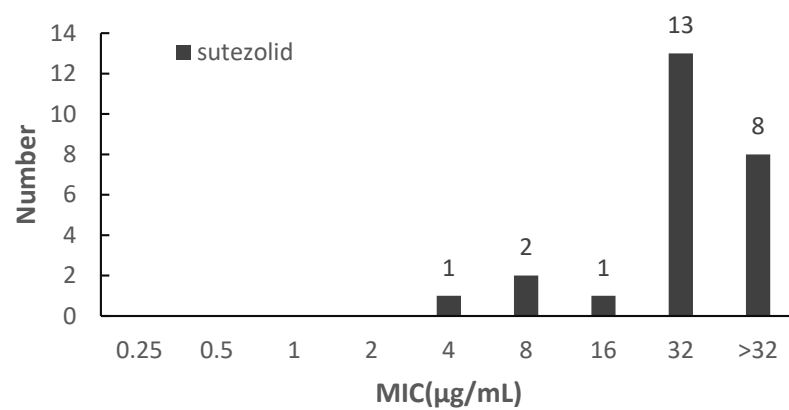
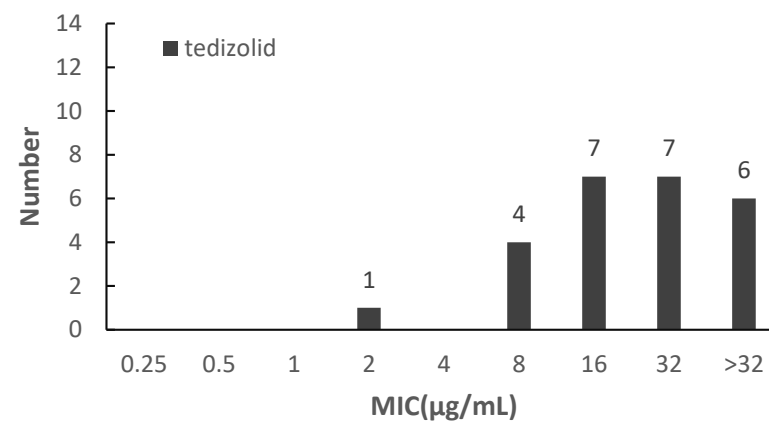
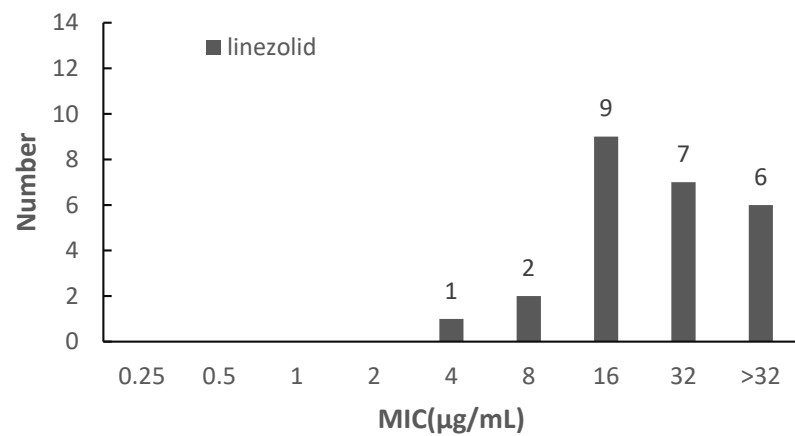
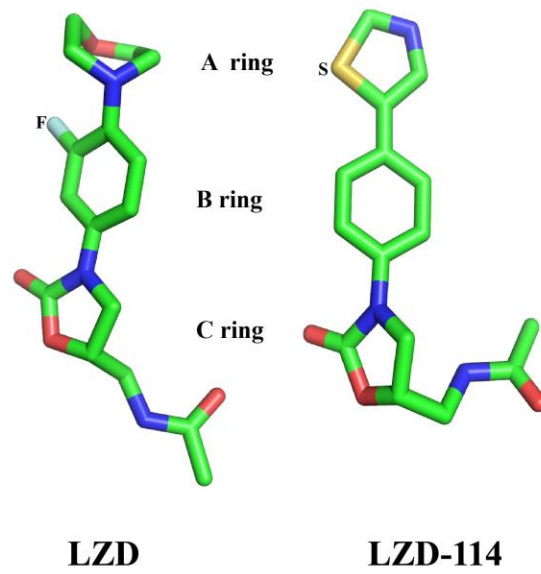


Figure.3

A



C



B

