In vitro Synergism of Six Antituberculosis Agents Against Drug-Resistant Mycobacterium tuberculosis Isolated from Retreatment Tuberculosis Patients

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Background: Retreatment tuberculosis (TB) has become a major source of drug-resistant TB. In contrast to the combination of isoniazid (INH) and rifampicin (RIF), that of pasiniazid (Pa) and rifabutin (RFB) or rifapentine (RFP) appears to have better activity in vitro against drug-resistant Mycobacterium tuberculosis (MTB), especially when combined with moxifloxacin (MXF). However, there has been limited study of potential synergism among Pa, RFB, RFP, and MXF, or simultaneous comparison with the standard INH and RIF combination.

Methods: In vitro synergism of four two-drug combinations (INH and RIF, Pa and RFB, Pa and RFP, MXF and Pa) and two three-drug combinations (MXF and Pa combined with RFB or RFP) was evaluated against 90 drug-resistant MTB strains isolated from retreatment TB patients by the checkerboard method. The fractional inhibitory concentration index (FICI) was calculated for each combination.

Results: The synergistic activity of the combination of Pa with RFB or RFP was higher than that of INH and RIF or MXF and Pa, and the synergistic activity of Pa in combination with RFP was even higher than that of RFB, although RFP yielded an MIC₉₀ of 64 mg/liter, higher than that of RFB of 8 mg/liter against 90 drug-resistant MTB strains. Meanwhile, for three-drug combinations, the synergistic effects of MXF and Pa combined with RFB or RFP were similar. Further stratification analysis showed that, for XDR-MTB strains, the synergistic effect of the Pa and RFP combination was also better than those of other two-drug combinations.

Conclusion: The combination of Pa with RFP shows better in vitro synergism than Pa with RFB and standard INH with RIF combinations, which can provide a reference for new regimens for retreatment TB patients.

Keywords: Retreatment tuberculosis, XDR-MTB, MDR-MTB, Combined drug sensitivity, FICI, MIC

Introduction

Tuberculosis (TB) is a chronic infectious disease caused by Mycobacterium tuberculosis (MTB), which requires a long course of treatment and can easily develop drug resistance.¹ Because of its serious harm to people’s health, TB is still a major public health problem throughout the world. Retreatment TB has become a major source of drug-resistant TB, which can evolve into multidrug-resistant TB (MDR-TB) or even extensively drug-resistant TB (XDR-TB) if not effectively treated.²,³ The current standard chemotherapy regimen for retreatment TB in China is the recombination and superposition of first-line anti-TB drugs, which is no longer...
suitable to deal with the high drug-resistance rates. Issues that arise from this regimen are low cure rate, aggravation of adverse reactions, long treatment course, and interruption of treatment with classic regimen, which contribute to the occurrence of MDR-TB. There is thus a need for new and more effective treatment combinations. In vitro drug susceptibility tests can accurately evaluate the inhibitory activity of different drug combinations against various drug-resistant MTB strains, thereby providing an effective basis for optimizing clinical treatment regimens.

As research on retreatment TB accelerates and more effective treatments are being sought, our previous work introduced a short-range treatment option by replacing isoniazid (INH), rifampicin (RIF), and streptomycin (SM) in the standard treatment regimen of INH, RIF, SM, ethambutol (EMB), and pyrazinamide (PZA) with pasiniazid (Pa), rifabutin (RFB), and moxifloxacin (MXF) to establish a new regimen of Pa, RFB, MXF, RFP, and RFP, was performed as described previously. In vitro activity and synergy between different antimicrobial agents against MTB have been reported. However, the synergism among Pa, RFB, RFP, and MXF against drug-resistant MTB has not been systematically analyzed. In this work, we calculated the antibacterial concentration index or fractional inhibitory concentration index (FICI) of the six core drugs in the two regimens on 90 drug-resistant MTB strains isolated from retreatment TB patients. Meanwhile, for patients unable to tolerate the side effects of RFB, rifapentine (RFP) can be used as an alternative. In vitro activity and synergy between different antimicrobial agents against MTB have been reported. However, the synergism among Pa, RFB, RFP, and MXF against drug-resistant MTB has not been systematically analyzed. In this work, we calculated the antibacterial concentration index or fractional inhibitory concentration index (FICI) of the six core drugs in the two regimens on 90 drug-resistant MTB strains isolated from retreatment TB patients in order to compare the antibacterial effects of the drug combinations of the standard and new regimens. Our results provide further support for the assertion that the combination of Pa with RFP exhibits better in vitro synergism than Pa with RFB, or INH with RIF, providing a reference for the development of new regimens for retreatment TB patients.

**Materials and Methods**

**Ethical Approval of the Study Protocol**

The study protocol was approved by the Ethics Committees of Shanghai Pulmonary Hospital of Tongji University (K19-008) in Shanghai, China. It was carried out in line with the ethical standards laid down in the 1964 Declaration of Helsinki and its later amendments. Patients gave their written informed consent to have their data included in this study.

**Mycobacterium tuberculosis Isolates**

This study included 90 drug-resistant clinical isolates obtained from retreatment TB patients, including 54 XDR-MTB, 29 MDR-MTB, 3 poly-resistant, and 4 INH-resistant isolates, at Shanghai Pulmonary Hospital affiliated to Tongji University in Shanghai, China, between January 2011 and December 2015. Drug susceptibility results of the 90 isolates were obtained by the proportion method using the BACTEC MGIT 960 system, as shown in Table S1. The H37Rv (number: ATCC27294) reference strain was also tested. These isolates were stored in 7H9 broth (Becton Dickinson, Franklin Lakes, NJ) containing 15% glycerol in a freezer at −80°C until analysis. The test isolates were grown at 37°C in Middlebrook 7H9 broth (Becton Dickson) supplemented with 10% ADC [5% bovine serum albumin (BSA), 2% dextrose, 5% catalase], and 0.05% Tween-80 (Sigma-Aldrich) to mid-log phase (OD$_{590}$ ≈ 0.4, ~2.5 × 10$^{5}$ CFU/mL) and diluted with broth medium to a final concentration of ~10$^{5}$ CFU/mL used for the drug susceptibility testing.

**Antimicrobial Agents**

INH, RIF, Pa, RFB, and RFP were purchased from Sigma Chemical Co. (St. Louis, MO); MXF was purchased from Med Chem Express (MCE). Drug configuration: All drugs were weighed to 10 mg; Pa dry powder was prepared with 1 mL of 4% NaOH to 10 mg/mL; INH dry powder was prepared with 1 mL of ultra-pure water to 10 mg/mL; RIF, RFB, and RFP dry powders were dissolved with 1 mL of dimethyl formamide to 10 mg/mL; and MXF dry powder was dissolved with 1 mL of dimethyl sulfoxide (DMSO) and prepared to 10 mg/mL. All drug solutions were filtered and sterilized with a 0.22 µm sterile filter, and stored separately in a freezer at −80°C. Further dilutions were made with coating buffer contained 30% ethanol and 0.5% sucrose.

**Drug Susceptibility Testing**

Broth microdilution minimum inhibitory concentration (MIC) testing of six anti-TB agents, namely, INH, RIF, Pa, MXF, RFB, and RFP, was performed as described previously. Serial double dilutions of the tested antimicrobial agents were prepared for each isolate using the following ranges of drug concentration: 0.125–8 mg/liter for INH, 0.03–32 mg/liter for RIF, 0.03–2 mg/liter for MXF, 0.0075–8 mg/liter for Pa, 0.03–2 mg/liter for RFB, and 0.25–16 mg/liter for RFP, as suggested by the Clinical and Laboratory Standards Institute (CLSI) with some adjustments. In brief,
if the standard highest concentration of each drug could not inhibit the growth of the tested isolates, the highest concentration would be doubled; if the tested isolates could not grow under the standard lowest concentration of each drug, the lowest concentration would be halved. The MIC test was performed in duplicate to ensure the reliability of the results.

The checkerboard titration method was used to test the combinations of INH with RIF, Pa with RFB, Pa with RFP, MXF with Pa, and MXF and Pa combined with RFB or RFP. For the combination of two agents, such as INH (drug A) with RIF (drug B), RIF was serially diluted along the abscissa, while INH was diluted along the ordinate. A1, H12 wells to a 96-well plate (rows A–H, columns 1–12) were prepared as a growth control with drug-free medium. The combination of Pa with RFB, Pa with RFP, and MXF with Pa were prepared as INH with RIF. Three-drug combination checkerboard titration methods were principally based on the standard two-drug (MXF and Pa) combination checkerboard assay. Two-drug combination checkerboard plates were prepared by dispensing the serially diluted Pa along the abscissa and MXF along the ordinate in a 96-well plate. The third drug (RFB or RFP) was then dispensed on the whole plate with one of the serial concentrations from the MIC range, except the A1 growth control well, as shown in Table 1. For the preparation of drug plates, sterile 96-well plates were added with 20 µL of drug-containing coating buffer using an automatic dispenser, air-dried, sealed, and stored at 4°C. For DST, 200 µL of mycobacterial suspension was added to each plate. After incubation at 37°C for 2 weeks, the growth status of the drug control strain (H37Rv) and growth control well (wells A1 and H12 in the dual-drug and well A1 in the triple-drug plates) of each plate was observed first to confirm that the drugs were effective and each strain grew normally, after which the MICs of all of the agents in the different combinations were observed and recorded.

The fractional inhibitory concentration index (FICI) of two drugs was calculated by the formula FICI = (MIC A + B/MIC A) + (MIC B + A/MIC B), where MIC A + B represents the MIC of compound A when combined with B; MIC B + A, the MIC of compound B when combined with A; and MIC A and MIC B, the MIC of compounds A and B tested alone, respectively. Synergy was defined as an FICI value ≤0.5, an FICI value between 0.5 and 4 was considered indifferent, and an FICI value >4 was considered to reflect antagonism.

The FICI of three drugs was calculated by the formula FICI = (MIC A + B + C/MIC A) + (MIC B +
A + C/MIC B) + (MIC C + A + B/MIC C), where MIC A + B + C represents the MIC of compound A when combined with B and C; MIC B + A + C, the MIC of compound B when combined with A and C; MIC C + A + B, the MIC of compound C when combined with A and B; and MIC A, MIC B, and MIC C, the MICs of compounds A, B, and C tested alone, respectively. An FICI value ≤0.75 was considered to indicate synergism, an FICI value between 0.75 and 4 was considered indifferent, and an FICI value >4 was considered to reflect antagonism.

**Statistical Analyses**

All experimental data were analyzed using GraphPad Prism software. The MIC$_{50}$ and MIC$_{90}$ were used to describe the centralized and discrete trends of the data, and the statistical significance of differences between the groups was determined using the Wilcoxon paired test. Synergy for counting data is expressed as a percentage (%). Chi-squared test or Fisher’s exact test was used to analyze the distribution of FICI values. A P value <0.05 was considered statistically significant.

**Results**

**Dual-Drug Susceptibility Testing of the Combinations of INH and RIF, Pa and RFB, Pa and RFP, as Well as MXF and Pa**

The single- and dual-drug susceptibility test results for the six antimicrobial agents and four combinations against 90 drug-resistant MTB clinical isolates from retreatment TB patients are shown in Table 2 and Tables S2 and S3. Except for the case of MXF, the MIC$_{50}$ and MIC$_{90}$ of INH, RIF, Pa, RFB, and RFP were all decreased after the combination of the other drug ($P < 0.001$), which suggests that the combinations of INH and RIF, Pa and RFB, and Pa and RFP exerted synergistic effects with each other’s antimicrobial activity. After Pa was combined with MXF, the MIC$_{50}$ and MIC$_{90}$ of Pa were reduced from 8 to 0.0075 mg/liter and 16 to 4 mg/liter ($P < 0.001$), respectively. However, the MIC$_{50}$ and MIC$_{90}$ of MXF did not decrease after it was combined with Pa, indicating that MXF has a significant unilateral synergistic effect on Pa.

The synergistic, indifferent, or antagonistic effects of each drug combination on the clinical isolates are shown in Table 3 and compared as described in reference 12. The combinations of Pa with RFP and Pa with RFB were more likely to show synergy than that of INH with RIF ($P = 0.0127$ and $P < 0.0001$, respectively), and MXF in combination with Pa ($P = 0.0004$ and $P < 0.0001$, respectively). Pa in combination with RFP was more likely to show synergy than the combination of Pa with RFB ($P = 0.0021$), although RFP yielded an MIC$_{90}$ of 64 mg/liter, higher than that of RFB of 8 mg/liter.

**Triple-Drug Susceptibility Testing (Based on the Combination of MXF and Pa)**

The dual-drug susceptibility testing provided stronger statistical evidence for synergistic effects for the

| Drug Combination | MIC (Mg/Liter) | P value |
|------------------|----------------|---------|
|                  | 50%            | 90%     | Range |
| INH+RIF          |                |         |       |
| INH              | 4              | 16      | 0.125 to 16 <0.001 |
| inh              | 2              | 8       | 0.125 to 16 <0.001 |
| RIF              | 24             | 64      | 0.03 to 64 <0.001 |
| rif              | 1              | 32      | 0.03 to 64 <0.001 |
| Pa+RFB           |                |         |       |
| Pa               | 2              | 16      | 0.015 to 32 <0.001 |
| pa               | 0.5            | 4       | 0.0075 to 16 <0.001 |
| RFB              | 4              | 8       | 0.0625 to 8 <0.001 |
| rfb              | 0.125          | 2       | 0.03125 to 4 <0.001 |
| Pa+RFP           |                |         |       |
| Pa               | 2              | 16      | 0.015 to 32 <0.001 |
| pa               | 0.25           | 4       | 0.0075 to 8 <0.001 |
| RFP              | 32             | 64      | 0.5 to 64 <0.001 |
| rfp              | 4              | 16      | 0.25 to 16 <0.001 |
| MXF+Pa           |                |         |       |
| MXF              | 0.5            | 1       | 0.03 to 4 0.012 |
| mxf              | 0.5            | 2       | 0.03 to 2 0.012 |
| Pa               | 8              | 16      | 0.015 to 32 <0.001 |
| pa               | 0.0075         | 4       | 0.0075 to 16 <0.001 |

Notes: INH, MIC of INH tested alone; inh, MIC of INH tested with RIF in combination; RIF, MIC of RIF tested alone; rif, MIC of RIF tested with INH in combination. The rest of the upper- and lower-case letters are the same (dual-drug susceptibility testing); data in bold indicate that the P values were statistically significant.

**Table 3** Distribution and Comparison of FICI of 90 Strains from Four Dual-Drug Combinations

| FICI Value | INH+RIF | Pa+RFB | Pa+RFP | MXF+Pa |
|------------|---------|--------|--------|--------|
| ≤0.5       | 0       | 0      | 0      | 0      |
| 0.5< and ≤4| 1       | 1      | 1      | 1      |
| >4         | 0       | 0      | 0      | 0      |

Notes: P value for a vs b comparison was 0.0127; P value for a vs c comparison was <0.0001; P value for a vs d comparison was 0.2807; P value for b vs c comparison was 0.0021; P value for b vs d comparison was 0.0004; P value for c vs d comparison was <0.0001; e, the data in and outside the brackets represented the percentages and frequencies.
combinations of Pa with RFP and Pa with RFB, while MXF displayed a significant unilateral synergistic effect on Pa. We thus first tested the individual MICs of RFB and RFP in the three-drug combinations (Table S3). The combination of RFB with MXF and Pa decreased the MIC$_{90}$ of RFB from 8 to 0.2375 mg/liter ($P < 0.001$), and the combination of RFP with MXF and Pa decreased the MIC$_{90}$ of RFP from 64 to 1 mg/liter ($P < 0.001$), as shown in Table 4. The MIC of Pa was also significantly decreased after combination with RFB and MXF, or RFP and MXF ($P < 0.001$). However, the synergistic rates of Pa and RFB or Pa and RFP after combination with MXF were decreased from 28.9% and 50% to 17.8% and 23.3%, respectively, and the difference between two three-drug combinations was not as significant as for the two-drug combinations ($P = 0.5319$), as shown in Table 5.

### Stratification Analysis

To evaluate the synergistic effects of different combinations on MTB strains with various drug resistance, further stratification analysis was performed against 54 strains of extensively drug-resistant MTB (XDR-MTB) and 29 strains of multidrug-resistant MTB (MDR-MTB).

Of the 54 XDR-MTB strains tested, the combination of Pa with RFP was more likely to show synergy than those of INH and RIF, MXF and Pa, and Pa and RFB ($P < 0.0001, P < 0.0001, P = 0.0004$, respectively). The combination of Pa with RFB was more likely to show synergy than that of MXF and Pa ($P = 0.0001$), but did not differ from that of INH in combination with RIF ($P = 0.4658$). The levels of synergism between two three-drug combinations, MXF and Pa combined with RFB or RFP, were similar ($P = 0.9646$), as shown in Tables 6 and 7.

Of the 29 MDR-MTB strains, the combinations of Pa with RFP and Pa with RFB were also more likely to show synergy than the combination of MXF and Pa ($P = 0.015$ and $P = 0.0074$, respectively), but did not differ from INH in combination with RIF ($P = 0.2079$ and $P = 0.125$, respectively). The levels of synergism between the two-drug combinations of Pa with RFP and Pa with RFB, or the three-drug combinations of MXF and Pa with RFB or RFP, were similar ($P > 0.99$ and $P = 0.3847$, respectively), as shown in Tables 8 and 9.

### Discussion

In this study, 90 strains of drug-resistant MTB isolated from sputum specimens of retreatment TB patients were tested for combined drug susceptibility to accurately evaluate the synergistic effects in vitro of the core drugs from the standard and new regimens. Via the comparison of MIC or FICI from drugs in different combinations, we want to provide some

**Table 4** Comparison of MICs for Each Drug Before and After Combinations from Triple-Drug Susceptibility Testing

| Drug Combination       | MIC (Mg/Liter) | P value |
|------------------------|----------------|---------|
|                        | 50%            | 90%     | Range     |
| MXF+Pa+RFB             |                |         |           |
| MXF                    | 0.5            | 1       | 0.03 to 4 | 0.129     |
| mxf                    | 0.25           | 2       | 0.03 to 2 |           |
| Pa                     | 2              | 16      | 0.015 to 32 | <0.001 |
| pa                     | 0.0075         | 0.5     | 0.0075 to 16 | <0.001 |
| RFB                    | 4              | 8       | 0.0625 to 8 | <0.001   |
| rfb                    | 0.125          | 0.2375  | 0.03125 to 1 |           |
| MXF+Pa+RFP             |                |         |           |
| MXF                    | 0.5            | 1       | 0.03 to 4 | 0.719     |
| mxf                    | 0.25           | 2       | 0.03 to 2 |           |
| Pa                     | 2              | 16      | 0.015 to 32 | <0.001 |
| pa                     | 0.0225         | 4       | 0.0075 to 16 | <0.001 |
| RFP                    | 32             | 64      | 0.5 to 64 |           |
| rfp                    | 1              | 1       | 0.5 to 4  | <0.001   |

**Notes:** MXF, MIC of MXF tested alone; mxf, MIC of MXF tested with the combination of Pa with RFB; Pa, MIC of Pa tested alone; pa, MIC of Pa tested with the combination of MXF with RFB; RFB, MIC of RFB tested alone; rfb, MIC of RFB tested with the combination of MXF with Pa. The rest of the upper- and lower-case letters are the same (triple-drug susceptibility testing); data in bold indicate that the P values were statistically significant.

**Table 5** Distribution and Comparison of FICI of 90 Strains from Two Triple-Drug Combinations

| FICI Value | MXF+Pa+RFB | MXF+Pa+RFP |
|------------|------------|------------|
| ≤0.75      | 16 (17.8)  | 21 (23.3)  |
| 0.75< and ≤4 | 64 (71.1) | 57 (63.3)  |
| >4         | 10 (11.1)  | 12 (13.3)  |

**Notes:** P value for a vs b comparison was 0.5319; c, the data in and outside the brackets represented the percentages and frequencies.

**Table 6** Distribution and Comparison of FICI of 54 XDR-MTB Strains from Four Dual-Drug Combinations

| FICI Value | INH+RIF | Pa+RFB | Pa+RFP | MXF+Pa |
|------------|---------|--------|--------|--------|
| ≤0.5       | 6 (11.1) | 11 (20.4) | 28 (51.9) | 3 (5.6) |
| 0.5< and ≤4 | 48 (88.9) | 41 (75.9) | 26 (48.1) | 51 (94.4) |
| >4         | 0        | 2 (3.7)  | 0      | 0      |

**Notes:** P value for a vs b comparison was 0.4658; P value for a vs c comparison was <0.0001; P value for a vs d comparison was 0.4862; P value for b vs c comparison was 0.0004; P value for b vs d comparison was 0.0001; P value for c vs d comparison was <0.0001; e, the data in and outside the brackets represented the percentages and frequencies.
Table 7 Distribution and Comparison of FICI of 54 XDR-MTB Strains from Two Triple-Drug Combinations

| FICI Value | MXF+Pa+RFB* | MXF+Pa+RFPb |
|------------|-------------|-------------|
| ≤0.75      | 8 (14.8)c   | 9 (16.7)    |
| 0.75< and ≤4 | 38 (70.4)  | 37 (68.5)   |
| >4         | 8 (14.8)    | 8 (14.8)    |

Notes: P value for a vs b comparison was 0.9646; c, the data in and outside the brackets represented the percentages and frequencies.

Table 8 Distribution and Comparison of FICI of 29 MDR-MTB Strains from Four Dual-Drug Combinations

| FICI Value | INH+RIFa | Pa+RFBb | Pa+RFPc | MXF+Pa d |
|------------|----------|---------|---------|----------|
| ≤0.5       | 4 (13.8)c| 9 (31.0)| 10 (34.5)| 1 (3.4) |
| 0.5< and ≤4 | 25 (86.2)| 20 (69.0)| 19 (65.5)| 28 (96.6) |
| >4         | 0        | 0       | 0       | 0        |

Notes: P value for a vs b comparison was 0.2079; P value for a vs c comparison was 0.125; P value for a vs d comparison was 0.1417; P value for b vs c comparison was >0.99; P value for b vs d comparison was 0.015; P value for c vs d comparison was 0.0076; c, the data in and outside the brackets represented the percentages and frequencies.

Table 9 Distribution and Comparison of FICI of 29 MDR-MTB Strains from Two Triple-Drug Combinations

| FICI Value | MXF+Pa+RFB* | MXF+Pa+RFPb |
|------------|-------------|-------------|
| ≤0.75      | 7 (24.1)c   | 10 (34.5)   |
| 0.75< and ≤4 | 20 (69.0)  | 15 (51.7)   |
| >4         | 2 (6.9)     | 4 (13.8)    |

Notes: P value for a vs b comparison was 0.3847; c, the data in and outside the brackets represented the percentages and frequencies.

Discussion and Conclusion

Strains from Two Triple-Drug Combinations

Table 9 shows that when two or more drugs are delivered synergistically to drug-resistant MTB, the MIC of the drug-resistant MTB on the MIC of the drug-resistant MTB of MXF and Pa combined with RFP was also better than those of MXF and Pa combined with RFP, although the difference was not significant (P = 0.5319). Based on our results, we recommend that the combination of MXF and Pa with RFP be used for retreatment TB, instead of the combination of MXF and Pa with RFP. These results further emphasize the importance of analyzing the synergistic effects of drugs in the regimen.

Table 7 shows the distribution and comparison of FICI of 54 XDR-MTB strains from two triple-drug combinations. The synergistic effect of the combination of MXF with Pa is higher than that of the combination of MXF with Pa and RFP. The combination of MXF with Pa and RFP displayed better antibacterial activity than the INH and RIF combination and that a third drug, especially a rifampin derivative, must be introduced.

Table 8 shows the distribution and comparison of FICI of 29 MDR-MTB strains from four dual-drug combinations. The combination of MXF with Pa is characterized by mild adverse reactions, high bioavailability, strong tissue permeability, long elimination half-life, and no cross-resistance with other antibacterial drugs, having been broadly used for drug-resistant TB. Pa is a composite preparation, which is a compound composed of INH and p-aminosalicylic acid; many MTB isolates resistant to INH or p-aminosalicylic acid are still susceptible to Pa. Both RFB and RFP are derivatives of RIF, and also show cross-resistance to RIF in vitro. However, studies have shown that RFB and RFP still retain certain bactericidal activity against MTB strains with low resistance to RIF, and RFB especially has a more significant effect.

Multidrug combination chemotherapy with fluoroquinolones is commonly used for MDR-TB or XDR-TB. In contrast to the synergistic activity of clofazimine with MXF or capreomycin against 30 MTB strains, we examined the in vitro activity of MXF combined with Pa, RFB and Pa, or RFP and Pa against 90 drug-resistant MTB isolates. After MXF and Pa were combined, the MIC value of MXF did not change markedly, but the MIC0 and MIC50 of Pa decreased from 8 to 0.0075 and 16 to 4 (P < 0.0001), respectively, indicating that MXF may be synergistic with the bacteriostatic effect of Pa. However, the synergy rate (5.6%) for the combination of MXF with Pa was similar to that of INH and RIF (11.1%), as shown in Table 3. This suggests that the combination of MXF with Pa could not achieve a better antibacterial effect than the INH and RIF combination and that a third drug, especially a rifampin derivative, must be introduced.
Stratified analysis further suggested the effectiveness of the combination of Pa with RFP for MTB with different drug resistance profiles. Interestingly, we found no antagonism for the combinations of INH and RIF, MXF and Pa, and Pa and RFP to all of the strains, while for Pa in combination with RFB, antagonism was seen only for two XDR-MTB strains. However, the antagonistic rates of three-drug combinations were significantly increased for XDR-MTB and MDR-MTB strains. We hypothesized that the number of drugs and the type of strain in the regimen are important, and that the number of superimposed drugs and different resistant strains might lead to antagonism between drugs in chemotherapy, which also reminds us of the need to consider the combinatorial effects of drugs before formulating a new drug combination regimen.

**Conclusion**

This study confirmed that the synergistic effects of the combinations of Pa with RFB and Pa with RFP in the new regimen for retreatment TB patients were better than those of INH and RIF or MXF and Pa against 90 drug-resistant MTB strains. The bacteriostatic effect of RFB was better than that of RFP, but the synergistic effect of the combination of Pa and RFP was better than that of Pa and RFB, which can provide a clue in vitro for the design of new regimens for retreatment TB patients. In vitro interactions between core antibacterial agents from the regimen should be detected to ensure the synergistic effects of the drug combinations.

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**Disclosure**

The authors report no conflicts of interest in this work.

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