Optimized duration of antituberculosis treatment for managing female genital tuberculosis: real-world experience from a high prevalence region in Eastern China

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Abstract

**Background** Duration of antituberculosis therapy (ATT) for managing female genital tuberculosis (FGTB) is controversial with the intermittent regimen no more advocated. We therefore conducted a prospective, real-world research to compare 6 months and 9 months of ATT.

**Methods** Between 2012 and 2018, 109 drug-susceptible patients newly diagnosed with FGTB and/or tuberculous peritonitis (genital, 13; peritoneal, 34; mixed, 62) received naïve treatment for 9-12 months and further 18-month follow-up. Data on disease features at baseline and long-term outcome (intent-to-treat) were compared between group A (aged 18-35 yr) and group B (aged 36-81 yr). Efficacy and side effects of treatment were compared within each group 6 months and 9 months from ATT initiation (per-protocol), respectively.

**Results** In contrast to group B at baseline, group A had more clinical evidence predicting active tuberculosis ($P < 0.05$), severer performance of genital lesions and pelvic adhesions ($P < 0.05$), more signs of active pulmonary tuberculosis ($P < 0.01$), and less performance of only TBP ($P < 0.01$). Intent-to-treat analysis showed higher incidence of overall single side effects and poor compliance in group B ($P < 0.05$), and similar recurrence rate between 2 groups. Per-protocol analysis showed increased complete response rate ($P < 0.01$) and similar incidence of side effects ($P > 0.05$) in group B, similar complete response rate ($P > 0.05$) and increased incidence of overall single side effects ($P < 0.05$) in group B at 9-month duration.

**Conclusions** Younger females with FGTB had a greater risk of systemic infection of tuberculosis compared to older ones. Nine-month ATT using daily therapy proved to be beneficial for younger patients at reproductive age. Six-month option was suitable for older patients for reducing side effects and poor compliance in the duration of treatment.

Background

Female genital tuberculosis (FGTB), an important type of extra-pulmonary tuberculosis (EPTB), is caused by the infection of mycobacterium tuberculosis (MTB) mainly disseminated from pulmonary tuberculosis (PTB) [1]. With continuous prevalence of TB and increased immigrants [2], FGTB causes significant morbidity, atypical clinical symptoms and multiple sequelae in infected females [1, 3-5].

Six-month antituberculosis therapy (ATT) using directly observed therapy (DOT) is the globally accepted treatment for drug-susceptible tuberculosis (DS-TB) [2, 6]. However, the duration of managing FGTB or abdominal TB continues to be controversial especially when the intermittent or shorter-course regimen, showing a high risk of treatment failure and disease relapse [7-10], is no more advocated by World Health Organization (2017 update guidelines for treatment of DS-TB) [6]. Despite 6-month ATT showing objective efficacy for managing FGTB or abdominal TB in some randomized controlled trials excluding older patients or patients with comorbidity of other types of TB [11, 12], there is lack of data especially real-world data supporting its efficacy. It brings a new debate, how to make a simplified, optimized...
strategy of ATT, for managing FGTB in patients at different age categories especially those with comorbidity of other types of TB and/or immune-deficient diseases.

We therefore conducted an open, real-world study to evaluate its efficacy using daily therapy in a large group of consecutive treatment-naive patients with FGTB. A secondary objective was to compare the disease expression, side effects and patients’ compliance among patients at different age categories.

**Methods**

**Patients**

All eligible females (aged $\geq 16$ yr) between 2012 and 2018, who were newly diagnosed as either FGTB or tuberculous peritonitis (TBP) in the region, were included in the study. In the specific period, 127 consecutive females (aged 18-81 yr) were diagnosed in our center with the disease. Among them, 7 patients unclearly diagnosed after re-evaluation, 9 patients suspected with DR-TB and additional 2 elderly patients with comorbidity of malignancy, were ineligible for receiving standard treatment and excluded. The remaining 109 patients were divided into group A (younger females at reproductive age, aged 18-35 yr) and group B (older females at reproductive age and postmenopausal period, aged 36-81 yr) (Figure 1).

**Clinical data collection**

Collected clinical data included epidemiological and clinical characteristics, hematological and biochemical tests, and Tuberculin Skin Test (TST) or T-SPOT.TB (T-SPOT.TB replaced TST from 2015). All patients took chest x-ray and 95 underwent chest computerized tomography (CT) scan during hospitalization. Patients with comorbidity of PTB, either active or non-active, were carefully evaluated with imaging data stored in our electronic image system whenever needed. Forty-two patients underwent laparotomy, laparoscopy or hysteroscopy. Surgical and histological findings were carefully recorded. Ultrasound-guided peritoneocentesis was performed in 59 of 86 patients with performance of ascites. Data of ascitic tests, including routine and biochemical tests, adenosine deaminase (ADA), acid-fast bacilli (AFB) on smear or culture, were also recorded. Positive definition of TST followed National Guidelines of Tuberculosis Diagnosis (WS 288-2017, China). Operative procedures and positive definition of T-SPOT.TB followed instructions of T-SPOT.TB assay kit (Oxford Immunotec, Abingdon, UK). According to relevant reports to T-SPOT.TB [13, 14] and its efficacy for interpreting active TB in the region, a response was considered stronger (moderate-to-strong) positive in this study if the number of spots per test well was $\geq 16$ (when the background control count was $< 5$), or at least triple the value found in the background control wells (when the background control count was $\geq 5$).

**Diagnostic criteria for FGTB and TBP**

A “definite” diagnosis of FGTB was considered in the presence of $\geq 2$ of the following: (i) epidemiological, clinical, and imaging or surgical evidence of FGTB involvement; (ii) positive result of AFB on smear/culture; (iii) stronger positive TST/T-SPOT.TB, (iv) histopathological presence of TB granulomas.
Presumptive diagnosis of FGTB was made if there was strong clinical suspicion, such as clinical or explorative features, genital lesions (tubo-ovarain mass) and/or PTB on imaging, and clinically confirmed when the diagnostic ATT proved to be effective.

A “definite” diagnosis of TBP was based on the presence of high-protein (>2.5 g/dL) ascites containing >250 white blood cells/mm$^3$ along with ≥2 of the following: (i) imaging evidence of peritoneal inflammation; (ii) ascetic ADA levels>35 u/L (iii) stronger positive TST/T-SPOTTB; (iv) positive AFB on smear/culture or presence of TB granulomas in the peritoneal biopsies. Presumptive diagnosis of TBP was similar to FGTB (except genital lesions).

**ATT protocol and management of side effects**

Intensive phase included 4 drugs (isoniazid 300 mg, rifampicin 450mg, pyrazinamide 1250 mg, and ethambutol 750mg) daily for 2 months. Patients weighing >50 kg received additional doses of rifampicin, pyrazinamide and ethambutol (150 mg, 250 mg and 250 mg, respectively). The continuation phase included isoniazid and rifampicin for 7-10 months according to individual response (9-12 months’ duration widely taken in EPTB management in the specific period). Drug-induced side effects were all recorded and managed in comprehensive measures relevant to individual performance.

**Follow-up and compliance to treatment**

Patients were registered and followed up at regular intervals at TB clinics of our center. DOT was administered by patients’ family members (trained by professional physicians) in the region. Whole blood tests, liver function tests, chest x-ray or chest CT, and abdominal ultrasonic scan/CT were done per month in the intensive phase and every 2-3 months in the duration phase. Patients who reduced doses of drug-intake for side effects received symptomatic treatment after careful counseling. Patients who took irregular protocol (taking drugs < 80% of intended days, or stopped drug-intake under no guidance) or lost to follow-up were defined as poorly compliant patients and excluded on intent-to-treat analysis.

**Assessment at two observation time-points**

Based on the aim of the study, the efficacy of treatment was assessed in 109 patients at 6 months and 9 months from the initiation of ATT respectively (per-protocol analysis). All patients underwent clinical evaluation and abdominal-pelvic imaging to demonstrate healing of lesions. Chest CT scan and AFB smear/culture of sputum were repeated in those who presented with PTB and regarded as an important measure to demonstrate the response to treatment. Twenty-three patients were persuaded to repeat hysteroscopy and/or endometrial biopsy. Surgical and histopathological findings were recorded for assessment.

**Outcome measures**

“Complete clinical response” was defined as complete resolution of symptoms and signs, normalization of biochemical and hematological tests, and healing of TB lesions on imaging or repeat surgical
procedures in this study. A partial response was defined as resolution of clinical manifestations and partial healing of TB lesions. Non-response was defined persistent clinical symptoms, persistence of TB granuloma on histopathology or AFB on microscopy/culture, persistence of TB lesions on imaging or repeat surgical procedures. Healing of PTB referred to healing of active PTB lesions (miliary tubercles, cavity, infiltrated lesions and tuberculous pleural effusion) along with no progression of non-active PTB.

Recurrence during 18-month follow-up

Eighty-eight patients persisted for 9-month treatment. Patients with clinical response were followed up 6-monthly for 18 months at TB clinics of our center, and those with no response every 1-3 months. Those who failed to visit the clinics were contacted telephonically and interviewed for recurrence. To compare the disparity of recurrence rate between 2 groups on intent-to-treat analysis, patients who lost to follow-up were defined as individuals with no recurrence.

Statistical analysis

Statistic analysis was conducted using SPSS version 22.0 (IBM, Armonk, New York, USA). Parametric results are expressed as mean (range) where appropriate. Non parametric category of variables was analyzed by χ² test, χ² with Yate's correction or Fisher's Exact test, where applicable. Two-sided P < 0.05 was considered as statistically significant.

Results

Characteristics of the patients at baseline

The main epidemiological, clinical and laboratory characteristics of 109 patients at baseline are shown in Table 1. Recent pregnancy or reproduction (31.7%:2.9%, P < 0.001), infertility (36.6%:11.8%, P=0.002), ascitic ADA>35 u/L (88.2%:47.6%, P = 0.010), and stronger positive TST (66.7%:23.3%, P = 0.022) or T.SPOTTB (96.4%:64.9%, P = 0.006) were all at higher incidence in Group A, whereas Group B had more presentation with immune-deficient diseases (16.2%:2.4%, P = 0.057) and obvious ascites (60.3%:22.0%, P < 0.001). There was no difference in others of the above parameters between 2 groups at baseline. 17 (41.5%) patients of group A and 27 (39.7%) of group B underwent surgical exploration respectively. Typical performance of FGTB, including thickening-stiff or tortuous tubes (76.5%:33.3%, P = 0.013), hydrosalpinx or pyosalpinx (29.4%:3.7%, P = 0.049), and pelvic adhesions (82.4%:44.4%, P = 0.030) were more seen in Group A, but no difference seen in other surgical or histopathological parameters (Table 2).

Predominant TB sites of disease and comorbidity of other TB sites

FGTB or TBP was definitively diagnosed in 38(92.7%) patients of group A and 60 (88.2%) of group B, respectively. The remaining 11 patients were presumptively diagnosed and clinically confirmed. Of 41 patients in group A, 6 (14.6%) had FGTB, 5 (12.2%) had TBP, and 30 (73.2%) had features of both. Of 68 patients in group B, 7 (10.3%) had FGTB, 29 (42.6%) had TBP, and 32 (47.1%) had both. Simple
performance of TBP was at higher incidence in Group B in contrast to group A (42.6%:12.2%, \( P = 0.004 \)). Both groups showed similar incidence of non-active PTB (26.8%:42.6%, \( P > 0.05 \)) and other EPTB (19.5%:25.0%, \( P > 0.05 \)), but active PTB was more seen in group A (43.9%:14.7%, \( P = 0.009 \)) (Table 3).

**Intent-to-treat analysis**

Twenty-one patients were lost to follow-up or took irregular protocol in 9-month duration after initiation of treatment. Of 12 patients (11 in group B) who took irregular protocol (almost all declined further consultation at TB clinics), 9 stopped drug-intake under no guidance and 3 took drugs < 80% of intended days. Of 9 patients lost to follow-up, 7 were from group B. Intent-to-treat analysis showed that Group B had higher incidence of poor compliance (26.5%:7.3%, \( P = 0.027 \)), and higher incidence of DILI (20.6%:4.9%, \( P = 0.049 \)) and overall single side effects (42.6%:17.1%, \( P = 0.006 \)) than group A. In addition, group B also showed higher incidence of anorexia (26.5%:12.2%), vomiting (14.7%:7.3%) and leucopenia/thrombocytopenia (7.4%:2.4%) than group A, but there was no significant difference in statistic analysis (\( P>0.05 \)).

**Per-protocol analysis**

Excluding 21 poorly compliant patients, the remaining 88 patients who persisted for \( \geq 9 \) months’ ATT were taken on per-protocol analysis (Table 5). Increased complete response was achieved in group A at 9-month time-point compared to their performance at 6-month one (92.1%:63.2%, \( P = 0.006 \)), whereas no disparity was shown in group B (88.0%:87.7%, \( P > 0.05 \)).

Equal side effects, either single or overall, were shown in Group A between two time-points (10.5%:15.8%, \( P > 0.05 \)), whereas incidence of anorexia (26.0%:8.8%, \( P = 0.016 \)), DILI (22.0%:7.0%, \( P = 0.033 \)) and overall single side effects (42.0%:22.8%, \( P = 0.033 \)), significantly elevated in group B at 9-month time-point compared to their performance at 6-month one. Despite no disparity, either single or overall side effects, was shown between 2 groups at the 6-mo time-point (\( P > 0.05 \)), group B had higher incidence of overall single side effects than group A at 9-mo time-point (42.0%:15.8%, \( P = 0.008 \)). None died during the period of treatment.

**18-month follow-up for recurrence**

During 18-month follow-up after 9-month duration, 7 patients (2 from group A and 5 from group B) were lost to follow up; 4 patients (1 from group A and 3 from group B) had recurrence of disease after standard treatment. Recurrence also occurred in additional 3 patients from group B who took irregular protocol within 6-mo duration. Of 6 patients with TB recurrence from group B, 3 had comorbidity of diabetes and lost control of blood glucose around the recurrence period; 2 had decompensated liver cirrhosis; 1 had sicca syndrome and long-term use of glucocorticoids. They were all considered as DS-TB, retreated with DOT category protocol for 6-9 months along with moderate doses of drugs and symptomatic treatment, and all responded.
The patient (aged 32 years at enrollment) with recurrence was diagnosed with DR-TB after careful re-evaluation, retreated with DOT category protocol for >18 months, received bilateral salpingectomy (denied by her family members last time) after completing 6-month intensive phase, and responded (in the continuation phase before the study completed). No recurrence was observed in the rest of the patients (except those lost to follow-up) at the end of 18-month follow-up.

Discussion

The present open real-world observational study assessed the natural history of FGTB in a cohort of consecutive drug-susceptible patients followed in a region of Eastern China for a long period. As FGTB usually co-occurs with TBP, eligible females newly diagnosed with FGTB or TBP in the region were all recruited for better analyzing the composition of disease. 41 (37.6%) of 109 patients were younger females at reproductive age. Of 68 older patients, 25 (36.8%) were older females at reproductive age (aged 36-49 yr) and 43 (63.2%) were postmenopausal females (aged ≥ 50 yr). Both of them were included in group B for no obvious discrepancy (disease expression or efficacy of treatment or side effects) found. Special emphasis was given in the association between the severity of disease and age composition of patients, and more specifically on the link between duration of daily ATT and outcome of patients.

Three major points are raised in this study: (i) disease expression and compliance to treatment are mainly dependent of age; (ii) 6-month ATT brings more benefits for older patients (similar response rate and fewer side effects); (iii) 9-month ATT using daily therapy proves to be a better option for younger patients at reproductive age (similar side effects but increased complete response rate).

The mean age of developing FGTB is 40 yr in developed nations [3, 5], whereas the disease presents in younger age group (20-30 yr) in Asia due to early marriage and child-bearing in them [1, 4, 15, 16]. This study confirms that FGTB can affect patients at any age with a peak incidence and severe disease presentation in younger females at reproductive age (18-35 yr). Of 75 females diagnosed with FGTB (or mixed with TBP) in the study, 36 (48.0%) were younger females (aged 18-34 yr) and 13 (36.1%) of them had pregnancy or reproduction within 1 year before onset of disease, confirming that FGTB more likely occurs in younger females who had increased blood supply in genital tract and/or damaged genital tract barrier due to invasive procedures [1, 16]. Interestingly, despite FGTB possibly omitted in some patients especially older ones (29 of 68 presented with only TBP) who declined surgical exploration, both composition and expression of disease seemingly turned milder with ascending age, indicating a closed link between haematogenous transmission of MTB and age of females. It was also verified in the following features of younger female group: (i) higher rate of stronger positive TST / T.SPOTTB; (ii) more performance of typical FGTB; (iii) higher incidence of comorbidity of active PTB.

Poor compliance, a major problem in managing TB and closely associated with side effects during treatment, may cause a series of adverse consequences including prolonged course of disease, drug resistance and recurrence [2, 6, 17-19]. Six-month ATT, especially intermittent 6-month therapy or even shorter one, therefore, mainly aims to manage the problem [7, 8, 10, 11, 17]. This study indicates that
older patients likely had higher incidence of poor compliance and side effects especially digestive symptoms and DILI (commonly seen in the region with high-burden of hepatitis B) in the longer duration compared to younger patients, whereas the efficacy of 6-month and 9-month ATT were equal. Although recurrence of disease occurred in 6 patients (3 obeyed to protocol and 3 not) with uncontrolled immuno-deficient diseases during 18-month follow-up, they were all diagnosed with DS-TB and responded to re-treatment after controlling respective comorbidity. According to 2017 update guidelines of WHO [6] and the results of this study, six-month ATT using daily therapy proved to be an appropriate option for managing FGTB in older females especially those with comorbidity of immuno-deficient diseases.

In contrast to older patients, 38 (92.7%) younger patients persisted for 9-month duration along with similar incidence of side effects and increased complete response rate. Of 29 (70.7%) younger patients with comorbidity of PTB, 8 (miliary PTB and infiltrating PTB mixed with tuberculous pleuritis presented in 6 and 2 patients, respectively) only achieved a partial healing of PTB lesions after 6-month treatment, but all achieved obvious healing after 9-month treatment, indicating that longer duration of ATT is helpful for managing systemic infection of TB. Nine-month ATT using daily therapy, therefore, may be a better option for managing FGTB in younger females at reproductive age.

There are a few weaknesses needed to mention in the study. Firstly, most of the older patients declined to receive or repeat hysteroscopy or endometrial biopsy for economic or personal reasons. It possibly reduced the diagnostic rate of FGTB in those presenting with simple performance of TBP, and not beneficial for accurate assessment of healing of FGTB lesions. Furthermore, AFB on polymerase chain reaction, beneficial for early diagnosis and efficacy assessment of FGTB treatment especially when typical TB granulomas was found in endometrium [12, 16, 20], was not taken in the study for our lacking of equipments in the specific period.

## Conclusions

The results of this study demonstrate that younger females with FGTB, in contrast to older ones, had a higher risk of systemic infection of TB and better compliance to ATT using daily therapy. Duration of ATT for managing FGTB can be simply optimized according to age category of a patient. Six-month ATT makes more benefits for older patients especially those with comorbidity of immune-deficient diseases. Nine-month ATT using daily strategy is a better option for younger patients at reproductive age.

## Declarations

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### Authors' contributions
JY design of the work, JS and WW analyzed the data and wrote the paper, JY, JS, WW and JD treated and followed the patients. JS, WW and JD collected the data, YC, YZ and MW validated imaging data. All authors have seen and approved the final draft of the paper.

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**Availability of data and materials**

All the dataset generated or analyzed during the current study are available from the corresponding author upon reasonable request.

**Ethics approval and consent to participate**

The study was approved by the ethical committee of First Hospital of Wannan Medical College. All patients gave written consent prior to antituberculosis treatment and any surgical procedure like laparotomy, laparoscopy and hysteroscopy. Informed consent for participating in the study was waived because it was an observational, real-world research and the regimen of antituberculosis treatment was based on national guidelines for treatment of extra-pulmonary tuberculosis, China.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

**Abbreviations**

FGTB: Female genital tuberculosis; EPTB: Extra-pulmonary tuberculosis; MTB: Mycobacterium tuberculosis; PTB: Pulmonary tuberculosis; DS-TB: Drug-susceptible tuberculosis; DR-TB: Drug-resistant tuberculosis; ATT: Antituberculosis therapy; DOT: Directly observed therapy; TBP: Tuberculous peritonitis; TST: Tuberculin Skin Test; CT: Computerized tomography; ADA: Adenosine deaminase; AFB: Acid-fast bacilli; TRS: Tuberculin reaction size; DILI: drug-induced liver injury.

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Tables

Due to technical limitations, the tables are only available as downloads in the supplementary files section.

Figures
Figure 1

Flow Chart of Patients with female genital tuberculosis or tuberculous peritonitis being followed at the Department of Infectious Diseases, the First Affiliated Hospital of Wanan Medical College, Wuhu, P.R. China. Abbreviations: ATT, anti-tuberculosis therapy; DR-TB, drug-resistant tuberculosis

Supplementary Files

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