Accepted Manuscript

Title: Management of MDR-TB in HIV co-infected patients in eastern europe: results from the TB:HIV study


PII: S0163-4453(17)30320-1
DOI: https://doi.org/doi:10.1016/j.jinf.2017.10.007
Reference: YJINF 4002

To appear in: Journal of Infection

Accepted date: 7-10-2017


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Management of MDR-TB in HIV co-infected patients in Eastern Europe: Results from the TB:HIV study

Management of HIV-positive patients with MDR-TB in Eastern Europe


for the TB:HIV study in EuroCoord

A full list of the TB:HIV Study Group investigators can be found in the acknowledgement section

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Keywords
Tuberculosis; HIV; MDR-TB; Eastern Europe

Word count manuscript: 2735; word count abstract: 200

Conference presentation

The results from this study were presented as a poster at the TB2016 conference, 16-17 July, Durban, South Africa (www.tb2016.org). Poster “Management of TB/HIV Co-infected Patients Including MDR-TB in Eastern Europe: Results from the TB:HIV Study”, number: P37.
Highlights

- Management of TB/HIV co-infected patients in Eastern Europe (EE) is suboptimal
- TB/HIV co-infected patients diagnosed with MDR-TB received inferior TB treatment with limited numbers of active drugs
- Access to rapid diagnostics for TB is urgently needed
- Performance of extended DST for all patients diagnosed with MDR-TB is required
- Integration of TB and HIV clinics can potentially ensure better management of patients in EE

Abstract

Objectives

Mortality among HIV patients with tuberculosis (TB) remains high in Eastern Europe (EE), but details of TB and HIV management remain scarce.

Methods

In this prospective study, we describe the TB treatment regimens of patients with multi-drug resistant (MDR) TB and use of antiretroviral therapy (ART).

Results

A total of 105 HIV-positive patients had MDR-TB (including 33 with extensive drug resistance) and 130 pan-susceptible TB. Adequate initial TB treatment was provided for 8% of patients with MDR-TB compared with 80% of those with pan-susceptible TB. By twelve months, an estimated 57.3% (95%CI 41.5-74.1) of MDR-TB patients had started adequate treatment. While 67% received ART, HIV-RNA suppression was demonstrated in only 23%.

Conclusions

Our results show that internationally recommended MDR-TB treatment regimens were infrequently used and that ART use and viral suppression was well below the target of 90%, reflecting the challenging patient population and the environment in which health care is provided. Urgent improvement of management of
patients with TB/HIV in EE, in particular for those with MDR-TB, is needed and includes widespread access
to rapid TB diagnostics, better access to and use of second-line TB drugs, timely ART initiation with viral
load monitoring, and integration of TB/HIV care.

Introduction

Although rates of tuberculosis (TB) have begun to decline in recent years, simultaneous rapid increases in
the relative contribution of multi-drug resistant TB (MDR-TB) and extensively drug resistant TB (XDR-TB) are
worrying (1-4). Eastern Europe (EE), together with central Asia, has the worlds’ highest proportions of MDR-
and XDR-TB with 9-35% of new TB cases and 49-77% of re-treatment cases being diagnosed with MDR-TB in
Belarus, Russia and Ukraine compared with 1-3% of new cases and 4-14% of re-treatment cases in Italy,
Switzerland and the United Kingdom (1, 5-8).

The MDR-TB epidemic in EE is further complicated by high rates of HIV co-infection. Whereas most other
regions have reported declining rates of new HIV infections over recent years, EE has experienced an
increase of 30% in annual number of new HIV infections since the turn of the millennium (9, 10). The
TB/HIV epidemic in EE is primarily driven by injection drug users (IDU) who often access health care late,
are co-infected with hepatitis C (HCV), and frequently have poor treatment compliance and retention in
care (11-14). We recently reported a one-year mortality rate of 27% for TB/HIV co-infected patients in EE,
and patients diagnosed with MDR-TB had roughly three-times higher mortality compared to those with
drug-susceptible TB. Further, patients from EE who initiated TB treatment with three or more active drugs
had significantly lower risk of death compared to patients who received less than three active TB drugs
(13% vs 34%) (15).

Treatment for MDR-TB is complex in terms of pill burden, drug-interactions and toxicity, and costly (1, 16-
18). MDR-TB therapy is prolonged (typically at least 20 months) although the 2016 WHO guidelines include
an option for shorter MDR-TB regimens in specific cases (16, 19). Preliminary results with a novel six
months regimen (consisting of pretomanid, bedaquiline and linezolid) may be highly effective to treat MDR-
TB (20). For now, standard of care consists of at least five drugs including a fluoroquinolone, a second-line
injectable, at least two other active drugs plus pyrazinamide during the intensive phase of treatment (19).
Reported MDR-TB treatment success rates, however, have been generally low, in particular for HIV-positive
individuals, and rarely exceed 50% (18, 21).

Epidemiological data and detailed descriptions of the clinical management of HIV-positive patients with
MDR-TB in EE remain relatively scarce (22). We prospectively studied patients with TB/HIV co-infection in
EE and in this paper report on the management of those with MDR-TB.

Methods

Study design and participants

The TB:HIV Study is a prospective cohort study including 62 TB and HIV clinics in 19 countries in EE, Western
Europe, Southern Europe, and Latin America. Adult (>16 years) HIV-positive patients with a TB diagnosis
were consecutively enrolled between 01/01/2011 and 31/12/2013. Demographic, clinical, and laboratory
data were collected on standardized case report forms at baseline and month 6, 12, and 24 (12); further
details are available at http://www.cphiv.dk/TBHIV. All participating clinics obtained ethical approval in
accordance with local rules and legislations, and the study was performed in accordance with the strobe
guidelines for observational studies (23).

Study definitions

Patients from EE were included in the present analyses if they had definite or probable TB. A diagnosis of
definite TB was based on positive culture or molecular diagnostics for Mycobacterium tuberculosis (Mtb),
and probable TB on the presence of acid fast bacilli and/or granulomatous inflammation on sputum smear or tissue biopsy specimens (12). Baseline was defined as the date TB treatment was initiated, and a baseline culture was defined as a culture obtained within one month of baseline. All TB drugs initiated within 10 days of baseline were considered to constitute initial TB treatment. Standard definitions of MDR-TB, pre-XDR-TB, and XDR-TB were used (see box), and pan-susceptible TB was defined as TB without documented drug resistance. TB treatment regimens were categorized in line with general recommendations (16, 24): 1: Treatment regimens containing RHZ (with or without ES) targeting drug-susceptible TB (24), 2: Treatment regimens containing RHZ plus a fluoroquinolone AND a second line injectable, providing empiric cover for both susceptible TB and MDR-TB, 3: Five or more MDR-TB drugs (including a fluoroquinolone AND a second line injectable) providing appropriate empiric cover for MDR-TB, 4: Regimens containing a fluoroquinolone OR a second line injectable (but not both), providing inadequate cover for MDR-TB, and 5: Any other drug combination (Fig. 1). For patients with MDR-TB, the number of active drugs in the TB regimen was calculated at various time points based on the susceptibility pattern of isolates obtained up to the given time point. If a Drug Susceptibility Test (DST) result was not available for a specific drug included in the regimen, the given \( Mtb \) isolate was either considered potentially susceptible to this drug (potentially active drugs) or resistant to this drug (known active drugs), thus representing a “best case” and “worst case” treatment scenario. Antiretroviral treatment (ART) was defined as a combination of >3 antiretroviral drugs from any ART class. Participants were considered lost to follow-up (LTFU) when no attendance was recorded at the relevant time points in those who were not known to have died.

Statistical analysis

Descriptive statistics were used for baseline characteristics of patients. Patients were stratified into three groups: MDR-TB at baseline, pan-susceptible TB at baseline, and other (no baseline DST results available or the presence of non-MDR-TB resistance patterns). MDR-TB patients who had extended DST available were
further classified as having pre-XDR or XDR-TB. Baseline characteristics among patients who had MDR-TB, those who were fully susceptible and those who did not have any baseline DST available were compared using the chi-squared test or Fisher’s exact test as appropriate for categorical variables, whereas the Kruskal-Wallis test was used to compare continuous variables.

TB and HIV treatment regimens were analyzed at months 0, 3, 7, 13, and 21. Kaplan-Meier estimates were used to estimate time to receiving adequate treatment for MDR-TB (category 3: five or more MDR-TB drugs including a fluoroquinolone AND a second line injectable), and time to starting ART censoring at last visit date (maximum 21 months) or death, whichever occurred first. Further, Kaplan-Meier estimates were also used to estimate the time to starting ART. All statistical analyses were performed using SAS (Statistical Analysis Software, Cary, NC, USA, version 9.3).

Results

Patient characteristics

Of 1406 patients enrolled into the TB:HIV study, 834 received care in EE, of these 485 had definite or probable TB and were included in this analysis (supplementary Fig. 1). A positive culture for Mtb was reported in 383 (79%) patients, 302 (79%) of these were tested for resistance to any drug, 270 (70%) for both R and H, and 68 (18%) for fluoroquinolone or second line injectables. Of the 485 patients, 105 (22%) had MDR-TB, 130 (27%) had pan-susceptible TB at baseline and 183 (38%) had no DST results; the remaining 67 (14%) had non-MDR-TB resistance patterns and were excluded from subsequent analyses. Baseline characteristics are shown in Table 1. The majority of patients were male and white, with a median age of 35 years. Recent incarceration was common, and previous TB, IDU, and HCV co-infection were significantly more common among MDR-TB patients. The median baseline CD4 cell count was 91 (interquartile range [IQR] 31-230) cells/mm³, and only 82 (17%) patients were on ART.
DST patterns for MDR-TB patients

Among the 105 patients with MDR-TB at baseline, DST results (obtained at any time during TB treatment) for other drugs ranged from 51-95% for group 1 drugs, 75-78% for group 2/3 drugs, and 49-69% for group 4 drugs (Table 2). The timing of DST testing for individual TB drugs is shown in Table 3a. Overall, 13 (12%) of MDR-TB patients had XDR-TB, 20 (19%) had pre-XDR, 35 (33%) had MDR, and 37 patients had insufficient data to determine pre-XDR/XDR-TB status.

Treatment regimen and outcomes

In Fig. 1, TB treatment regimens and outcomes are depicted from baseline through 21 months of follow-up. WHO recommended regimens were initiated in the majority of individuals with drug-susceptible TB but maintained well beyond the recommended 6 months. Despite the high population prevalence of MDR-TB, very few subjects received empiric therapy that provided cover against drug-susceptible and (multi)drug resistant TB (category 2). High rates of death were observed among MDR-TB patients and those without DST. Loss to follow-up (from both TB and HIV care) was high, ranging from 8% of MDR-TB patients to 19-20% among patients with pan-susceptible TB or no DST.

TB treatment for MDR-TB patients

Only 8 (8%) of MDR-TB patients initiated an adequate empiric MDR-TB regimen (category 3), which increased to 35 (44% of patients still under follow-up) after three months. A total of 49 MDR-TB patients ever received category 3 treatment after a median of 1.2 months (IQR 0.4-2.3), and by 12 months, an estimated 57.3% (95%CI=41.5-74.1%) had started adequate MDR-TB treatment. Similar results were obtained when excluding patients with known pre-XDR-TB or XDR-TB (data not shown).

Fluoroquinolones and second-line injectables were uncommon components of initial treatment and, if used, commonly used without adequate companion drugs (Table 3b/c). By three months, a quarter of MDR-TB patients still had not initiated a fluoroquinolone or a second-line injectable. Group 4 drugs were
variably used, a single patient had received linezolid (introduced late during the course of treatment), and no patients had received bedaquiline or delamanide. Culture conversion was documented in 45/105 (43%) of MDR-TB patients and 77/130 (59%) of patients with pan-susceptible TB after a median of 3.0 (IQR 1.8-5.1) and 2.3 (IQR 1.7-4.9, p=0.30) months, respectively.

**Activity of TB treatment for MDR-TB patients**

Fig. 2a-c illustrates the total number of drugs and the total number of “potentially” active drugs and “known” active drugs, respectively, in the MDR-TB treatment regimens. While most patients received 4 or more drugs in their regimen during the first year of treatment, more than 70% received 0, 1 or 2 drugs with demonstrated activity. A regimen containing 5 or more known active drugs was received by less than 15% of patients and a regimen containing 5 or more potentially active drugs by less than 40% of patients.

**HIV treatment for MDR-TB patients**

Despite 77% of MDR-TB patients at the time of their TB diagnosis already having an HIV diagnosis and the majority being severely immunosuppressed with a median CD4 cell count of 100 cells/mm³, only 13% were on ART at baseline (Table 1). The proportion on ART among those who remained in care increased to more than 60% (Figure 3). The majority of those on ART did not have regular HIV-RNA monitoring, and less than 20% of MDR-TB patients were documented to have had an undetectable HIV-RNA (<500 copies/mL) at any stage during follow up. The Kaplan-Meier estimate of starting ART by eight weeks was only 23%; similar results were observed for patients with drug-susceptible TB and those with no DST results (data not shown).

**Discussion**
This study shows suboptimal management of MDR-TB in HIV-positive patients in EE. Despite the high prevalence of MDR-TB, full DST testing was often restricted and delayed, resulting in prolonged use of inappropriate or failing regimens that contained few active drugs and suboptimal intensification strategies, with only half of patients ultimately receiving recommended MDR-TB therapy. In addition, we observed a low uptake of ART, inadequate viral load monitoring and low rates of HIV-RNA suppression. All of these factors are likely to have contributed to the mortality rate of nearly 50% at two years. These data point towards opportunities to improve management of HIV-positive individuals with TB in parts of EE.

The similar characteristics of patients with and without MDR-TB preclude public health strategies based on risk-stratification of patients and argue for routine DST in all TB patients. Rapid TB diagnostics and DST such as line-probe assays or cartridge-based molecular tests for \textit{Mtb} and rifampicin resistance (16) should be introduced as a priority (25), with routine evaluation of genotypic or phenotypic DST for fluoroquinolones and second line injectables of all rifampicin-resistant isolates. Thirty-nine percent of those tested for MDR-TB in our cohort had MDR-TB which is consistent with other recent reports from the EE region (8, 26). Of the patients with MDR-TB, approximately one-third had MDR, one-third had pre-XDR or XDR-TB, and one-third did not have sufficient DST data to assess whether they had (pre-)XDR-TB. We also found common co-resistance to other drugs, as described in another European multicenter study (27). Despite the setting of high MDR-TB prevalence in EE, empiric therapy rarely provided cover for MDR-TB. This is also problematic as selection for additional drug-resistance is more likely to occur early on when the bacillary burden is high. It is worth noting, however, that four in five patients who later were shown to have fully susceptible TB, initially received adequate RHZ-based treatment, as recently described (28).

In patients with MDR-TB, there were significant delays to initiate treatment for MDR-TB, and many patients never received adequate MDR-TB treatment as defined by international guidelines (19). The number of active drugs remained low at all time points; only one-third of patients were treated at some stage with at least five potentially active drugs, and only one-sixth of patients received regimens with 5 drugs known to
have activity against their Mtb isolate. This is of concern as recent studies demonstrate improved outcomes with optimal MDR-TB regimens (≥5 likely effective drugs) (29-31). Optimal MDR-TB treatment requires continued, unrestricted access to high quality medications; in a recent survey we documented that this was not the case in many clinics in EE (32). In agreement with this, linezolid, bedaquiline and delamanid were generally not used although widespread use in the current context raises significant opportunities for these drugs to be rapidly lost due to the emergence of resistance.

There is now solid evidence that ART should be initiated shortly after TB diagnosis, especially in patients with MDR-TB and those with CD4 cell counts below 50 cells/mm$^3$ (33-36). In our study, patients were severely immunosuppressed, and two-thirds of those who remained under follow-up at three months had initiated ART. However, less than 20% had documented suppressed viral loads at any time which is well short of the WHO goal of virological suppression in 90% of patients receiving ART (37).

We documented a mortality rate of nearly 50% within the first two years following initiation of TB treatment, which is similar to the mortality observed in a retrospective study from our group of HIV-patients with MDR-TB in EE which was conducted in the period 2004-2006 (14). Retrospective African studies reported mortality rates among HIV-positive patients with MDR-TB of 15-31% (38-40) and a meta-analysis of outcomes found a pooled mortality rate of 38% (41). The vast majority of deaths in our retrospective cohort were due to TB (42), which underscores the particular need for high-quality TB management for people with HIV. Other circumstances may have played a role in determining outcome of MDR-TB patients in EE. A disintegrated health care system with separated TB and HIV management remains common (32) and is likely to contribute to poor coordination of care to this often very vulnerable patient population. In particular, opioid substitution therapy for IDU is severely restricted or even prohibited in some countries despite documented positive effects on adherence to treatment and health care in general (43).
Our observational study has several limitations including the intrinsic risk of selection bias. The participating clinics were primarily located in major cities and may not be representative of all clinics across EE. DST was done locally and methods for performing DSTs varied between clinics, and when calculating the number of active drugs, we counted all drugs as equally effective which is likely to be an over-simplification. A large number of patients had missing DST results, CD4 cell counts and HIV-RNA measurements precluding a more robust evaluation of immune-virological outcomes. Nonetheless, the circumstances described reflect the difficulties for many HIV and TB clinicians working in EE, although substantial regional variability in the prevalence of MDR-TB and availability of DST, TB and ART drugs, and HIV-RNA measurements exists (12). Due to limited number of patients with MDR-TB we did not assess intraregional variability.

Conclusion

In EE, there is an urgent need for access to rapid diagnostics to guide initial TB treatment, to extended DST to all patients diagnosed with rifamycin-resistant or MDR-TB and to provide better access to second line drugs to allow the administration of optimally active MDR-TB regimens. Integration of TB and HIV services can ensure better management and support for people with HIV who frequently have IDU and HCV coinfection, including rapid initiation of fully suppressive ART.

Acknowledgements

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**Funding**

Funding: This work was supported by EU – 7th Framework (FP7/2007-2013, EuroCoord n 260694) programme. The funding source had no role in study design, the collection, analysis and interpretation of data; in the writing of the report; and in the decision to submit the article for publication.

**Disclosure of Potential Conflicts of Interest**
RFM reports personal fees from Gilead, ViiV, Merck, and Janssen outside the submitted work (non-promotional lectures on clinical aspects of HIV infection). JMM reports grants and personal fees from Abbvie, BMS, Cubist, Genentech, Gilead, Medtronik, MSD, Novartis, and ViiV Healthcare outside the submitted work. EG reports personal fees from Otsuka Novel products, Angelini, Gilead, Janssen outside the reported work. HJF reports grants from Abbvie, Gilead, MSD, BMS, ViiV, all paid to his institution outside the submitted work. OK reports personal fees and non-financial support from Gilead, personal fees from ViiV, non-financial support from Bristol Myers-Squibb outside the submitted work. All other authors report no conflict of interest.

References


Figure 1: TB treatment and outcomes over time among 105 patients with MDR-TB (left), 183 with no DST (middle) and 130 patients with susceptible TB (right) from time of start of TB treatment.

- Cat 1: RHZ (+/- ES)
- Cat 2: RHZ+FQ+2L-inj
- Cat 3: FQ+2L-inj + ≥3 MDR drugs
- Cat 4: RHZ+FQ or 2L-inj
- Cat 5: Other
- Not on treatment
- LTFU
- Dead

Variables:
- R = Rifampicin
- H = Isoniazid
- Z = Pyrazinamide
- E = Ethambutol
- S = Streptomycin
- FQ = Fluoroquinolone
- 2L-inj = Second-line injectables
- LTFU = Loss to follow-up
- DST = Drug-susceptibility test
Figure 2a-c: Number of total TB drugs (a), potentially active* (b), and known active** (c) TB drugs at each time point for 105 MDR-TB patients under follow-up.

*Potentially active: Assuming susceptibility where DST for a specific drug was missing.

**Known active: Only including known DST results and not assuming susceptibility for missing DST.
Figure 3. Antiretroviral (ART) status over time for MDR-TB patients (N=105)*

*ART and viral load (VL) was calculated +/- 1 month of the various time-points.

*ART and viral load (VL) was calculated +/- 1 month of the various time-points.
Table 1. Baseline characteristics of the included individuals (N=485)

<table>
<thead>
<tr>
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<th>All other groups (N=380)</th>
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<td>24 (23)</td>
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<td>327 (67)</td>
<td>80 (76)</td>
<td>247 (65)</td>
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<td>0 (0)</td>
<td>6 (2)</td>
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<td>59 (56)</td>
<td>195 (51)</td>
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</tr>
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<td>45 (43)</td>
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<tr>
<td>Extra pulmonary</td>
<td>24 (5)</td>
<td>4 (4)</td>
<td>20 (5)</td>
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<tr>
<td>Disseminated</td>
<td>285 (59)</td>
<td>56 (53)</td>
<td>229 (60)</td>
<td></td>
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<td>HIV+ &gt;3 months before TB diagnosis</td>
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<td></td>
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<tr>
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<td>354 (73)</td>
<td>81 (77)</td>
<td>273 (72)</td>
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<td>ART</td>
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<td>82 (17)</td>
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<td>68 (18)</td>
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<tr>
<td>Median (IQR)</td>
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<td></td>
<td></td>
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</tr>
<tr>
<td>Age</td>
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</tr>
<tr>
<td>Year</td>
<td>35 (31-40)</td>
<td>35 (31-40)</td>
<td>35 (31-40)</td>
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<td>Weight</td>
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<tr>
<td>Kg</td>
<td>60 (53-68)</td>
<td>63 (55-68)</td>
<td>59 (53-68)</td>
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<td></td>
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<td></td>
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<tr>
<td>(cells/mm³)</td>
<td>91 (31-230)</td>
<td>100 (32-267)</td>
<td>89 (31-220)</td>
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<tr>
<td>HIV-RNA</td>
<td></td>
<td></td>
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<tr>
<td>Log10 copies/ml</td>
<td>5.25 (4.40-5.75)</td>
<td>5.42 (4.46-5.91)</td>
<td>5.22 (4.35-5.70)</td>
<td>0.13</td>
</tr>
</tbody>
</table>

1. 9 individuals had missing data on previous TB.
2. 17 individuals had missing data on ethnicity.
3. 148 individuals had missing data on hepatitis C.
4. 300 individuals had missing weight at baseline.
5. 87 individuals had missing baseline CD4.
6. 211 individuals had missing baseline HIV-RNA.
7. P-values compare the characteristics between individuals with MDR-TB and all other individuals in included in the study. The chi-squared test, fischer’s exact test, or Mann-Whitney U test was used as appropriate.
8. OST = Opioid substitution therapy.
9. ART = antiretroviral therapy.
Table 2. Results from drug susceptibility test performed at any time point in follow-up among all patients with MDR-TB (N=105)

<table>
<thead>
<tr>
<th>Drug</th>
<th>N (%) with data</th>
<th>N (% of those tested) with resistance</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Group 1</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pyrazinamide</td>
<td>54 (51)</td>
<td>30 (56)</td>
</tr>
<tr>
<td>Ethambutol</td>
<td>100 (95)</td>
<td>72 (72)</td>
</tr>
<tr>
<td>Streptomycin</td>
<td>75 (71)</td>
<td>72 (96)</td>
</tr>
<tr>
<td><strong>Group 2</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fluoroquinolones</td>
<td>79 (75)</td>
<td>22 (28)</td>
</tr>
<tr>
<td><strong>Group 3</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Second-line injectables</td>
<td>82 (78)</td>
<td>35 (43)</td>
</tr>
<tr>
<td><strong>Group 4</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ethionamide/Prothionamide</td>
<td>72 (69)</td>
<td>23 (32)</td>
</tr>
<tr>
<td>Cycloserine/Terizidone</td>
<td>51 (49)</td>
<td>8 (16)</td>
</tr>
<tr>
<td>Para-Aminosalicylic Acid (PAS)</td>
<td>61 (58)</td>
<td>10 (16)</td>
</tr>
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</table>
Table 3. DST performance and TB drug use among patients with MDR-TB (N=105)

3a: N (%) of patients who had DST performed per drug up to the indicated time point among those under follow-up at given time point

<table>
<thead>
<tr>
<th>N</th>
<th>R</th>
<th>H</th>
<th>Z</th>
<th>E</th>
<th>FQ</th>
<th>2L-inj</th>
<th>PT/ET</th>
<th>CS/TZ</th>
<th>PAS</th>
<th>BDQ</th>
<th>DLM</th>
<th>LNZ</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>105</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 month</td>
<td>88</td>
<td>88 (100)</td>
<td>88 (100)</td>
<td>47 (53)</td>
<td>82 (93)</td>
<td>63 (72)</td>
<td>66 (75)</td>
<td>57 (65)</td>
<td>40 (45)</td>
<td>47 (53)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2 months</td>
<td>79</td>
<td>79 (100)</td>
<td>79 (100)</td>
<td>44 (56)</td>
<td>74 (94)</td>
<td>58 (73)</td>
<td>64 (81)</td>
<td>54 (68)</td>
<td>40 (51)</td>
<td>45 (57)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>3 months</td>
<td>72</td>
<td>72 (100)</td>
<td>72 (100)</td>
<td>40 (56)</td>
<td>68 (94)</td>
<td>54 (75)</td>
<td>61 (85)</td>
<td>51 (71)</td>
<td>37 (51)</td>
<td>41 (57)</td>
<td>0</td>
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</tr>
<tr>
<td>7 months</td>
<td>56</td>
<td>56 (100)</td>
<td>56 (100)</td>
<td>42 (57)</td>
<td>53 (95)</td>
<td>43 (77)</td>
<td>47 (84)</td>
<td>40 (71)</td>
<td>30 (54)</td>
<td>33 (59)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>13 months</td>
<td>44</td>
<td>44 (100)</td>
<td>44 (100)</td>
<td>25 (57)</td>
<td>42 (95)</td>
<td>35 (80)</td>
<td>37 (84)</td>
<td>30 (70)</td>
<td>22 (50)</td>
<td>26 (59)</td>
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</tr>
<tr>
<td>21 months</td>
<td>31</td>
<td>31 (100)</td>
<td>31 (100)</td>
<td>16 (52)</td>
<td>29 (94)</td>
<td>23 (74)</td>
<td>25 (81)</td>
<td>23 (74)</td>
<td>16 (52)</td>
<td>22 (71)</td>
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</table>

3b: N (%) of patients under follow-up who received selected anti-TB medications at the indicated time point

<table>
<thead>
<tr>
<th>N</th>
<th>R</th>
<th>H</th>
<th>Z</th>
<th>E</th>
<th>FQ</th>
<th>2L-inj</th>
<th>PT/ET</th>
<th>CS/TZ</th>
<th>PAS</th>
<th>BDQ</th>
<th>DLM</th>
<th>LNZ</th>
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<tbody>
<tr>
<td>0</td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 month</td>
<td>88</td>
<td>86 (82)</td>
<td>89 (85)</td>
<td>90 (86)</td>
<td>84 (80)</td>
<td>16 (15)</td>
<td>23 (22)</td>
<td>14 (13)</td>
<td>10 (10)</td>
<td>4 (4)</td>
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<tr>
<td>2 months</td>
<td>79</td>
<td>57 (65)</td>
<td>59 (67)</td>
<td>76 (86)</td>
<td>68 (77)</td>
<td>32 (36)</td>
<td>38 (43)</td>
<td>31 (35)</td>
<td>23 (26)</td>
<td>17 (19)</td>
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<tr>
<td>3 months</td>
<td>72</td>
<td>39 (49)</td>
<td>37 (47)</td>
<td>66 (84)</td>
<td>53 (67)</td>
<td>43 (54)</td>
<td>46 (58)</td>
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<td>30 (38)</td>
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<td>23 (32)</td>
<td>55 (76)</td>
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<td>51 (71)</td>
<td>48 (67)</td>
<td>49 (68)</td>
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<td>33 (46)</td>
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<tr>
<td>13 months</td>
<td>44</td>
<td>6 (11)</td>
<td>5 (9)</td>
<td>42 (75)</td>
<td>21 (38)</td>
<td>50 (89)</td>
<td>39 (70)</td>
<td>47 (84)</td>
<td>37 (66)</td>
<td>31 (55)</td>
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<tr>
<td>21 months</td>
<td>31</td>
<td>2 (6)</td>
<td>1 (3)</td>
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<td>7 (23)</td>
<td>25 (81)</td>
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<td>19 (61)</td>
<td>22 (71)</td>
<td>15 (48)</td>
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3c: N (%) of patients under follow-up who had ever received selected TB medications at the indicated time point

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<th>H</th>
<th>Z</th>
<th>E</th>
<th>FQ</th>
<th>2L-inj</th>
<th>PT/ET</th>
<th>CS/TZ</th>
<th>PAS</th>
<th>BDQ</th>
<th>DLM</th>
<th>LNZ</th>
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</tr>
<tr>
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<td>88</td>
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<td>75 (85)</td>
<td>81 (92)</td>
<td>76 (86)</td>
<td>33 (38)</td>
<td>40 (45)</td>
<td>32 (36)</td>
<td>23 (26)</td>
<td>17 (19)</td>
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<tr>
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<td>79</td>
<td>68 (86)</td>
<td>66 (84)</td>
<td>72 (91)</td>
<td>68 (86)</td>
<td>45 (60)</td>
<td>49 (62)</td>
<td>41 (52)</td>
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<tr>
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<td>62 (86)</td>
<td>61 (85)</td>
<td>66 (92)</td>
<td>61 (85)</td>
<td>54 (75)</td>
<td>55 (76)</td>
<td>51 (71)</td>
<td>40 (56)</td>
<td>36 (50)</td>
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<td>56</td>
<td>47 (84)</td>
<td>47 (84)</td>
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<td>51 (91)</td>
<td>49 (88)</td>
<td>48 (86)</td>
<td>40 (71)</td>
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<td>42 (95)</td>
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<td>40 (91)</td>
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<td>37 (84)</td>
<td>31 (70)</td>
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<td>30 (97)</td>
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<td>26 (84)</td>
<td>23 (74)</td>
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*DST = Drug Susceptibility Test
*R = Rifampicin*
H = Isoniazid
Z = Pyrazinamide
E = Ethambutol
FQ = Fluoroquinolone
2L-inj = Second-line injectables
PT/ET = Prothionamide/Ethionamide
CS/TZ = Cycloserine/Terizidone
PAS = Para-aminosalicylic acid
BDQ = Bedaquiline
DLM = Delamanide
LNZ = Linezolid
**Definitions of tuberculosis resistance**

<table>
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<th>Definition</th>
<th>Definition</th>
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<td>Multidrug-resistant tuberculosis (MDR-TB)</td>
<td>Tuberculosis resistant to rifampicin and isoniazid</td>
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<tr>
<td>Pre-extensively drug-resistant tuberculosis (pre-XDR-TB)</td>
<td>MDR-TB plus additional resistance against a fluoroquinolone or a second-line injectable</td>
</tr>
<tr>
<td>Extensively drug-resistant tuberculosis (XDR-TB)</td>
<td>MDR-TB plus additional resistance against a fluoroquinolone and a second-line injectable</td>
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