

UCSF

UC San Francisco Previously Published Works

Title

Fluoroquinolones and isoniazid-resistant tuberculosis: implications for the 2018 WHO guidance

Permalink

<https://escholarship.org/uc/item/9493293s>

Journal

European Respiratory Journal, 54(4)

ISSN

0903-1936

Authors

Stagg, Helen R
Bothamley, Graham H
Davidson, Jennifer A
et al.

Publication Date

2019-10-01

DOI

10.1183/13993003.00982-2019

Peer reviewed



Fluoroquinolones and isoniazid-resistant tuberculosis: implications for the 2018 WHO guidance

Helen R. Stagg ^{1,2}, Graham H. Bothamley ^{3,21}, Jennifer A. Davidson^{4,21}, Heinke Kunst^{5,21}, Maeve K. Lalor^{1,4,21}, Marc C. Lipman^{6,7,21}, Miranda G. Loutet^{4,21}, Stefan Lozewicz^{8,21}, Tehreem Mohiyuddin^{4,21}, Aula Abbara^{9,22}, Eliza Alexander^{10,22}, Helen Booth^{11,22}, Dean D. Creer^{12,22}, Ross J. Harris^{13,22}, Onn Min Kon ^{14,22}, Michael R. Loebinger^{15,22}, Timothy D. McHugh ^{16,22}, Heather J. Milburn^{17,22}, Paramita Palchadhuri^{18,22}, Patrick P.J. Phillips ^{19,22}, Erik Schmok^{3,22}, Lucy Taylor^{10,22} and Ibrahim Abubakar¹, on behalf of the London INH-R TB study group²⁰

@ERSpublications
WHO has assessed regimen recommendations for isoniazid-resistant TB to be of very low certainty. The addition of fluoroquinolones to a 12-month (isoniazid, rifamycin, ethambutol, short-duration pyrazinamide) regimen may be unnecessary in certain settings. <http://bit.ly/2XoTgNL>

Cite this article as: Stagg HR, Bothamley GH, Davidson JA, *et al.* Fluoroquinolones and isoniazid-resistant tuberculosis: implications for the 2018 WHO guidance. *Eur Respir J* 2019; 54: 1900982 [<https://doi.org/10.1183/13993003.00982-2019>].

ABSTRACT

Introduction: 2018 World Health Organization (WHO) guidelines for the treatment of isoniazid (H)-resistant (Hr) tuberculosis recommend a four-drug regimen: rifampicin (R), ethambutol (E), pyrazinamide (Z) and levofloxacin (Lfx), with or without H ([H]RZE-Lfx). This is used once Hr is known, such that patients complete 6 months of Lfx (≥ 6 [H]RZE-6Lfx). This cohort study assessed the impact of fluoroquinolones (Fq) on treatment effectiveness, accounting for Hr mutations and degree of phenotypic resistance.

Methods: This was a retrospective cohort study of 626 Hr tuberculosis patients notified in London, 2009–2013. Regimens were described and logistic regression undertaken of the association between regimen and negative regimen-specific outcomes (broadly, death due to tuberculosis, treatment failure or disease recurrence).

Results: Of 594 individuals with regimen information, 330 (55.6%) were treated with (H)RfZE (Rf=rifamycins) and 211 (35.5%) with (H)RfZE-Fq. The median overall treatment period was 11.9 months and median Z duration 2.1 months. In a univariable logistic regression model comparing (H)RfZE with and without Fqs, there was no difference in the odds of a negative regimen-specific outcome (baseline (H)RfZE, cluster-specific odds ratio 1.05 (95% CI 0.60–1.82), $p=0.87$; cluster NHS trust). Results varied minimally in a multivariable model. This odds ratio dropped (0.57, 95% CI 0.14–2.28) when Hr genotype was included, but this analysis lacked power ($p=0.42$).

Conclusions: In a high-income setting, we found a 12-month (H)RfZE regimen with a short Z duration to be similarly effective for Hr tuberculosis with or without a Fq. This regimen may result in fewer adverse events than the WHO recommendations.

This article has supplementary material available from erj.ersjournals.com

Received: 15 May 2019 | Accepted after revision: 01 July 2019

Copyright ©ERS 2019. This version is distributed under the terms of the Creative Commons Attribution Non-Commercial Licence 4.0.

Introduction

Isoniazid (H) is a key drug used in the treatment of both tuberculosis disease (TB) and latent TB infections. Research into H-resistant (Hr) TB has been neglected in favour of studies of simultaneous Hr and rifampicin (R) resistance (Rr), *i.e.* multidrug resistance (MDR) [1]. Globally, 7.1% of new incident TB patients between 2003 and 2017 had Hr disease without associated Rr (henceforth known as “Hr TB”), as did 7.9% of previously treated patients [2]. The distribution of Hr TB varies substantially by country [1, 3].

Hr has been associated with poor treatment outcomes, the need to tailor treatment regimens and the development of additional drug resistance during treatment [1]. A meta-analysis of randomised controlled trial (RCT) data, controlling for regimen, demonstrated that incidence rates of treatment failure were 10.9 times higher in Hr TB *versus* drug-sensitive disease (95% CI 5.9–20) [4]. In the same study, relapse rates in Hr TB were 1.8-fold higher (1.2–2.6) and acquired drug resistance 5.1 times higher (2.3–11.0).

Given these concerns, policymakers have issued specific treatment guidance for Hr TB. In 2018 the World Health Organization (WHO) conditionally recommended a regimen of R, ethambutol (E), pyrazinamide (Z) and levofloxacin (Lfx) with or without H ([H]RZE-Lfx), to be initiated once Hr is confirmed [5]. If treatment starts before Hr is known, it is continued until Lfx is used for 6 months, even if the duration of the other drugs is therefore longer (≥ 6 [H]RZE-6Lfx). In the absence of rapid molecular testing for Hr, overall treatment duration is thus 7.5–9 months, depending upon whether liquid or solid culture is used [6]. WHO has assessed the evidence underlying this regimen to be of very low certainty [5]. Within the UK, the National Institute for Health and Care Excellence recommends a 9-month regimen of 2 months of RZE, followed by 7 months of RE [7]. This can be extended to 12 months’ duration (10-month continuation phase), if disease is extensive. The American Thoracic Society is currently revising its guidance [8]. In 2003, they recommended a 6-month regimen of RZE, plus a fluoroquinolone (Fq) for extensive disease [9]. All bodies acknowledge the need for future studies to optimise regimens, *e.g.* to determine the implications of the resistance-causing Hr mutation(s).

In light of the 2018 WHO recommendations, we undertook a retrospective cohort study to identify the treatment regimens currently being used for Hr TB in a high-income setting with universal healthcare (London, UK). We assessed the importance of including Fqs during treatment, accounting for baseline Hr phenotype and genotype.

Methods

Study population

We included all patients aged ≥ 18 years notified in England (as a statutory requirement) to Public Health England (PHE)’s Enhanced TB Surveillance system (ETS) between January 1, 2009 and December 31, 2013 with disease caused by phenotypically Hr *Mycobacterium tuberculosis*. Baseline demographic and basic clinical and microbiological data were available from PHE. Individuals notified in London formed the retrospective cohort; additional data collection for these individuals is described below.

Treatment regimens

Detailed regimen, adherence and regimen-specific outcome information was gathered from clinical notes at the last hospital to treat the patient recorded by PHE (supplementary file 1).

Affiliations: ¹Institute for Global Health, University College London, London, UK. ²Usher Institute of Population Health Sciences and Informatics, University of Edinburgh, Edinburgh, UK. ³Respiratory Medicine, Homerton University Hospital, London, UK. ⁴Tuberculosis Unit, National Infection Service, Public Health England, London, UK. ⁵Blizard Institute, Barts and The London School of Medicine and Dentistry, Queen Mary University of London, London, UK. ⁶Respiratory Medicine, Royal Free Hospital, London, UK. ⁷UCL Respiratory, Division of Medicine, University College London. ⁸Respiratory Medicine, North Middlesex University Hospital, London, UK. ⁹Infectious Diseases, London North West University Healthcare NHS Trust, London, UK. ¹⁰National Mycobacterial Reference Service South, Public Health England, London, UK. ¹¹Tuberculosis Service, University College London Hospitals/Whittington Health, London, UK. ¹²Respiratory Medicine, Barnet General Hospital, Royal Free London NHS Foundation Trust, London, UK. ¹³Statistics, Modelling and Economics Department, Public Health England, London, UK. ¹⁴TB Service, Imperial College Healthcare, London, UK. ¹⁵Respiratory Medicine, Chelsea and Westminster Hospital, London, UK. ¹⁶Centre for Clinical Microbiology, University College London, London, UK. ¹⁷Respiratory Medicine, Guy’s and St Thomas’ Hospital, London, UK. ¹⁸Respiratory Services, Queen Elizabeth Hospital, London, UK. ¹⁹Dept of Medicine and Dept of Epidemiology and Biostatistics, University of California San Francisco, San Francisco, CA, USA. ²⁰Additional London INH-R TB study group members are listed in the acknowledgements section. ²¹These authors contributed equally to this manuscript and are presented alphabetically. ²²These authors contributed equally to this manuscript and are presented alphabetically.

Correspondence: Helen R. Stagg, Usher Institute of Population Health Sciences and Informatics, University of Edinburgh, Edinburgh, EH8 9DX, UK. E-mail: helen.stagg@ed.ac.uk

Regimens were described and categorised. The rifamycins (Rf) R and rifabutin were grouped together, as were the injectables, Fqs other than moxifloxacin (M), and the previously named group 4/5 drugs [10, 11]. A binary regimen variable was created of RfZE regimens in the presence or absence of H with or without an additional Fq: (H)RZE *versus* (H)RZE-Fq/M. If additional drugs were included, the regimen was not counted within the binary variable.

The presence of high-dose H within the regimen was documented, as was whether Rf, Z or E were dosed thrice weekly (as opposed to more frequently). The length of time a patient was treated before the regimen was adapted to account for Hr (which was dependent on the duration of drug sensitivity testing (DST)) was grouped 0 to <2, 2 to <6 and \geq 6 months.

Genotyping and phenotyping

Phenotypic DSTs for first-line drugs were conducted on baseline samples. DSTs for second-line drugs were conducted if resistance to R or two or more other first-line drugs (but not H alone, although this could be requested) was detected. These results were recorded within ETS. Patients were grouped according to the baseline drug resistance pattern of their disease.

The degree of phenotypic resistance to H was extracted from the National Mycobacterium Reference Service (NMRS)-South system (supplementary file 1).

Whole-genome sequencing (WGS) to detect resistance mutations was undertaken for a subset of patients among those notified 2012–2013 using an Illumina HiSeq (San Diego, CA, USA) at the PHE central sequencing unit [12].

Other exposure variables

Age, sex, being born in the UK, ethnic group, social risk factors (homelessness, problematic drug use, problematic alcohol use and imprisonment), previous diagnosis of TB and inpatient information came from ETS. Decisions surrounding the grouping of these variables are documented in supplementary file 1.

An outbreak of Hr TB has been present in (mainly north) London since 1995 [13, 14]. Due to awareness of this outbreak among clinicians, patients with epidemiological risk factors consistent with the outbreak (in which nonadherence was common and treatment outcomes poor) may have been treated differently from other patients.

An additional variable documented if a patient had issues adhering to treatment, according to their clinical notes (supplementary file 1).

Outcomes

A patient's treatment period for Hr TB is made up of up to three components: the regimen used prior to Hr being known; the regimen used once Hr is known; and (potentially) a further regimen or regimens if the Hr regimen is insufficiently effective. Overall treatment outcomes (available in ETS) capture this entire period. Regimen-specific outcomes, taken from clinical notes, document the effectiveness of the Hr regimen and thus capture only the first two components (table 1). For the regression model, the neutral and positive groups were merged to create a binary outcome.

Analysis

Data were cleaned in Microsoft Excel (Redmond, WA, USA) and analysed in Stata 15 (StataCorp, College Station, TX, USA).

The characteristics of the London cohort were assessed. Descriptive analyses of the regimens used were undertaken, followed by regression analyses. Initially, individuals with additional phenotypic drug resistance identified in baseline samples taken were excluded from the regression models, unless resistance was to streptomycin (S). This was because S is not routinely used in the treatment of drug-sensitive or MDR TB in the UK [7, 16]. Random-effects univariable logistic regression models were built to examine the impact of different factors on the likelihood of negative regimen-specific outcomes, with a random effect included on National Health Service (NHS) trust to adjust for clustering.

A multivariable logistic regression model was then built, using the binary regimen categorisation as the main exposure and including a random effect on NHS trust. Details of confounder selection, *etc.* are presented in supplementary file 1.

Sensitivity and extended analyses

Four additional logistic regression models were run. The first included Hr genotyping results. Next, adherence was substituted for thrice-weekly dosing. The third included all patients, regardless of whether

TABLE 1 Classification of regimen-specific outcomes

Components	Comments
<p>Negative Treatment completed, followed by recurrence; outcome missing, but recurrence; or neutral outcome, followed by recurrence</p> <p>Died due to TB or TB-associated death ≥ 2 weeks after starting treatment</p> <p>Treatment stopped early or regimen changed due to worsening/not improving, treatment failure, adverse events or the development of additional drug resistance</p> <p>Additional drug resistance developed during treatment</p>	<p>Recurrence of disease ≥ 12 months after notification; recurrences documented until the end of 2015 (the most recent available data at the time of analysis); if disease recurred after the end of treatment at any time and the patient re-presented to the same hospital, this was classified as a negative outcome</p> <p>Death before 2-week threshold considered to be too early to be influenced by the treatment [15]</p> <p>Length extended, antibiotics added/removed, frequency altered, dose altered, treatment stopped</p>
<p>Neutral Died from TB or TB-associated death within 2 weeks of starting treatment</p> <p>Died from non-TB related causes, or cause unknown</p> <p>Treatment stopped early or regimen changed due to non-adherence, loss to follow-up, patient choosing to cease their medication, pregnancy or comorbidities</p> <p>Patient transferred to another hospital during their treatment</p>	<p>To any drug</p> <p>Death before 2-week threshold considered to be too early to be influenced by the treatment [15]</p> <p>No further documentation; transfer before any negative outcomes occurred</p>
<p>Positive Treatment completed as initially prescribed (once Hr known); or treatment completed, no recurrence</p>	

Regimen-specific outcomes (extracted from clinical notes) presented in detail. The first outcome arising per patient was documented, unless a negative outcome occurred after one that is neutral. TB: tuberculosis; Hr: isoniazid resistance.

they were resistant to drugs in addition to H (and S). The fourth was a *post hoc* model adjusting for factors associated with the use of Fqs.

Ethical permissions

PHE is legislated by the National Information Governance Board for Health and Social Care to hold and analyse surveillance data for public health purposes under section 251 of the NHS Act 2006. This retrospective cohort study was approved by the London Camberwell St Giles research ethics committee (16/LO/1269) and, in addition, given permission to undertake data extraction without consent under section 251 (Confidentiality Advisory Group reference 16/CAG/0092).

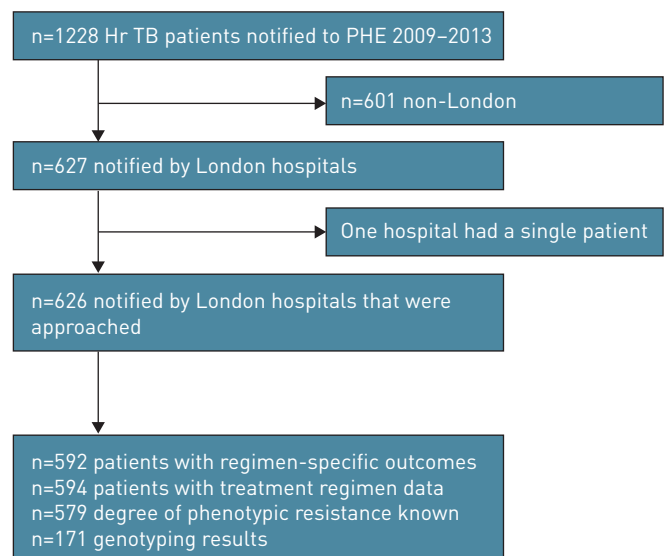


FIGURE 1 Flow chart of participants. Hr: isoniazid-resistant; TB: tuberculosis; PHE: Public Health England.

TABLE 2 Demographic and clinical baseline characteristics of the 626 individuals in the London cohort

Overall	626 (100)
Year	
2009	137 (21.9)
2010	118 (18.8)
2011	141 (22.5)
2012	125 (20.0)
2013	105 (16.8)
Missing	0 (0.0)
Sex	
Male	380 (60.7)
Female	246 (39.3)
Missing	0 (0.0)
Age years	
18–37	358 (57.2)
38–57	199 (31.8)
58–77	62 (9.9)
≥78	7 (1.1)
Missing	0 (0.0)
UK born	
No	497 (79.4)
Yes	121 (19.3)
Missing	8 (1.3)
Ethnic group	
White	97 (15.5)
Black African	125 (20.0)
Black Other	45 (7.2)
Indian subcontinent	270 (43.1)
Other	85 (13.6)
Missing	4 (0.6)
Social risk factors	
No or unknown	510 (81.5)
One or more ever	37 (5.9)
One or more current	79 (12.6)
Previous TB diagnosis	
No	575 (91.9)
Yes	20 (3.2)
Missing	31 (5.0)
Inpatient	
No	422 (67.4)
Yes	190 (30.4)
Missing	14 (2.2)
Site of disease	
Pulmonary±extrapulmonary, smear +ve	194 (31.0)
Pulmonary±extrapulmonary, smear -ve	159 (25.4)
Meningeal TB or other CNS involvement	24 (3.8)
Other extrapulmonary	249 (39.8)
Missing	0 (0.0)
Part of outbreak	
No	501 (80.0)
Yes	65 (10.4)
Missing	60 (9.6)
Any additional drug resistance	
No	453 (72.4)
Yes	173 (27.6)

Data are presented as n (column %). TB: tuberculosis; CNS: central nervous system; ±: with or without; -ve: negative; +ve: positive.

Results

Patient population

1228 individuals with Hr TB were notified in England between 2009 and 2013 (figure 1). Of these, 626 (51.0%) were notified by 31 hospitals (supplementary file 2) in London (19 NHS trusts). One hospital

TABLE 3 Regimen-specific outcomes and availability of regimen data

	Negative	Neutral	Positive
Frequency of outcome n (% of 592)	97 (16.4)	87 (14.7)	408 (68.9)
Details of outcome	n=3 recurrences after treatment was completed n=2 recurrences after an otherwise neutral or missing outcome n=3 developed additional drug resistance (two to R and one to clarithromycin; additionally, one patient developed resistance to E and one to R, but this was predated by other negative outcomes) n=1 stopped treatment for negative reasons n=7 had the length of their treatment extended for negative reasons n=78 treatment regimen changes by other means for negative reasons n=3 deaths from TB >2 weeks after treatment started		
Regimen data available n (column % within outcome)	95 (97.9)	79 (90.8)	408 (100.0)
Details of regimen data			n=374 with regimen data for the full duration of treatment n=7 partial uncertainties surrounding the drugs present in the regimen n=27 some date information missing

Regimen-specific outcomes and treatment regimen availability for the 592 (94.6%) out of 626 of individuals with an outcome recorded in the London cohort. E: ethambutol; R: rifampicin; TB: tuberculosis.

had only a single patient and was not approached for local approvals. The baseline characteristics of the London cohort are described in table 2.

Phenotypic testing for non-H drug resistance revealed that 173 (27.6%) out of 626 patients within the London cohort had additional drug resistance at baseline (supplementary file 3). The most common resistance was towards S (139 (22.2%) out of 626).

The majority of samples were documented in the NMRS system as highly Hr at baseline (495 (79.1%) out of 626). Three (0.5%) displayed borderline results; one was listed as drug sensitive (0.2%); and 35 (5.6%) were present in the system, but did not have their Hr levels logged. 47 individuals could not be found within NMRS, but were recorded as Hr within ETS.

Regimen-specific outcomes

Regimen-specific outcomes were available for 592 (94.6%) out of 626 patients (table 3). 97 (16.4%) had a negative outcome.

Relationship between treatment regimens and regimen-specific outcomes

Of the 626 patients, 582 (93.0%) had both a regimen-specific outcome recorded and treatment information. Of these, 538 (92.4%) were not resistant to drugs in addition to H, apart from S, and 84 had a negative regimen-specific outcome (three of which were recurrences). 498 (92.6%) out of 538 were treated with (H)RfZE or (H)RfZE-Fq/M (table 4). For a more detailed description of the treatment regimens, see supplementary file 4.

Differences in the odds of a negative regimen-specific outcome were not detected between patients treated with (H)RfZE (baseline) and (H)RfZE-Fq/M (cluster-specific OR 1.02, 95% CI 0.59–1.77; $p=0.93$; table 4). None of the other treatment regimens or associated factors were found to be associated with the odds of negative outcomes (table 4, supplementary file 5). We observed more negative outcomes with the use of thrice weekly dosing *versus* more frequent dosing (OR 1.81, 95% CI 0.83–3.94), but this may have been a chance finding ($p=0.15$).

Seven exposure variables/confounders were included in the multivariable model: regimen, thrice-weekly dosing, Hr phenotype, sex, age (linear variable), ethnic group and previous TB diagnosis. Evidence for effect modification was not found. In the final multivariable model of 435 patients (table 5), there was no

TABLE 4 Univariable logistic regression of treatment regimen and associated factors as predictors of negative outcomes

	Overall n (column %)	Negative outcome n (row %)	OR (95% CI)	p-value
Overall	538 (100)	84 (15.6)		
Regimen				
(H)RfZE	306 (56.9)	46 (15.0)		0.93
(H)RfZE-Fq/M	192 (35.7)	30 (15.6)	1.02 (0.59–1.77)	
Missing	40 (7.4)	8 (20.0)		
Thrice-weekly dosing				
More frequent	464 (86.2)	66 (14.2)		0.15
Thrice weekly	53 (9.9)	12 (22.6)	1.81 (0.83–3.94)	
Missing	21 (3.9)	6 (28.6)		
Time before Hr known				
0 to <2 months	325 (60.4)	56 (17.2)		0.27
2 to <6 months	159 (29.6)	18 (11.3)	0.62 (0.34–1.13)	
≥6 months	10 (1.9)	2 (20.0)	1.11 (0.22–5.66)	
Missing	44 (8.2)	8 (18.2)		
Phenotype				
Highly resistant	442 (82.2)	69 (15.6)		0.73
Resistant	36 (6.7)	5 (13.9)	0.88 (0.32–2.39)	
Borderline, sensitive or results not logged	29 (5.4)	6 (20.7)	1.46 (0.55–3.88)	
Missing	31 (5.8)	4 (12.9)		
Adherence issues or treatment gaps				
No or unknown	425 (79.0)	64 (15.1)		0.29
Not severe or of unknown severity	56 (10.4)	13 (23.2)	1.62 (0.80–3.28)	
Severe	57 (10.6)	7 (12.3)	0.72 (0.30–1.73)	

Univariable logistic regression of treatment regimen and associated factors as predictors of negative regimen-specific outcomes. Included patients were notified in London, had regimen-specific outcome and regimen information, and their disease was without additional drug resistance, unless to streptomycin. Each model contains the patients without missing data. H: isoniazid; Rf: rifamycin; Z: pyrazinamide; E: ethambutol; Fq: fluoroquinolones; M: moxifloxacin; Hr: isoniazid resistance.

discernible difference in the odds of a negative outcome between the two regimens (OR 0.99, 95% CI 0.53–1.85; $p=0.97$). The association between thrice-weekly dosing and negative outcomes was slightly strengthened in terms of the effect estimate (OR 2.34, 95% CI (0.90–6.09), although the association observed could still have been due to chance ($p=0.09$).

Impact of genotype and other sensitivity analyses

The most common Hr genotypes observed were *fabG1* C-15 T (87 (50.9%) out of 171) and *katG* S315 T (75 (43.9%) out of 171; supplementary file 6). For 10 (5.8%) out of 171 strains, sequencing either failed, no resistance mutations were detected or it was not known whether the single nucleotide polymorphisms (SNPs) found generate drug resistance.

In a univariable model, no difference was seen in the likelihood of a negative treatment outcome between the *katG* S315 T/N genotypes and a *fabG1* C-15 T baseline (OR 1.17, 95% CI 0.42–3.31; $p=0.76$). In a multivariable model, evidence for effect modification by genotype was not detected. Genotype was not independently associated with the outcome (supplementary file 7). In this model, there was a suggestion that the odds of a negative regimen-specific outcome were reduced for (H)RfZE-Fq/M versus (H)RfZE (OR 0.57, 95% CI 0.14–2.28), but we were underpowered for this analysis ($p=0.42$).

Inclusion of other potential confounder sets in the multivariable model did not impact our findings (supplementary file 8).

Discussion

In this analysis of Hr TB patients notified by London hospitals between 2009 and 2013, 16.4% of individuals had a negative outcome. (H)RfZE and (H)RfZE-Fq/M regimens were taken by 92.6% of individuals without additional drug resistance (apart from to S) and with both regimen and regimen-specific outcome data. Among these patients, we found no discernible difference in the odds of a negative regimen-specific outcome between (H)RfZE and (H)RfZE-Fq/M regimens. Examining individuals

TABLE 5 Multivariable logistic regression of treatment regimen as a predictor of negative outcomes

	OR (95% CI)	p-value
Regimen		
(H)RfZE		0.97
(H)RfZE-Fq/M	0.99 [0.53–1.85]	
Thrice-weekly dosing		
More frequent		0.09
Thrice weekly	2.34 [0.90–6.09]	
Phenotype		
Highly resistant		0.66
Resistant	0.64 [0.17–2.43]	
Borderline, sensitive or results not logged	1.40 [0.45–4.31]	
Missing		
Sex		
Male		0.02
Female	2.05 [1.13–3.71]	
Age years		
18–37		0.46
Per 20-year increase	1.18 [0.75–1.86]	
Ethnic group		
White		0.15
Black African	0.42 [0.15–1.18]	
Black Other	0.33 [0.08–1.39]	
Indian subcontinent	0.58 [0.23–1.45]	
Other	1.10 [0.42–2.92]	
Previous TB diagnosis		
No		0.13
Yes	3.12 [0.75–12.91]	

Multivariable logistic regression of treatment regimen as a predictor of negative regimen-specific outcomes in patients without additional drug resistance, unless to streptomycin, adjusted for all variables in the table. Model contains 435 patients. H: isoniazid; Rf: rifamycin; Z: pyrazinamide; E: ethambutol; Fq: fluoroquinolones; M: moxifloxacin; TB: tuberculosis.

with a positive treatment outcome, the overall duration of treatment was generally 12 months, with Z durations of 2 months in the initiation phase. After adjustment for Hr genotype, the likelihood of a negative outcome was found to be lower among individuals treated with (H)RfZE-Fq/M, but this analysis was underpowered.

Our findings sit in the context of preceding work on the relative efficacy and effectiveness of different regimens for Hr TB, including four meta-analyses [17–20]. FREGONESE *et al.*'s [17] individual-level patient meta-analysis, the foundation of the 2018 WHO guidelines, showed a value for including a Fq in continuous (H)RZE regimens, and suggested equivalence between 6 and 8–9 months of (H)RZE. The WHO acknowledges that overall treatment length findings may be subject to confounding by indication, due to patients with more complex sites of disease receiving longer regimens [5].

Notably, global RCT evidence for the effectiveness of Fqs in non-MDR-TB derive solely from the Rifaquin trial, as ReMox did not demonstrate non-inferiority when H was replaced with M for non-MDR TB [21, 22]. When considering the choice of Fq, although the WHO recommends the use of Lfx, M was generally used in our study. Within FREGONESE *et al.* [17], roughly equal numbers of studies used these two drugs, which were not directly compared. However, comparative data are available from a MDR-TB trial (no difference in treatment outcomes when comparing the two drugs; fewer adverse events for M) [23], and rabbit and mouse models (M broadly superior over Lfx) [24, 25]. Lfx doses in such studies may have been too low [26, 27]. Further RCTs are required.

The above meta-analyses were unable to thoroughly consider the role of Hr genotype and phenotype in treatment decisions; the evidence from previous observational studies is unclear [1, 28]. Where adjustment for genotype in observational studies has been undertaken, it was largely for *inhA* and *katG*. In our cohort, with a very high prevalence of *fabG1* in addition to *katG* mutations, we find an indication that the Hr genotype is influential. *fabG1* is part of the *inhA* operon and is involved in fatty acid synthesis; SNPs within the gene are known to confer Hr [29, 30].

The evidence currently underpinning global treatment guidelines for Hr TB is limited. Our study adds to this discussion, including consideration of the effect of resistance phenotype and genotype on the regimen–outcomes relationship. Importantly, in our core analysis, 192 patients received a Fq in addition to (H)RfZE, which provides substantial new evidence to that presented by FREGONESE *et al.* [17], whose analysis of treatment success included 251 patients receiving a Fq. Our findings did not differ when site of disease was adjusted for as a confounder (including meningeal TB or other central nervous system involvement; data not shown) and when patients with additional drug resistance were included.

Within this study, actual rather than intended treatment durations were captured, which prevented us from undertaking analyses of the impact of overall or drug-specific durations. Importantly, however, when considering 9 *versus* 12 months of treatment, the majority of negative outcomes occurred before 9 months and the number of relapses was small, with two of the three occurring after >15 months of treatment. Thus our data may indicate the potential to shorten treatment to 9 months in our setting. Some patient notes could not be accessed as patients had died. This was unlikely to have been of a magnitude sufficient to bias our findings. We did not differentiate between recurrence due to relapse *versus* reinfection, and thus may have overestimated the number of negative outcomes (nondifferential misclassification). Gaps in phenotypic data arose due to 1) missing records within NMRS from a specific period and reference laboratory; and 2) incomplete data entry into NMRS from the reference laboratory (cross-tabulations against patient characteristics did not indicate that this particularly affected any specific patient groups). The phenotypic and genotypic Hr patterns documented summarise that of the overall bacterial population; the presence of minor strains will not have been captured. Our findings about thrice-weekly dosing may represent the use of such a dosing pattern specifically among patients where directed observation of treatment was deemed necessary. HIV status, a potential confounder, was not obtainable during data collection.

Despite these limitations, there are important ramifications for our findings both nationally and internationally. We document a drug combination that differs from that recommended (with very low certainty) by the WHO [5], which may be as effective. We note that, if the overall duration of treatment is long enough (12 months), a Fq may not be necessary in certain settings, even with relatively short durations (median 2 months in the initiation phase) of Z. Notably, in settings where DST occurs *via* phenotyping from cultures, the WHO regimen $\geq 6[H]RZE-6Lfx$ is likely to have total duration of 7.5–9 months, when time to result is considered. This also affects the longer regimen in their 6- *versus* 8–9-month duration comparison; the latter translates to 9.5–12 months. By comparison, in settings undertaking rapid genotyping directly from patient samples, the WHO regimen duration would be 6 months and the average duration documented here ~ 10 months.

Global regimen choices will depend upon the trade-off between patient desire for regimens of minimal length, adherence concerns, adverse events, ease of administration and cost. Costs are raised if fixed dose combination pills cannot be used and Fqs are added in. When it comes to comparing the likelihood of adverse events, the trade-off would be between a longer duration of E, but shorter duration of Z in our predominantly used regimen, *versus* continued Z and the addition of Lfx, as per the WHO recommendations. Each of these drugs has its own distinct adverse event profile [23, 31].

Fq DST results are important when deciding on Fq use within a Hr regimen. Only 48 individuals in the London cohort had their baseline samples tested for resistance to M. In 2018, PHE rolled-out prospective WGS to provide routine resistance predictions and mutation identification, thus improving the rapidity of DST and coverage of second-line testing. New molecular Hr tests can also aid rapidity, as the use of WGS still depends on culture [32].

Within the limitations of an observational study, where the use of Fqs was not randomised, we find in a high-income setting with comprehensive patient management, a 12-month (H)RfZE regimen with a short Z duration to be similarly effective for Hr TB, with or without a Fq. Hr genotype may influence these findings. In the absence of Fqs and long durations of Z, this regimen may have fewer adverse events than the WHO recommended $\geq 6[H]RZE-6Lfx$. RCTs should be undertaken to provide stronger global recommendations.

Acknowledgements: H.R. Stagg would also like to acknowledge the National Health Service and Research and Development office staff at different hospital sites, who are too numerous to name individually, but without whom this study would not have been possible. The authors also wish to thank Patryk Solinski, a Patient and Public Involvement representative on H.R. Stagg's Fellowship Advisory Panel and to acknowledge the following clinicians for their help with data collection: David Adeboyeke (Central Middlesex Hospital) and Devan Vaghela (Barts Health NHS Trust).

Additional London INH-R TB study group members are as follows (please note that these individuals contributed equally and are presented in alphabetical order): Lucy V. Baker, Respiratory Medicine, University Hospital Lewisham, London, UK; Jessica C. Barrett, Infectious Diseases, London North West University Healthcare NHS Trust, London, UK; Helen Burgess, Respiratory Medicine, West Middlesex University Hospital, London, UK; Catherine Cosgrove,

Infectious Diseases, St George's Hospital, London, UK; Anne Dunleavy, Respiratory Medicine, St George's Hospital, London, UK; Marie Francis, Institute for Global Health, University College London, London, UK; Urmi Gupta, Respiratory Medicine, King George Hospital, London, UK; Shahid Hamid, Respiratory Medicine, Princess Royal University Hospital, London, UK; Brigitte M. Haselden, Respiratory Medicine, The Hillingdon Hospital NHS Foundation Trust, London, UK; Emma Holden, Respiratory Medicine, Kingston Hospital, London, UK; Vanessa Kahr, Respiratory Services, St Helier Hospital, London, UK; William Lynn, Infectious Diseases, London North West University Healthcare NHS Trust, London, UK; Felicity M. Perrin, Respiratory Medicine, King's College Hospital NHS Foundation Trust, London, UK; Ananna Rahman, Respiratory Medicine Service, Barts Health NHS Trust, London, UK; and Mohammad R. Soobratty, Respiratory Services, Croydon University Hospital, London, UK.

Support statement: This report is independent research supported by the National Institute for Health Research (Post Doctoral Fellowship, Helen R. Stagg, PDF-2014-07-008). The views expressed in this publication are those of the author (s) and not necessarily those of the NHS, the National Institute for Health Research or the Department of Health. The sponsors had no role in the study design; collection, analysis and interpretation of the data; writing of the report; and in the decision to submit the paper for publication. Funding information for this article has been deposited with the Crossref Funder Registry.

Conflict of interest: H.R. Stagg reports grants from National Institute for Health Research, UK (PDF-2014-07-008), during the conduct of the study; grants from Medical Research Council, UK (MC_PC_17101) and Korea Health Industry Development Institute, outside the submitted work. G.H. Bothamley has nothing to disclose. J.A. Davidson has nothing to disclose. H. Kunst has nothing to disclose. M.K. Lalor has nothing to disclose. M.C. Lipman has nothing to disclose. M.G. Loutet has nothing to disclose. S. Lozewicz has nothing to disclose. T. Mohiyuddin has nothing to disclose. A. Abbara has nothing to disclose. E. Alexander reports personal fees for advisory board work from Inamed, outside the submitted work. H. Booth has nothing to disclose. D.D. Creer has nothing to disclose. R.J. Harris has nothing to disclose. O.M. Kon has nothing to disclose. M.R. Loebinger has nothing to disclose. T.D. McHugh has nothing to disclose. H.J. Milburn has nothing to disclose. P. Palchaudhuri has nothing to disclose. P.P.J. Phillips has nothing to disclose. E. Schmok has nothing to disclose. L. Taylor has nothing to disclose. I. Abubakar reports grants from NIHR and MRC, outside the submitted work.

References

- 1 Stagg HR, Lipman MC, McHugh TD, *et al.* Isoniazid-resistant tuberculosis: a cause for concern? *Int J Tuberc Lung Dis* 2017; 21: 129–139.
- 2 World Health Organization (WHO). Global Tuberculosis Report 2018. Geneva, WHO. www.who.int/tb/publications/global_report/en/ Date last updated: September 2018. Date last accessed: April 2018.
- 3 Jenkins HE, Zignol M, Cohen T. Quantifying the burden and trends of isoniazid resistant tuberculosis, 1994–2009. *PLoS One* 2011; 6: e22927.
- 4 Menzies D, Benedetti A, Paydar A, *et al.* Effect of duration and intermittency of rifampin on tuberculosis treatment outcomes: a systematic review and meta-analysis. *PLoS Med* 2009; 6: e1000146.
- 5 World Health Organization (WHO). WHO Treatment Guidelines For Isoniazid-Resistant Tuberculosis: Supplement to the WHO Treatment Guidelines for Drug-Resistant Tuberculosis. Geneva, WHO. www.who.int/tb/publications/2018/WHO_guidelines_isoniazid_resistant_TB/en/ Date last updated: July 2018. Date last accessed: July 2018.
- 6 Drobniewski F, Cooke M, Jordan J, *et al.* Systematic review, meta-analysis and economic modelling of molecular diagnostic tests for antibiotic resistance in tuberculosis. *Health Technol Assess* 2015; 19: 1–188.
- 7 National Institute for Health and Care Excellence. Tuberculosis. www.nice.org.uk/guidance/ng33/resources/tuberculosis-prevention-diagnosis-management-and-service-organisation-1837390683589 Date last updated: January 2016. Date last accessed: April 2018.
- 8 Nahid P, Dorman SE, Alipanah N, *et al.* Official American Thoracic Society/Centers for Disease Control and Prevention/Infectious Diseases Society of America Clinical Practice Guidelines: Treatment of Drug-Susceptible Tuberculosis. *Clin Infect Dis* 2016; 63: e147–e195.
- 9 Blumberg HM, Burman WJ, Chaisson RE, *et al.* American Thoracic Society/Centers for Disease Control and Prevention/Infectious Diseases Society of America: treatment of tuberculosis. *Am J Respir Crit Care Med* 2003; 167: 603–662.
- 10 Tiberi S, Scardigli A, Centis R, *et al.* Classifying new anti-tuberculosis drugs: rationale and future perspectives. *Int J Infect Dis* 2017; 56: 181–184.
- 11 World Health Organization (WHO). WHO Treatment Guidelines for Drug-Resistant Tuberculosis. Date last updated: October 2016. Date last accessed: April 2018. <https://apps.who.int/iris/bitstream/handle/10665/250125/9789241549639-eng.pdf;jsessionid=605E8FC225283D2EA72288>.
- 12 Walker TM, Kohl TA, Omar SV, *et al.* Whole-genome sequencing for prediction of *Mycobacterium tuberculosis* drug susceptibility and resistance: a retrospective cohort study. *Lancet Infect Dis* 2015; 15: 1193–1202.
- 13 Maguire H, Brailsford S, Carless J, *et al.* Large outbreak of isoniazid-monoresistant tuberculosis in London, 1995 to 2006: case-control study and recommendations. *Euro Surveill* 2011; 16: 19830.
- 14 Smith CM, Trienekens SC, Anderson C, *et al.* Twenty years and counting: epidemiology of an outbreak of isoniazid-resistant tuberculosis in England and Wales, 1995 to 2014. *Euro Surveill* 2017; 22: 30467.
- 15 Stagg HR, Abubakar I, Brown J, *et al.* Towards better guidance on caseload thresholds to promote positive tuberculosis treatment outcomes: a cohort study. *BMC Med* 2016; 14: 52.
- 16 Potter JL, Capstick T, Ricketts WM, *et al.* Streptomycin. TB Drug Monographs. www.tbdrugmonographs.co.uk/streptomycin.html Date last accessed: July 2019. Date last updated: August 2018.
- 17 Fregonese F, Ahuja SD, Akkerman OW, *et al.* Comparison of different treatments for isoniazid-resistant tuberculosis: an individual patient data meta-analysis. *Lancet Respir Med* 2018; 6: 265–275.
- 18 Gegia M, Winters N, Benedetti A, *et al.* Treatment of isoniazid-resistant tuberculosis with first-line drugs: a systematic review and meta-analysis. *Lancet Infect Dis* 2017; 17: 223–234.
- 19 Menzies D, Benedetti A, Paydar A, *et al.* Standardized treatment of active tuberculosis in patients with previous treatment and/or with mono-resistance to isoniazid: a systematic review and meta-analysis. *PLoS Med* 2009; 6: e1000150.

- 20 Stagg HR, Harris RJ, Hatherell HA, *et al.* What are the most efficacious treatment regimens for isoniazid-resistant tuberculosis? A systematic review and network meta-analysis. *Thorax* 2016; 71: 940–949.
- 21 Gillespie SH, Crook AM, McHugh TD, *et al.* Four-month moxifloxacin-based regimens for drug-sensitive tuberculosis. *N Engl J Med* 2014; 371: 1577–1587.
- 22 Jindani A, Harrison TS, Nunn AJ, *et al.* High-dose rifapentine with moxifloxacin for pulmonary tuberculosis. *N Engl J Med* 2014; 371: 1599–1608.
- 23 Kang YA, Shim TS, Koh WJ, *et al.* Choice between levofloxacin and moxifloxacin and multidrug-resistant tuberculosis treatment outcomes. *Ann Am Thorac Soc* 2016; 13: 364–370.
- 24 Maitre T, Petitjean G, Chauffour A, *et al.* Are moxifloxacin and levofloxacin equally effective to treat XDR tuberculosis? *J Antimicrob Chemother* 2017; 72: 2326–2333.
- 25 Sarathy J, Blanc L, Alvarez-Cabrera N, *et al.* Fluoroquinolone efficacy against tuberculosis is driven by penetration into lesions and activity against resident bacterial populations. *Antimicrob Agents Chemother* 2019; 63: e02516-18.
- 26 Al-Shaer MH, Alghamdi WA, Alsultan A, *et al.* Fluoroquinolones in drug-resistant tuberculosis: culture conversion and pharmacokinetic/pharmacodynamic target attainment to guide dose selection. *Antimicrob Agents Chemother* 2019; 63: e00279-19.
- 27 Deshpande D, Pasipanodya JG, Mpagama SG, *et al.* Levofloxacin pharmacokinetics/pharmacodynamics, dosing, susceptibility breakpoints, and artificial intelligence in the treatment of multidrug-resistant tuberculosis. *Clin Infect Dis* 2018; 67: S293–S302.
- 28 Thai PVK, Ha DTM, Hanh NT, *et al.* Bacterial risk factors for treatment failure and relapse among patients with isoniazid resistant tuberculosis. *BMC Infect Dis* 2018; 18: 112.
- 29 Marrakchi H, Ducasse S, Labesse G, *et al.* MabA (FabG1), a *Mycobacterium tuberculosis* protein involved in the long-chain fatty acid elongation system FAS-II. *Microbiology* 2002; 148: 951–960.
- 30 Seifert M, Catanzaro D, Catanzaro A, *et al.* Genetic mutations associated with isoniazid resistance in *Mycobacterium tuberculosis*: a systematic review. *PLoS One* 2015; 10: e0119628.
- 31 Zellweger JP. Treatment of tuberculosis. *Expert Rev Respir Med* 2007; 1: 85–97.
- 32 Xie YL, Chakravorty S, Armstrong DT, *et al.* Evaluation of a rapid molecular drug-susceptibility test for tuberculosis. *N Engl J Med* 2017; 377: 1043–1054.