Original article



# Introducing the Best Six Loci in Mycobacterial Interspersed Repetitive Unit-Variable-Number Tandem Repeat (MIRU-VNTR) Typing for *Mycobacterium Tuberculosis* Genotyping

Mahdis Ghavidel<sup>1, 2</sup>, Keyvan Tadayon<sup>3</sup>, Nader Mosavari<sup>4</sup>, Kimiya Nourian<sup>5</sup>, Hamid Reza BahramiTaghanaki<sup>6</sup>, Gholam Reza Mohammadi<sup>7</sup>, Mohammad Rashtibaf<sup>8</sup>, Kiarash Ghazvini<sup>\*1, 2</sup>

## Abstract

**Background:** Tuberculosis (TB) still remains endemic worldwide making epidemiological studies essential to mitigating efforts implicated in identifying its source, controlling, and preventing the spread of dangerous strains amongst humans such as *Mycobacterium tuberculosis (Mtb)*.

*Methods:* In this study, we sought to determine the 6 Mycobacterial Interspersed Repetitive Unit-Variable-Number Tandem Repeat (MIRU-VNTR) loci with high discriminatory powers for *Mtb* genotyping as well as the loci with the highest and the lowest discriminatory powers for MIRU-VNTR. To conduct our search, we used several databases such as science direct, Embase (Elsevier), Web of Science, Scopus and Medline via PubMed. Searches were performed using key words including: *Mycobacterium tuberculosis*, MIRU-VNTR, Allele diversity, Genetic diversity and human patient. Finally, 56 articles were selected after filtering out titles, abstracts and full texts.

*Results:* Loci with high discriminatory powers included MIRU10 and MIRU26, while MIRU2, MIRU20, MIRU24 and ETRD had poor discriminatory powers. According to previous data in the literature, the loci MIRU10, MIRU26, MIRU40, QUB 26, QUB 11b and Mtub21 have high discriminatory powers.

*Conclusions:* Therefore, these loci recommended for genotyping *Mtb* to save time and cost and to ensure the production of reliable results.

Keywords: Discriminatory power, Genotyping, MIRU-VNTR, Mycobacterium tuberculosis.

## Introduction

In spite of recent efforts to control and eliminate TB, this highly infectious disease still remains the second leading cause of death worldwide (1, 2). In 2018, WHO predicted that 10 million patients (ranging

from 9 to 11.1 million) were stricken by TB. *Mtb* is a member of the TB complex which causes TB and can be transmitted via aerosolization of bodily fluids from coughing, sneezing or speaking (2).

<sup>1:</sup> Antimicrobial Resistance Research Center, Mashhad University of Medical Sciences, Mashhad, Iran.

<sup>2:</sup> Department of Microbiology and Virology, School of Medicine, Mashhad University of Medical Sciences, Mashhad, Iran.

<sup>3:</sup> Department of Microbiology, Razi Vaccine and Serum Research Institute (RVSRI), Agricultural Research, Education and Extension Organization (AREEO), Karaj, Iran.

<sup>4:</sup> PPD Tuberculin Department, Razi Vaccine and Serum Research Institute, (RVSRI), Agricultural Research, Education and Extension Organization (AREEO), Karaj, Iran.

<sup>5:</sup> Doctor of Veterinary Medicine, Graduate Student of School of Veterinary Medicine, Ferdowsi University of Mashhad, Mashhad, Iran.

<sup>6:</sup> School of Persian Medicine, Mashhad University of Medical Sciences, Mashhad, Iran.

<sup>7:</sup> Department of Clinical Sciences, School of Veterinary Medicine, Ferdowsi University of Mashhad, Mashhad - Iran.

<sup>8:</sup> Deputy of Veterinary administration of Khorasan Razavi Province, Mashhad, Iran.

<sup>\*</sup>Corresponding author: Ghazvini Kiarash; Tel: +98 51 38012589, Fax: +98 51 38409612; E-mail: Ghazvinik@mums.ac.ir. Received: 17 Jun, 2019; Accepted: 30 Jun, 2019

To effectively control TB, preventive policies based on transmission routes are required. Molecular epidemiology of *Mtb* involves monitoring special strains like multi-drug-resistant (MDR) *Mtb* during periods of increasing prevalence, inspecting regions of latest and potential outbreaks, locating the source and route of transmission and transmission gene sequence, discovering hidden strains and tracking circulating immigrant strains (3).

Various molecular methods are used in epidemiological studies about Mtb strains. Each method has a particular specificity and sensitivity. Some of these approaches include IS6110, IS6110restriction fragment length polymorphism (RFLP), MIRU-VNTR, Spoligotyping, whole genome sequencing (WGS), Random Amplification of Polymorphic DNA PCR (RAPD\_PCR), Repetitive element sequence-based PCR (rep-PCR), Pulsedfield gel electrophoresis (PFGE), Next Generation Sequencing (NGS) and finally, a combination of two or more techniques listed above will be applied. Among current typing methods, a test has to be chosen according to its feasibility, cost-benefit and discriminatory powers (4-7)

In a survey comparing PFGE, 24 locus MIRU-VNTR and IS6110-RFLP, it was revealed that the 24 locus MIRU-VNTR method was the preferred method due to a high power of discrimination and time management during epidemiological investigations (7).

This study was performed to review published applications of the Mycobacterial Interspersed Repetitive Unit-Variable-Number Tandem Repeat (MIRU-VNTR) method in *Mtb* genotyping and to introduce the best 6 loci for MIRU-VNTR in typing *Mtb* isolated from human patients along with determining the loci with highest and lowest discriminatory powers for MIRU- VNTR.

## Materials and methods

To identify the best loci of the 6 loci in MIRU-VNTR method for *Mtb* genotyping, the literature search was performed using several databases including science direct, Embase (Elsevier), Web of Science, Scopus, ISC and Medline via PubMed. Chosen keywords were: *Mtb*, MIRU–VNTR, Allele diversity, genetic diversity and human patient. Inclusion and exclusion criteria were determined by the following: 1. Isolation of *Mtb* from human patients.

2. Investigating the genetic diversity among *Mtb* just based on MIRU-VNTR.

3. Excluding the studies where lineage determination was based on MIRU-VNTR where Allele diversity "*h*" was not measured for each locus.

4- Excluding the studies in which MIRU-VNTR ability in cluster analysis and Hunter-Gaston discriminatory index (HGDI) for this method was investigated and h was not estimated for each of their locus.

Several articles were excluded from our study since allele diversity (h) was not mentioned for each locus. Screening the articles was done in 3 steps: 1. Title screening, 2. Abstract evaluation, 3. Full text evaluation based on these criteria.

#### Results

A total number of 228 articles were found collectively amongst the databases. As the title screening was performed, 90 articles were removed. Abstract screening resulted in 82 more studies to be omitted during the search. Finally, after full text screening, 56 articles remained. In the remaining articles, genotyping for *Mtb* using MIRU-VNTR was investigated from 2002 to 2019. Allele diversity (h) was evaluated amongst the 56 articles for each locus separately (Table 1).

*Mycobacterium tuberculosis* genotyping was performed on 56 studies using the MIRU-VNTR technique; 39 of which were conducted in Asia, seven in America, six in Africa, three in Europe and one in a different country. The location and number of studies are shown in Figure 1. Each individual study employed a different number of loci. The results revealed that MIRU10 and MIRU26 had the highest discriminatory powers while MIRU2, MIRU20, MIRU24 and ETRD had the lowest discriminatory powers, respectively.

Table 2 shows both the number of studies in which MIRU10, MIRU26, QUB26, MIRU40, QUB11b and Mtub21 was reported to be the loci with the highest discriminatory powers (h>0.6), including the range of *h* for the remaining loci.

MIRU2 and MIRU20 (each in 21 studies), MIRU24 (17 studies) and ETRD (13 studies) were suggested as the loci with the lowest discriminatory powers (h<0.3).

No.	Authors reference	year	Geographical region	Continent	Locus of MIRU-VNTR	Numbers of locus	High power	Low power
.1	Cowan et al (8)	2002	United States (Michigan)	America	ETR D,E MIRU2,10,16,20,23,24,26,27,39 ,40	12	MIRU40	MIRU2,2 0
.2	Sola et al (9)	2003	Different regions such as; USA, Sicily, Guadeloupe and Russia	Thailand,	ETR A-E MIRU2,10,16,20,23,24,26,27,39 ,40	15	EIRA, MIRU40,26,10	MIRU2,2 0,27
.3	Sun et al (10)	2004	Singapore	Asia	ETR D,E MIRU2,10,16,20,23,24,26,27,39 ,40	12	MIRU26,10, EIRE	MIRU2,2 0
.4	Kremer et al (11)	2005	China	Asia	ETR A-E MIRU10,16,26,39,40 QUB 11a,11b,26,1895	14	QUB 11b,11a MIRU10	MIRU16 EIRC
.5	Kovalev et al (12)	2005	Russian Federation	Asia	ETR D,E MIRU2,10,16,20,23,24,26,27,39 ,40	12	MIRU26, ETRE	MIRU24 27
.6	Asgarzadeet al (13)	2007	Iran	Asia	ETR D,E MIRU2,10,16,20,23,24,26,27,39 ,40	12	MIRU26,40	MIRU16 39
.7	Çavuşoğlu et al (14)	2007	Turkey	Asia	ETR D,E MIRU2,10,16,20,23,24,26,27,39 ,40	12	MIRU16,40,26	MIRU24 27
.8	Maes et al (15)	2008	Venezuela	America	ETR A-E MIRU2,10,16,20,23,24,26,27,39 ,40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	MIRU40,26, EIRB	MIRU20 2,24
9	Alonso-Rodríguez et al (16)	2008	Spain	Europe	EIRA,C-E MIRU10,16,26,40 Mtub4,21,30,39 QUB 11b,26,4156	15	QUB, 26,11b, MIRU40, 10	EIRD
0	Yunetal(17)	2009	Korea	Asia	EIRA-F MIRU2,10,16,20,23,24,26,27,39 ,40	6	MIRU26, ETRE,F	MIRU24 20 EIRD
1	Stavnum et al (18)	2009	South Africa	Africa	EIRA,C-E MIRU10,16,26,40 Mtub4,21,30,39 QUB 11b,26,4156	15	QUB 11b	EIRD
2	Shamputa et al (19)	2010	South Korea	Asia	ETR A-E MIRU2,10,1620,23,24,26,27,39 40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	QUB 11b,26, Mtub4	MIRU2,2 4,23 EIRB,D
.3	Noguti et al (20)	2010	Brazil	America	EIRD,E MIRU2,10,1620,23,24,26,27,39 ,40	12	MIRU40,23,10	MIRU24 39 ETRD
4	Jafarian et al (21)	2010	Iran	Asia	EIR D,E MIRU2,10,1620,23,24,26,27,39 ,40	12	MIRU 26, 10,16	MIRU2, 24 EIRD
5	Zhang et al (22)	2011	Cambodia	Asia	ETR A-E MIRU2,10,16,20,23,24,26,27,39 40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	ETRD,Mtub 39, QUB26	Mtub 34, MIRU2,2 0
6	Asgarzad et al (23)	2011	Iran	Asia	EIR A-E MIRU2,10,1620,23,24,26,27,39 ,40	15	MIRU10,26,40	MIRU39 2,4
7	Bidovec-Stojkovi et al (24)	2011	Slovenia	Europe	ETR A-E MIRU2,10,1620,23,24,26,27,39 40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	QUB 26, 11b, MIRU40,10	MIRU24 39
8	Cerezo et al (25)	2012	Colombia	America	ETR D,E MIRU2,10,16,20,23,24,26,27,39 ,40	12	MIRU10,40	MIRU24 39
.9	Chatterjee et al (26)	2013	India	Asia	EIR D,E MIRU2,10,1620,23,24,26,27,39 ,40	12	MIRU26, 10	MIRU2,2 0

Table 1. Fifty-six studies after full-text evaluation that MIRU-VNTR were used.

.20	Zamani et al (27)	2013	Iran	Asia	ETRA,C-E MIRU10,16,26,40 Mtub4,21,30,39 QUB 11b,26,4156	15	MIRU16 EIRA	MIRU26, Mtub 21, 30, 39 QUB4156
.21	Joseph et al (28)	2013	India	Asia	EIRA,C-E MIRU10,1626,40 Mitub4,21,30,39 QUB 11b,26,4156	15	ETRA,B MIRU40	MIRU2,2 0
.22	Yasmin et al (29)	2014	Pakistan	Asia	EIR A-E MIRU2,10,16,20,23,24,26,27,39 40 Mub4,21,29,30,34,39 QUB 11b,26,4156	24	MIRU26 QUB26 MIRU10	MIRU2,2 0,27,24
.23	Alietal (30)	2014	Pakistan	Asia	ETRA,C-E MIRU10,16,26,40 Mub4,21,30,39 QUB 11b,26,4156	15	QUb26 ,MIRU10,26	EIRD
.24	Chaoui et al (31)	2014	Maracco	Africa	ETR D,E MIRU2,10,16,20,23,24,26,27,39 ,40	12	MIRU40,23,10	MIRU24, 39
.25	Zheng et al (32)	2014	China	Asia	FIR A-E MIRU2,10,16,20,23,24,26,27,39 40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	QUB11b,26 Mtub21 MIRU26	MIRU24 Mtub 34
.26	Vasconcellos et al (33)	2014	Brazil	America	EIR A-E MIRU2,10,16,20,23,24,26,27,39 40 Mub4,21,29,30,34,39 QUB 11b,26,4156	24	QUB 4156, 11b MIRU10	MIRU24, 39
.27	Rindi et al (34)	2014	Italy	Ешоре	EIRA,D,E MIRU10,16,26,40 Mtub21 QUB 11b, 26 VNIR 42,43,47,52,53	15	QUB 26 MIRU 40 QUB 11b	MIRU04 ETRE
.28	Bouklata et al (35)	2015	Maracco	Africa	EIR A-E MIRU2,10,16,20,23,24,26,27,39 ,40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	QUB 26, MIRU40,26	MIRU20, 27
.29	Devictal (36)	2015	India	Asia	EIR A-E MIRU2,10,16,20,23,24,26,27,39 40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	Qub26, 11b MIRU10 QUB 4156 MIRU26 Mtub 21	MIRU2,2 0,27
.30	Zamani et al (37)	2016	Iran	Asia	EIRA,C-E MIRU10,16,26,40 Mtub4,21,30,39 QUB 11b,26,4156	15	MIRU40	MIRU 10 QUB 4156
.31	Hoza et al (38)	2016	Tanzania	Africa	EIRAC-E MIRU2,10,16,20,23,24,26,27,39 40 Mtub4,21,29,30,39 QUB 11b,26,4156	22	MIRU26,10,16 ETRAE QUB 26,4156	MIRU27, 2,20
.32	Cheng et al (39)	2016	China	Asia	EIR A-E MIRU10,16,20,26,27,39,40 Mtub31 QUB 11a,11b	15	QUB11a	ETRB,C
.33	Bhembe et al (40)	2017	South Africa (Eastern Cape)	Africa	ETR D,E MIRU2,10,16,20,23,24,26,27,39 ,40	12	ETRE, MIRU27,24	MIRU40
.34	Zhang et al (41)	2017	China	Asia	ETR A-E MIRU 10,16,23,26,27,39,40 Mtub21,30,39	15	Mtub 21, MIRU 26,10	ETRC,B
.35	Liuetal (42)	2017	China	Asia	ETRA-E MIRU10,16,23,26,27,39,40 Mtub21,30,39	15	MIRU26 Mtub21	MIRU27, 23 EIRD
.36	Pasechnik et al (43)	2017	West Siberia	Asia	ETRA-D MIRU2,10,16,20,23,24,26,27,39 ,40	15	MIRU26	MIRU24 EIRB

.37	Pan et al (44)	2017	China	Asia	EIRA,D,E MIRU10,16,26,39,40 Mtub4,21,24,30,39 QUB11a,11b,18,26,3232,1895,4 156 VNIR3820,4120	22	VNIR3820 QUB3232	QUB4156 Mtub 24
.38	Khosravi et al (45)	2017	Iran	Asia	EIRD,E MIRU2,10,16,20,23,24,26,27,39 40	12	MIRU 10,26	MIRU2,2 0
.39	Ravansalar et al (46)	2017	Iran	Asia	EIRA-F MIRU10,16,26,39,40 QUB 11b	12	MIRU10,26 ETRF	EIRD
.40	Shah et al (47)	2017	Nepal	Asia	ETRA-F MIRU2,10,16,20,23,24,26,27,39 40 Mtub21,30,39 QUB 11b,11b,26,4156	24	QUB 26 MIRU10	EIR B,C
.41	Rascaha et al (48)	2017	Madagascar	Africa	ETRA,C-E MIRU10,16,26,40 Mtub4,21,30,39 QUB 11b,26,4156	15	MIRU26 QUB 11b Mtub 21 QUB 26	ETRD,C Mtub4
.42	Chen et al (49)	2017	Asian countries (Cambodia, Singapore and Taiwan)	Asia	ETRA-E MIRU2,10,16,20,23,24,26,27,39 ,40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	Mtub21 QUB 11b	MIRU2 QUB 4156
.43	Lietal(50)	2018	China	Asia	EIRA-E MIRU2,10,16,20,23,24,26,27,39 ,40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	Mtub4 MIRU40,10	Mtub21 MIRU27
.44	Xuetal(51)	2018	China	Asia	ETRA,C-E MIRU10,16,26,40 Mtub4,21,30,39 QUB 11b,26,4156	15	QUB 11b Mtub21 MIRU26	EIRC MIRU16 QUB 4156
.45	Esteves et al (52)	2018	Brazil	America	EIRA-E MIRU2,10,16,20,23,24,26,27,39 ,40 QUB11,26 VNIR42,1955,47,52,53,49	24	QUB26 QUB11 VNIR42	MIRU39, 24 EIRD
.46	Augusto et al (6)	2018	Brazil	America	ETRD,E MIRU2,10,16,20,23,24,26,27,39 ,40	12	MIRU16, 10,26	MIRU20, 24
.47	Riyahi Zaniani et al (53)	2018	lan	Asia	ETRA-E MIRU2,10,16,20,23,24,26,27,39 40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	MIRU 10 QUB 26 Mtub 4,34	EIRD, MIRU 20, Mtub 29
.48	Azimi et al (54)	2018	lian	Asia	ETRA,C-E MIRU10,16,26,40 Mtub4,21,30,39 QUB11b,26,4156	15	MIRU26, 10 Mtub 21 QUB 26	EIRD,E
.49	Liletal (3)	2018	China	Asia	EIRA-E MIRU16,23,26,27,39,40 Mtub21,30,39	15	EIRE, MIRU10, 26,39,40 Mtub 21	EIRB,C MIRU16, 23
.50	Mansoori et al (55)	2018	Iran	Asia	ETRA-E MIRU2,10,16,20,23,24,26,27,39 ,40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	MIRU 10, 16, 26,	MIRU2, 20 EIRD
.51	Chawla et al (56)	2018	India	Asia	EIRD,E MIRU2,10,16,20,23,24,26,27,39 40	12	MIRU 39,10,26	MIRU 2,20
.52	Shietal (57)	2018	China	Asia	ETRA-F MIRU2,10,16,20,23,24,26,27,39 ,40 Mtub4,21,29,30,34,38,39 QUB 11b,26,4156	26	QUB 11b Mtub 21 MIRU26	MIRU24, 2,20

.53	Eietal (58)	2018	Myanmar	Asia	EIRA-E MIRU2,10,16,20,23,24,26,27,39 ,40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	QUB 26 QUB 11b MIRU26	Mtu34 MIRU20
.54	Weenasekera et al (59)	2019	Sri Lanka	Asia	EIRA-E MIRU2,10,16,20,23,24,26,27,39 ,40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	Mtub 21,39 QUB 11b	MIRU20, 2,16
.55	Zarzour et al (60)	2019	Syria	Asia	EIRA-E MIRU2,10,16,20,23,24,26,27,39 ,40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	QUB 26 MIRU10,26	MIRU24, 20, Mtub 29
.56	Lietal(61)	2019	China	Asia	EIRA-E MIRU2,10,1620,23,24,26,27,39 ,40 Mtub4,21,29,30,34,39 QUB 11b,26,4156	24	QUB11b MIRU 26 Mtub21	MIRU2, 20,24

 Table 2. Six MIRU-VNTR loci for Mycobacterium tuberculosis based on the number of studies and the reported allele diversity (h) range for each locus.

locus	Number of studies	h range
MIRU10	28	0.61 to 0.82
MIRU26	32	0.61 to 0.81
QUB26	18	0.604 to 0.89
MIRU40	17	0.604 to 0.76
QUB11b	17	0.72 to 0.84
Mtub21	12	0.64 to 0.83



Fig. 1. Location and numbers of studies was shown in the world from 2002 to 2019. Two studies were performed in different regions: Studies No. 2 and 42.

#### Discussion

An alarming 62% of emerging TB cases occurred in the South-East Asia and Western Pacific regions, followed by Africa, which accounts for 25% of new cases. Countries such as India, China, Indonesia, the Philippines, Pakistan, Bangladesh, Nigeria and South Africa hold a prevalence rate greater than 60% for TB. It is believed that the surge in TB research in Asia could be that TB is highly prevalent in this continent based on 2018 WHO report (2). Of the 56 studies extracted, 39 studies were conducted within Asia. More specifically, 11 were performed in China, ten in Iran, four in India, two in Pakistan, two in Korea and only one in Nepal, Singapore, Myanmar, Turkey, Cambodia, Sri Lanka, Syria, Russia, Siberia, Taiwan, and Cambodia. Among the 39 studies conducted in Asia, MIRU26 and MIRU10 was reported having high discriminatory power loci. Among the 7 studies conducted in the Western Hemisphere, four studies were carried out in Brazil, and one study was performed in Michigan, Venezuela and Colombia. Both MIRU40 and MIRU10 had high power discriminatory power loci in these ancients.

Among the six studies performed in Africa, four studies were conducted in South Africa and Morocco, respectively. One study, performed in Madagascar and Tanzania, reported that QUB26 and MIRU26 had high discriminatory power loci in this continent. Several investigations in this study used PCR-based techniques such as spacer (spoligotyping) oligonucleotide typing and mycobacterial interspersed repetitive unitsvariable-number of tandem repeats (MIRUanalyses. Although spoligotyping VNTR) benefits from genetic diversity, it can minimize Mtb clonal diversity. This method uses one direct repeat (DR) which incorporates identical alternate and variable spacers and the results are represented as a single digit pattern. The basis of MIRU-VNTR depends on the (variable) number of tandem repeat elements called mycobacterial interspersed repetitive units (MIRU). The results are represented as a code, and offers a low discriminatory power when used alone. The MIRU-VNTR method analyzes DNA segments

which involves tandem repeat sequences and the copy number which differs amongst various strains. This method depends on PCR efficiency, specifically the quantity of repeats which is based on the size of the amplified product. The results are illustrated as characters that range from 15 -24 characters, in which each character represents the number of repeats at a single locus (3, 62).

These results are evaluated in comparison to a strain database on the web-based tool MIRU-VNTR plus. Therefore, MIRU-VNTR typing is considered the gold-standard for genotypic analysis of *Mtb* (7, 9). The discriminatory power of these methods was found to be different which was determined by the Hunter and Gaston Index factor .Several studies reported different Hunter-Gaston Index for MIRU-VNTR ranging from 0.951 to 0.999 (10, 13, 45, 47, 48, 53-56).

The allele diversity index (h) is used to describe the discriminatory power of MIRU-VNTR loci. If the index is greater than 0.6 (h>0.6), the discriminatory power of the locus is high. If the index lies between 0.3 and 0.6 (0.3<h <0.6), the locus has medium discriminatory power, however, if the index is less than 0.3 (0.3>h), the discriminatory power is considered weak. Our results suggest that among the 24 defined loci for MIRU-VNTR introduced by the MIRU-VNTR plus database, the MIRU26, MIRU10, MIRU40, QUB26, QUB11b and Mtub21 loci had the highest discriminatory powers, in contrast to the MIRU2, MIRU20, MIRU24 and ETRD loci which yielded low discriminatory powers.

*Mycobacterium bovis* (M. *bovis*) is the causative agent of TB in humans. When comparing the discriminatory powers of loci between *Mtb* and M. *bovis*, the loci QUB 11b and QUB 3232 have the highest discriminatory powers (for M. *bovis*), whereas ETRD (for both strains) and MIRU10 (for M. *bovis*) had the lowest discriminatory powers. Therefore, it can be concluded from our study that QUB 11b in both *Mtb* and *M. bovis* has a high discriminatory power while ETRD for both strains is considered a low power locus. However, a difference exists between the two

strains in the MIRU10 locus which has a high discriminatory power in *Mtb* but low in *M. bovis* (63).

While the 24 loci MIRU-VNTR has been introduced as the best typing method for *Mtb* based on multiple comparisons between various molecular techniques, it is, therefore, highly recommended to include the following loci, QUB26, QUB11b, MIRU10, MIRU26, MIRU40 and Mtub21, provided that MIRU-VNTR is done with less than 24 loci, in order to obtain the best results when genotyping *Mtb* isolates. In this regard, data extraction becomes inexpensive and time efficient.

Among typing methods, MIRU-VNTR is considered to be one of the best. Further, the 6

#### References

1. Kaufmann SH, Schaible UE. 100th anniversary of Robert Koch's Nobel Prize for the discovery of the tubercle bacillus. Trends in microbiology. 2005;13(10):469-75.

2. WHO. Global tuberculosis report 2018. World Health Organization Geneva, 2018.

3. Lil B, Ma Y, Liu H, Shenl X, Ma B, Jiang M, et al. Molecular typing of Mycobacterium tuberculosis isolates from Qinghai province of northwest china by spoligotyping and 15-locus MIRUVNTR. Journal of Microbiology and Biotechnology Reports. 2018;2(1):1-4.

4. Ravansalar H, Tadayon K, Ghazvini K. Molecular typing methods used in studies of Mycobacterium tuberculosis in Iran: a systematic review. Iranian journal of microbiology. 2016;8(5):338.

5. Ei PW, Aung WW, Lee JS, Choi G-E, Chang CL. Molecular strain typing of Mycobacterium tuberculosis: a review of frequently used methods. Journal of Korean medical science. 2016;31(11):1673-83.

6. Augusto CJ, Carvalho WdS, Almeida INd, Figueiredo LJdA, Dantas NGT, Suffys PN, et al. Comparative study of RFLP-IS6110 and MIRU-VNTR from Mycobacterium tuberculosis isolated in the state of Minas Gerais, Brazil. brazilian journal of microbiology. 2018;49(3):641-6.

7. Jeon S, Lim N, Park S, Park M, Kim S. Comparison of PFGE, IS6110-RFLP, and 24-Locus

loci including MIRU10, MIRU26, MIRU40, QUB 26, QUB 11b and Mtub21, all of which have high discriminatory powers, are recommended in *Mtb* genotyping to save time and cost. Indeed, epidemiological studies are critical for disease surveillance of TB as they enable us to track circulating strains on a global scale, as well as for identifying important risk factors necessary for implementing control measures in vulnerable populations worldwide.

## Acknowledgment

We thank Antimicrobial Resistance Research Center and Mashhad University of Medical Sciences The authors of this study declare no conflicts of interest.

MIRU-VNTR for Molecular Epidemiologic Typing of Mycobacterium tuberculosis Isolates with Known Epidemic Connections. Journal of microbiology and biotechnology. 2018;28(2):338-46.

8. Cowan LS, Mosher L, Diem L, Massey JP, Crawford JT. Variable-number tandem repeat typing of Mycobacterium tuberculosis isolates with low copy numbers of IS6110 by using mycobacterial interspersed repetitive units. Journal of clinical microbiology. 2002;40(5):1592-602.

9. Sola C, Filliol I, Legrand E, Lesjean S, Locht C, Supply P, et al. Genotyping of the Mycobacterium tuberculosis complex using MIRUs: association with VNTR and spoligotyping for molecular epidemiology and evolutionary genetics. Infection, Genetics and Evolution. 2003;3(2):125-33.

10. Sun Y-J, Bellamy R, Lee AS, Ng ST, Ravindran S, Wong S-Y, et al. Use of mycobacterial interspersed repetitive unit-variable-number tandem repeat typing to examine genetic diversity of Mycobacterium tuberculosis in Singapore. Journal of clinical microbiology. 2004;42(5):1986-93.

11. Kremer K, Arnold C, Cataldi A, Gutiérrez MC, Haas WH, Panaiotov S, et al. Discriminatory power and reproducibility of novel DNA typing methods for Mycobacterium tuberculosis complex strains. Journal of clinical microbiology. 2005;43(11):5628-38.

12. Kovalev S, Kamaev E, Kravchenko M, Kurepina N, Skorniakov S. Genetic analysis of

Mycobacterium tuberculosis strains isolated in Ural region, Russian Federation, by MIRU-VNTR genotyping. The International Journal of Tuberculosis and Lung Disease. 2005;9(7):746-52.

13. Asgharzadeh M, Khakpour M, Salehi TZ, Kafil HS. Use of mycobacterial interspersed repetitive unit-variable-number tandem repeat typing to study Mycobacterium tuberculosis isolates from East Azarbaijan province of Iran. Pakistan journal of biological sciences. 2007;10(21):3769-77.

14. Cavuşoğlu C, Karataş E, Soeyler I. Genotyping of rifampin-resistant Mycobacterium tuberculosis isolates by mycobacterial interspersed repetitive unit-variable number tandem repeat (MIRU-VNTR) analysis. Mikrobiyoloji bulteni. 2007;41(3):385-93.

15. Maes M, Kremer K, van Soolingen D, Takiff H, de waeerd JH. 24-locus MIRU-VNTR genotyping is a useful tool to study the molecular epidemiology of tuberculosis among Warao Amerindians in Venezuela. Tuberculosis (Edinburgh, Scotland). 2008;88(5):490-4.

16. Alonso-Rodríguez N, Martínez-Lirola M, Herránz M, Sanchez-Benitez M, Barroso P, Bouza E, et al. Evaluation of the new advanced 15-loci MIRU-VNTR genotyping tool in Mycobacterium tuberculosis molecular epidemiology studies. BMC microbiology. 2008;8(34):1-9.

17. Yun KW, Song EJ, Choi GE, Hwang IK, Lee EY, Chang CL. Strain typing of Mycobacterium tuberculosis isolates from Korea by mycobacterial interspersed repetitive units-variable number of tandem repeats. The Korean journal of laboratory medicine. 2009;29(4):314-9.

18. Stavrum R, Mphahlele M, Øvreås K, Muthivhi T, Fourie PB, Weyer K, et al. High diversity of Mycobacterium tuberculosis genotypes in South Africa and preponderance of mixed infections among ST53 isolates. Journal of clinical microbiology. 2009;47(6):1848-56.

19. Shamputa IC, Lee J, Allix-Béguec C, Cho E-J, Rajan V, Lee EG, et al. Genetic diversity of Mycobacterium tuberculosis isolates from a tertiary care tuberculosis hospital in South Korea. Journal of clinical microbiology. 2010;48(2):387-94.

20. Noguti EN, Leite CQF, Malaspina AC, Santos ACB, Hirata RDC, Hirata MH, et al. Genotyping of Mycobacterium tuberculosis isolates from alowendemic setting in northwestern state of Paraná in Southern Brazil. Memorias do Instituto Oswaldo Cruz. 2010;105(6):779-85.

21. Jafarian M, Aghali-Merza M, Farnia P, Ahmadi M, Masjedi MR, Velayati AA. Synchronous comparison of Mycobacterium tuberculosis epidemiology strains by "MIRU-VNTR" and "MIRU-VNTR and spoligotyping" technique. Avicenna journal of medical biotechnology. 2010;2(3):145-52.

22. Zhang J, Heng S, Le Moullec S, Refregier G, Gicquel B, Sola C, et al. A first assessment of the genetic diversity of Mycobacterium tuberculosis complex in Cambodia. BMC infectious diseases. 2011;11(42):1-7.

23. Asgharzadeh M, Kafil HS, Roudsary AA, Hanifi GR. Tuberculosis transmission in Northwest of Iran: using MIRU-VNTR, ETR-VNTR and IS6110-RFLP methods. Infection, Genetics and Evolution. 2011;11(1):124-31.

24. Bidovec-Stojkovic U, Zolnir-Dovc M, Supply P. One year nationwide evaluation of 24-locus MIRU-VNTR genotyping on Slovenian Mycobacterium tuberculosis isolates. Respiratory medicine. 2011;105:S67-S73.

25. Cerezo I, Jiménez Y, Hernandez J, Zozio T, Murcia MI, Rastogi N. A first insight on the population structure of Mycobacterium tuberculosis complex as studied by spoligotyping and MIRU-VNTRs in Bogotá, Colombia. Infection, Genetics and Evolution. 2012;12(4):657-63.

26. Chatterjee A, Mistry N. MIRU–VNTR profiles of three major Mycobacterium tuberculosis spoligotypes found in western India. Tuberculosis. 2013;93(2):250-6.

27. Zamani S, Aflaki M, Fooladi AAI, Darban-Sarokhalil D, Bameri Z, Khazaee S, et al. MIRU-VNTR analysis of the Mycobacterium tuberculosis isolates from three provinces of Iran. Scandinavian journal of infectious diseases. 2013;45(2):124-30.

28. Joseph BV, Soman S, Hill V, Kumar RA, Rastogi N, Mundayoor S. Efficient discrimination by MIRU-VNTRs of Mycobacterium tuberculosis clinical isolates belonging to the predominant SIT11/EAI3-IND ancestral genotypic lineage in Kerala, India. International journal of mycobacteriology. 2013;2(4):244-7.

29. Yasmin M, Gomgnimbou MK, Siddiqui RT, Refrégier G, Sola C. Multi-drug resistant Mycobacterium tuberculosis complex genetic diversity and clues on recent transmission in Punjab, Pakistan. Infection, Genetics and Evolution. 2014;27:6-14.

30. Ali A, Hasan Z, Jafri S, Inayat R, Hasan R. Mycobacterium tuberculosis Central Asian Strain (CAS) lineage strains in Pakistan reveal lower diversity of MIRU loci than other strains. International journal of mycobacteriology. 2014;3(2):108-16.

31. Chaoui I, Zozio T, Lahlou O, Sabouni R, Abid M, El Aouad R, et al. Contribution of spoligotyping and MIRU-VNTRs to characterize prevalent Mycobacterium tuberculosis genotypes infecting tuberculosis patients in Morocco. Infection, Genetics and Evolution. 2014;21:463-71.

32. Zheng C, Zhao Y, Zhu G, Li S, Sun H, Feng Q, et al. Suitability of IS6110-RFLP and MIRU-VNTR for differentiating spoligotyped drug-resistant Mycobacterium tuberculosis clinical isolates from Sichuan in China. BioMed research international. 2014;2014:1-13.

33. Vasconcellos S, Acosta C, Gomes L, Conceiçao E, Lima K. Strain Classification of Mycobacterium tuberculosis Isolates in Brazil Based on Genotypes Obtained by Spoligotyping, Mycobacterial Interspersed Repetitive Unit Typing and the Presence of Large Sequence and Single Nucleotide Polymorphism. PloS one. 2014;9(10):e107747.

34. Rindi L, Medici C, Bimbi N, Buzzigoli A, Lari N, Garzelli C. Genomic Variability of Mycobacterium tuberculosis Strains of the Euro-American Lineage Based on Large Sequence Deletions and 15-Locus MIRU-VNTR Polymorphism. PloS one. 2014;9(11):e114676.

35. Bouklata N, Supply P, Jaouhari S, Charof R, Seghrouchni F, Sadki K ea. Molecular Typing of Mycobacterium tuberculosis complex by 24-locus based MIRU-VNTR typing in conjunction with spoligotyping to assess genetic diversity of strains circulating in Morocco. PloS one. 2015;10(8):e0135695.

36. Devi KR, Bhutia R, Bhowmick S, Mukherjee K, Mahanta J, Narain K. Genetic diversity of Mycobacterium tuberculosis isolates from Assam, India: dominance of Beijing family and discovery of two new clades related to CAS1\_Delhi and EAI family based on spoligotyping and MIRU-VNTR typing. PloS one. 2015;10(12):e0145860. 37. Zamani S, Haeili M, Nasiri MJ, Fooladi I, Ali A, Javadpour S, et al. Genotyping of Mycobacterium tuberculosis isolates from Hormozgan province of Iran based on 15-locus MIRU-VNTR and spoligotyping. International journal of bacteriology. 2016;2016:1-8.

38. Hoza A, Mfinanga S, Moser I, König B. Molecular characterization of Mycobacterium tuberculosis isolates from Tanga, Tanzania: First insight of MIRU-VNTR and microarray-based spoligotyping in a high burden country. Tuberculosis. 2016;98:116-24.

39. Cheng X-f, Jiang C, Zhang M, Xia D, Chu L-l, Wen Y-f, et al. Mycobacterial interspersed repetitive unit can predict drug resistance of Mycobacterium tuberculosis in China. Frontiers in Microbiology. 2016;7:378.

40. Bhembe NL, Nwodo UU, Okoh AI, Obi CL, Mabinya LV, Green E. Clonality and genetic profiles of drug-resistant Mycobacterium tuberculosis in the Eastern Cape Province, South Africa. MicrobiologyOpen. 2017:1-10.

41. Zhang H, Huang H, Liu C, Jia T, Zhang L, Zhou D, et al. Genotyping and drug-resistance epidemiology of mycobacterium tuberculosis in Xuzhou, China. International Journal of Clinical and Experimental Pathology. 2017;10(9):9675-86.

42. Liu J, Li J, Liu J, Zhao X, Lian L, Liu H, et al. Genotypic diversity of mycobacterium tuberculosis clinical isolates in the multiethnic area of the Xinjiang Uygur autonomous region in China. BioMed research international. 2017;2017:1-8.

43. Pasechnik O, Dymova MA, Stasenko VL, Blokh AI, Tatarintseva MP, Kolesnikova LP, et al. Molecular & genetic characteristics of Mycobacterium tuberculosis strains circulating in the southern part of West Siberia. The Indian journal of medical research. 2017;146(1):49-55.

44. Pan X-L, Zhang C-L, Nakajima C, Fu J, Shao C-X, Zhao L-N, et al. A quantitative and efficient approach to select MIRU–VNTR loci based on accumulation of the percentage differences of strains for discriminating divergent Mycobacterium tuberculosis sublineages. Emerging microbes & infections. 2017;6(7):e68.

45. Khosravi AD, Shahraki AH, Dezfuli SK, Hashemzadeh M, Goodarzi H, Mohajeri P. Genetic diversity of multidrug-resistant Mycobacterium tuberculosis strains isolated from tuberculosis patients in Iran using MIRU-VNTR technique. Kaohsiung Journal Medical Science. 2017;33(11):550-7.

46. Ravansalar H, Tadayon K, Mosavari N, Derakhshan M, Ghazvini k. Genetic Diversity of Mycobacterium tuberculosis Complex Isolated from Patients in the Northeast of Iran by MIRU-VNTR and Spoligotyping Jundishapur Journal of Microbiology. 2017;10(4):e39568.

47. Shaha Y, Maharjana B, Thapaa J, Poudelc A, Diabd H. High diversity of multidrug-resistant Mycobacterium tuberculosis Central Asian Strain isolates in Nepal. International Journal of Infectious Diseases. 2017;63:13-20.

48. Rasoahanitralisoa R, Rakotosamimanana N, Stucki D, Sola C, Gagneux S, Razanamparany VR. Evaluation of spoligotyping, SNPs and customised MIRU-VNTR combination for genotyping Mycobacterium tuberculosis clinical isolates in Madagascar. PloS one. 2017;12(10):e0186088.

49. Chen Y-Y, Chang J-R, Huang W-F, Hsu C-H, Cheng H-Y, Sun J-R, et al. Genetic diversity of the Mycobacterium tuberculosis East African–Indian family in three tropical Asian countries. Journal of Microbiology, Immunology and Infection. 2017;50(6):886-92.

50. Li Y, Hu Y, Mansjö M, Zhao Q, Jiang W, Ghebremichael S, et al. The Epidemiological Significance and Temporal Stability of Mycobacterial Interspersed Repetitive Units-Variable Number of Tandem Repeats-Based Method Applied to Mycobacterium tuberculosis in China. International journal of environmental research and public health. 2018;15(4):782-93.

51. Xu G, Mao X, Wang J, Pan H. Clustering and recent transmission of Mycobacterium tuberculosis in a Chinese population. Infection and drug resistance. 2018;11:323-30.

52. Esteves LS, Dalla Costa ER, Vasconcellos SEG, Vargas A, Junior SLMF, Halon ML, et al. Genetic diversity of Mycobacterium tuberculosis isoniazid monoresistant and multidrug-resistant in Rio Grande do Sul, a tuberculosis high-burden state in Brazil. Tuberculosis. 2018;110:36-43.

53. Zaniani FR, Moghim S, Esfahani BN. Genetic Diversity of Drug-resistant Mycobacterium tuberculosis Isolates in Isfahan Province of Iran. Advanced Biomedical Research. 2018;7(23):1-6. 54. Azimi T, Nasiri MJ, Zamani S, Hashemi A, Goudarzi H, Fooladi AAI, et al. High genetic diversity among Mycobacterium tuberculosis strains in Tehran, Iran. Journal of Clinical Tuberculosis and Other Mycobacterial Diseases. 2018;11:1-6.

55. Mansoori N, Yaseri M, Vaziri F, Douraghi M. Genetic diversity of Mycobacterium tuberculosis complex isolates circulating in an area with high tuberculosis incidence: Using 24-locus MIRU-VNTR method. Tuberculosis. 2018;112:89-97.

56. Chawla K, Kumar A, Shenoy VP, Chauhan D, Sharma P. Strain diversity and relative transmission of Mycobacterium tuberculosis in south coastal Karnataka, India. The International Journal of Tuberculosis and Lung Disease. 2018;22(8):878-83.

57. Shi J, Zheng D, Zhu Y, Ma X, Wang S, Li H, et al. Role of MIRU-VNTR and spoligotyping in assessing the genetic diversity of Mycobacterium tuberculosis in Henan Province, China. BMC infectious diseases. 2018;18(447):1-12.

58. Ei PW, Aung WW, Nyunt WW, Swe TL, Aung ST, Htwe MM, et al. Evaluation Of Mycobacterial Interspersed Repetitive Unit-Variable Number Tandem Repeat Typing To Discriminate Mycobacterium Tuberculosis Strains From Myanma. The Southeast Asian journal of tropical medicine and public health 2018;49(2):276-84.

59. Weerasekera D, Pathirane H, Madegedara D, Dissanayake N, Thevanesam V, Magana-Arachchi DN. Evaluation of the 15 and 24-loci MIRU-VNTR genotyping tools with spoligotyping in the identification of Mycobacterium tuberculosis strains and their genetic diversity in molecular epidemiology studies. Infectious Diseases. 2019:1-10.

60. Zarzour H, Madania A, Ghoury I, Habous M. High-resolution genotyping of Mycobacterium tuberculosis isolates from syria using mycobacterial interspersed repetitive unitvariable-number tandem repeat. Biomedical and Biotechnology Research Journal (BBRJ). 2019;3(1):1-8.

61. Li D, Song Y, Yang P, Li X, Zhang AM, Xia X. Genetic diversity and drug resistance of Mycobacterium tuberculosis in Yunnan, China.

Journal of clinical laboratory analysis. 2019:e22884.

62. Jagielski T, Minias A, Van Ingen J, Rastogi N, Brzostek A, Żaczek A, et al. Methodological and clinical aspects of the molecular epidemiology of Mycobacterium tuberculosis and other mycobacteria. Clinical microbiology

reviews. 2016;29(2):239-90.

63. Ghavidel M, Mansury D, Nourian K, Ghazvini K. The most common spoligotype of Mycobacterium bovis isolated in the world and the recommended loci for VNTR typing; A systematic review. Microbial pathogenesis. 2018;118:310-5.