Quantitative Drug Susceptibility Testing of *Mycobacterium tuberculosis* by Use of MGIT 960 and EpiCenter Instrumentation[∇]

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Since numbers of drug-resistant *Mycobacterium tuberculosis* strains are on the rise, the simple classification into "susceptible" and "resistant" strains based on susceptibility testing at "critical concentrations" has to be reconsidered. While future studies have to address the correlation of phenotypic resistance levels and treatment outcomes, a prerequisite for corresponding investigations is the ability to exactly determine levels of quantitative drug resistance in clinical *M. tuberculosis* isolates. Here we have established the conditions for quantitative drug susceptibility testing for first- and second-line agents using MGIT 960 instrumentation and EpiCenter software equipped with the TB eXiST module. In-depth comparative analysis of a range of well-characterized susceptible and resistant clinical isolates has allowed us to propose conditions for testing and to develop criteria for interpretation.

Tuberculosis (TB) is a leading cause of morbidity and mortality worldwide. Developing countries are the most vulnerable, with more than 95% of the cases (32, 33). The present trend is characterized by an alarming emergence of drug resistance (7, 8, 30). Much attention has focused on the burden of multidrug-resistant (MDR) TB, i.e., resistance to the firstline drugs isoniazid and rifampin (rifampicin) (32, 33), and the emergence of extensively drug resistant TB (6, 26). The rise of drug-resistant TB and the increased susceptibility of the human population to TB due to coinfection with human immunodeficiency virus are driving the worldwide TB pandemic and will worsen the situation in the years ahead, with devastating effects in poor countries, whose economies suffer most from this development (19, 20).

In the diagnostic laboratory, testing of mycobacteria for drug susceptibility is substantially different from the general testing procedures used in bacteriology. Rather than determining MICs, a single drug concentration, termed the critical concentration, is usually used to categorize a clinical isolate as susceptible or resistant. This "critical concentration" is more an epidemiological parameter (to distinguish "wild-type" strains from "non-wild-type" strains that are able to grow in the presence of higher drug concentrations [5]) than a clinical cutoff value established to guide treatment decisions (14). With growing knowledge about the mechanisms that underlie drug resistance, it has become evident that drug resistance is multifaceted and that different mutations may lead to different levels of resistance. The acquisition of a resistance mutation leading to a decrease in drug susceptibility should not inevitably exclude

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an anti-TB drug from a treatment regimen, since low-level resistance does not necessarily imply clinical resistance (3). However, until now, different levels of phenotypic resistance have only rarely been taken into account in the procedures used for in vitro drug susceptibility testing (DST) of mycobacteria (4).

"Critical concentration"-based DST of primary and secondary drugs has been established for the radiometric Bactec 460 instrumentation (Becton Dickinson Microbiology Systems, Sparks, MD) and is considered the "gold standard" in the testing of second-line drugs (22, 25). However, the Bactec 460 system has several drawbacks: (i) it involves the use of sharps and radioisotopes with the need for disposal; (ii) it is only semiautomated; and (iii) it needs considerable hands-on time. The nonradiometric MGIT (mycobacterial growth indicator tube) 960 platform (Becton Dickinsion) has been evaluated extensively for DST of first-line drugs (1, 2, 15, 29) and has recently also been evaluated for second-line DST (16, 25). In contrast to the Bactec 460 instrumentation, the MGIT 960 platform is a fully automated system that uses a fluorescencequenching-based oxygen sensor for growth detection.

We have previously characterized quantitative drug resistance levels in clinical strains of drug-susceptible and drugresistant *Mycobacterium tuberculosis* using radiometric Bactec 460 measurements (28). However, for widespread implementation of quantitative DST in diagnostic mycobacteriology, the technique chosen should be fully automated, compatible with a computerized expert system for interpretation (so as to avoid individual errors and subjectivity), and safe and reliable (e.g., if possible, the system should not use radioactive material nor needles, such as syringes, for inoculation nor needles inside the instrument). To this end, we have subjected a carefully chosen subset of a previously described and well-characterized collection of clinical *M. tuberculosis* isolates (28) to quantitative measurements of drug susceptibility using the MGIT 960 plat-

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TABLE 1. Concentrations of drugs used for susceptibility testing

Drug	Concn(s) (mg/liter) followi	used for testing in the ng system:
U	Bactec 460	MGIT 960
Isoniazid Rifampin Ethambutol Streptomycin Amikacin Ethionamide Ofloxacin	0.1, 0.4, 1.0, 3.0, 10.0 1.0, 3.0, 10.0, 50.0 2.5, 5.0, 12.5, 50.0 1.0, 10.0, 50.0 ^b 1.0, 10.0, 50.0 2.5, 12.5 ^a 2.0, 10.0, 20.0	$\begin{array}{c} 0.1, 0.4, 1.0, 3.0, 10.0\\ 1.0, 3.0, 10.0, 50.0\\ 2.5, 5.0, 12.5, 25.0, 50.0\\ 1.0, 4.0, 20.0^{b}\\ 1.0, 4.0, 20.0\\ 2.5, 12.5^{a}\\ 2.0, 10.0, 50.0 \end{array}$
Linezolid Capreomycin	0.4, 4.0 5.0	$1.0, 4.0 \\ 1.25, 5.0, 25.0$

 a Additional testing for selected strains at concentrations of 1.25, 2.5, 5.0, 12.5, and 25.0 mg/liter.

^b Additional data points were determined for selected strains.

form in conjunction with EpiCenter software equipped with the TB eXiST module.

MATERIALS AND METHODS

Susceptibility testing using the Bactec 460 system. DST for first- and secondline anti-TB drugs was performed using the Bactec 460 system as recommended by the manufacturer (27). A 0.1-ml volume from a positive MGIT tube was inoculated into Bactec 460 12B vials; the final concentrations of drugs are given in Table 1. For the drug-free growth control, the bacterial suspension was diluted 1:100 before inoculation (proportion testing). The vials were incubated at 37° C and read daily until the growth of the control reached a growth index (GI) of >500. Test results were interpreted by comparing the changes in the GI of the growth control with the flasks containing the test drugs by using the standard interpretation procedure (if the GI of the drug is greater than or equal to the GI of the control, the strain is resistant [R]; if the GI of the drug is less than the GI of the control, the strain is susceptible [S]).

Susceptibility testing using the MGIT 960 system with EpiCenter TB eXiST software. The MGIT 960 system was used for primary isolation and standard susceptibility testing of first-line drugs (10) as recommended by the manufacturer. MGIT tubes supplemented with 0.8 ml of supplement (MGIT 960 SIRE supplement: Becton Dickinson) were inoculated with 0.1 ml of the drug solution and 0.5 ml of the test strain suspension. For preparation of the drug-free growth control tube, the organism suspension was diluted 1:100 with sterile saline, and then 0.5 ml was inoculated into the tube (proportion testing). For quantitative DST using the MGIT 960 instrumentation, we requested Becton Dickinson to develop custom-designed software with the following characteristics: automated recording of the readings, additional incubation time beyond the time to positivity of the drug-free control, minimization of the number of drug-free control tubes required, graphical representation of the growth unit (GU) value increase and storage of data, and easy handling and documentation. The software is EpiCenter, version 5.53, equipped with the TB eXiST module and available from Becton Dickinson. Table 1 lists the drug concentrations used for comparative analysis. The susceptibility testing sets were placed in the MGIT 960 instrument and continuously monitored using EpiCenter (version 5.53) TB eXiST software.

Results were interpreted as follows. At the time when the GU of the drug-free control tube was >400, if the GU of the drug-containing tube to be compared was \geq 100, the strain was R. If the GU of the drug-containing tube was <100, it was incubated for a further 7 days. If it was still <100, the strain was S. If the GU of the drug-containing tube was \geq 100 during this further 7 days of incubation after the GU of the drug-free control tube reached >400, the strain was intermediate (I).

Strains and molecular detection of resistance mutations. The isolates investigated in this analysis represent a subset of 29 clinical *M. tuberculosis* strains from a previously reported study (28).

For the identification of resistance mutations, the GenoType MTBDRplus assay (Hain Lifescience, Nehren, Germany) was used (11, 17). The GenoType MTBDRplus assay is a reverse hybridization line probe assay designed for rapid detection of *rpoB* and *katG* gene mutations. In addition, the strip harbors two wild-type probes covering the promoter region of the *inhA* gene, and four mutations in this region are targeted using mutated probes (-8T/C, -8T/A, -15C/T, -16A/G). The assay was performed as recommended by the manufac-

turer. Either the absence of a wild-type probe or a hybridization signal of a mutant probe is an indication of resistance.

For the detection of mutational alterations associated with resistance to ethambutol, amikacin, or streptomycin, i.e., *embB* position 306 (13), 16S rRNA position 1408 (23), and *rpsL* positions 42 and 87 (9), PCR-driven gene amplification and nucleic acid sequence determination were applied. Amplified gene fragments were sequenced using the BigDye Terminator cycle-sequencing ready reaction kit (Applied Biosystems Inc., Foster City, CA) and an ABI 3130 DNA genetic analyzer (Applied Biosystems).

RESULTS

A representative set of well-characterized clinical M. tuberculosis strains, with drug resistance defined at the genotypic level, was used to establish conditions for quantitative DST on the basis of the MGIT 960 instrumentation and EpiCenter, version 5.53, TB eXiST software. A total of 29 clinical isolates of M. tuberculosis, categorized as resistant to one or more first-line drugs on the basis of standard critical-concentration testing, were tested for susceptibility at higher drug concentrations using Bactec 460 and MGIT 960 instrumentation. Table 1 lists the drug concentrations used for comparative analysis. The EpiCenter software allows continuous monitoring of fluorescence, thus enabling a precise assessment of bacterial growth. Continuous growth monitoring, together with an extended incubation period of a further 7 days following the positivity of the drug-free control, facilitated the development of the term "intermediate growth" as an additional characteristic in data interpretation. The phenotypic heterogeneity in isoniazid resistance expression with the katG S315T mutation has been described previously (28) and may serve to illustrate the term "intermediate" (Fig. 1).

To determine the accuracy of quantitative DST, we compared the results of MGIT 960 testing with those of radiometric Bactec 460 testing and related them to the molecular resistance determinants identified. For molecular profiling, the GenoType MTBDRplus assay (Hain Lifescience, Nehren, Germany) was used (rifampin, isoniazid, and ethionamide); in part, mutations were assessed by nucleic acid sequencing (ethambutol, streptomycin, and amikacin).

For isoniazid, we included 11 strains with low-level drug resistance (MICs, ≥ 0.1 mg/liter and <1 mg/liter) (Table 2) and 18 strains with resistance levels of ≥ 1 mg/liter (Table 3). All low-level resistant isolates harbored an *inhA* promoter mutation (C15T). In 14/18 isolates with resistance levels of ≥ 1 mg/liter, a *katG* S315T mutation was present. The results obtained with the MGIT 960 platform correlated well with those generated by radiometric Bactec 460 testing over the whole range of drug concentrations tested (Table 3; Fig. 2). The only major discrepancy (S versus R) was observed at a test concentration of 3 mg/liter. Here the Bactec 460 system consistently reported susceptibility for a single isolate, while the MGIT 960 system gave a resistant test result; the two systems gave identical results for this isolate at test concentrations of 1 mg/liter (resistant) and 10 mg/liter (susceptible).

For rifampin, 11 isolates with resistance levels of ≥ 1 mg/liter were included in the analysis. Complete agreement between the two test systems was observed (Table 3; Fig. 2). Of the 11 isolates tested, 10 showed resistance at >50 mg/liter, and all 10 harbored well-described resistance mutations in the *rpoB* gene (Table 3). One isolate showed low-level resistance (MIC > 1



FIG. 1. Phenotypic heterogeneity in the isoniazid resistance of isolates with the *katG* S315 mutation and the term "intermediate" in MGIT 960 testing with EpiCenter (version 5.53) TB eXiST software. (A) Isolate 177836, resistant at 0.1, 1.0, 3.0, and 10.0 mg/liter; (B) isolate 186137, resistant at 0.1, 1.0, and 3.0 mg/liter and intermediate at 10.0 mg/liter; (C) isolate 186069, resistant at 0.1, 1.0, and 3.0 mg/liter and susceptible at 10.0 mg/liter; (D) isolate 176291, resistant at 0.1 and 1.0 mg/liter, intermediate at 3.0 mg/liter, and susceptible at 10.0 mg/liter. Drug concentrations are represented by colors as follows: blue, 0.1 mg/liter; green, 1.0 mg/liter, yellow, 3.0 mg/liter, red, 10.0 mg/liter; black, drug-free growth control.

mg/liter and < 3 mg/liter) with no detectable *rpoB* alteration; intermediate test results were not observed.

Ethambutol test results correlated well between the Bactec 460 and MGIT 960 (Table 3; Fig. 2), with three intermediate test results each obtained with the MGIT 960 at 5 mg/liter and

at 12.5 mg/liter. Comparison of the test concentration of 25 mg/liter (MGIT 960) with that of 50 mg/liter (Bactec 460) pointed to one discrepancy: the Bactec 460 system reported a sensitive result, while in MGIT 960 testing, the isolate showed resistance at a concentration of 25 mg/liter. Testing of the

			Sus	ceptibility	to the fo	ollowing di	rug at the	indicated	concn (mg/	liter):		
Strain	Method			Isoniazio	1				Ethionamic	le		Genetic resistance
		0.1	0.4	1.0	3.0	10.0	1.25	2.5	5.0	12.5	25.0	
TBC 2694	Bactec 460	R	S	S	S	S	R	R	ND^{a}	S	S	katG, wt ^b ; inhA, C15T
	MGIT 960	R	Ι	Ι	S	S	R	R	R	Ι	S	
TBC 4269	Bactec 460	R	S	S	S	S	R	R	R	R	R	katG, wt; inhA, C15T
	MGIT 960	R	Ι	Ι	S	S	R	R	R	R	R	
NZM 117	Bactec 460	R	S	S	S	S	R	R	ND	S	S	katG, wt; inhA, C15T
	MGIT 960	R	Ι	S	S	S	R	R	R	S	S	
NZM 150	Bactec 460	R	S	S	S	S	R	R	R	R	ND	katG, wt; inhA, C15T
	MGIT 960	R	Ι	S	S	S	R	R	R	R	Ι	
NZM 188	Bactec 460	R	S	S	S	S	R	R	ND	S	S	katG, wt; inhA, C15T
	MGIT 960	R	Ι	S	S	S	R	R	R	S	S	
TBC 2339	Bactec 460	R	S	S	S	S	R	R	ND	S	S	katG, wt; inhA, C15T
	MGIT 960	R	S	S	S	S	R	R	R	R	Ι	
TBC 179320	Bactec 460	R	S	S	S	S	R	R	R	R	ND	katG, wt; inhA, C15T
	MGIT 960	R	Ι	S	S	S	R	R	R	R	R	
TBC 179987	Bactec 460	R	S	S	S	S	R	R	ND	S	S	katG, wt; inhA, C15T
	MGIT 960	R	S	S	S	S	R	R	R	S	S	
NZM 186008	Bactec 460	R	S	S	S	S	R	R	ND	S	S	katG, wt; inhA, C15T
	MGIT 960	R	S	S	S	S	R	R	R	S	S	
NZM 186016	Bactec 460	R	S	S	S	S	R	R	ND	S	S	katG, wt; inhA, C15T
	MGIT 960	R	Ι	S	S	S	R	R	R	S	S	
TBC 176592	Bactec 460	R	S	S	S	S	R	S	ND	S	S	katG, wt; inhA, C15T
	MGIT 960	R	Ι	S	S	S	R	R	R	S	S	

TABLE 2. Clinical strains with low-level isoniazid resistance

^a ND, not done.

^b wt, wild type.

								j	5			UCICHINDIN SHIAIT	IS WIL	in m m	CTCOT 5	ומוועי	.10111 .	egu tu gilli-							
					Isol	niazic	q				Rifan	pin				Etha	mbuto						Strepto	omycin	
Strain	Method	Sus fc	cept ollow (mį	ibilit /ing (g/lite	y at t concr xr):	1 I	Genetic resistance	Su 1	Scepti ollow (mg	bility ing co //liter)	at the ncn :	Genetic resistance		Suscept follov (m	ibility /ing co g/liter)	at the mcn):		Genetic resistance		follo (r	tibilit wing ng/lite	y at th concn ar):	e	0 5	enetic iistance
		0.1	0.4	-	б	10		-	3	10	50		2.5	5 1	2.5 2	25	0			4	10	20	50		
NZM 130	Bactec 460 MGTT 960	22 22	2 2	2 2	2 2	2-	katG, S315T1 ^d ; inh 4 wt ^b	а К	2 2	2 2	2 2	S531L	R R	R R	D ^a		u en	tbB codon 306,	~~~	g,	s	s	sv	rpsL pos. ^c	42, AAG (wt);
NZM 179	Bactec 460	4 22	4 22	4 22	4 22	- R	katG, wt; inhA,	4 22	4 22	4 22	4 22	S531L plus	4 22	R S	nΖ	D	n eu	$bB \mod 306$,	4 22	, 2	, 2	2 M	2 M	pus. o., rpsL pos. 4	2, AGG (Arg)
	MGIT 960	2	2	Ч	Ч	2 2	C15T	2	К	К	ц	2nd mutation	ч К	RI	ŝ	Zc	D	CTG (valine)	К	ц	ц	К	Q a	pos. 87,	AAG (wt)
NZM 186006 (NZM 154/04)	Bactec 400 MGIT 960	* *	x x	* ~	^ –	n vo	<i>kat</i> G, wt; <i>mnA</i> , wf	х х	x x	トス	* *	TICCO	x x	x x v v	Zv	n∠ ⊇	D em	<i>tob</i> codon 300, ATA (isolencine)	トト	* *	* *	× ×	чg	rpsL pos. 4	2, AUU (Arg) AAG (wt)
NZM 186137	Bactec 460	Ч	Ч	Ч	К	s	katG, S315T1;	Ч	Ч	Ч	Ч	S531L	Ч	RS	Z	D	en	<i>bB</i> codon 306,	Ч	Ч	Ч	К	Ч	rpsL pos. 4	2, AGG (Arg)
(NZM 152/04)	MGIT 960	24	24	2	2	н с	inhA, wt	24	2	2	Ч		2	R I	ŝ	2 کے ب	D	wt (ATG)	2	2	2	2	g.	pos. 87,	AAG (wt)
1DC 0093	MGIT 960	2 2	2 2	2 2	0 F	- 0 U	(110100, 1111); inh 4 wt	2 2	00	00	00	۱M	00	0 U 0 U	ζυ	o ≥ ⊇	L L	at (ATG)	2 2		, E	00	00	rpsic pus. 4	2, AAG (wt); A AG (wt)
TBC 3106	Bactec 460	4 22	4 22	4 22	ч Ч	s s	katG, wt; inhA,	4 M	2 22	n M	n M	S531L	2 22	R	'nΖ	D S	en en	$bB \mod 306$,	3	ŝ	s	ŝ	ŝ	ND	(1)() (1)()
	MGIT 960	Ч	К	Ч	I	s	wt	Я	К	К	К		К	R S	S	Z	D	wt (ATG)	S	s	s	s	S		
NZM 186038	Bactec 460	Ч	Я	Ч	К	s	katG, S315T1;	Я	Ч	К	К	S531L	К	R S	Z	Ð	en	<i>bB</i> codon 306,	Ч	К	Ч	К	К	rpsL pos. 4	2, AGG (Arg)
(NZM 3/05)	MGIT 960	ц	Ч	ц	ц	- u	inhA, wt	ц	ц	ц	ц		ц	RS	ŝ	د ہے ہ	D	ATC (isoleucine)	ц	ч,	ч,	Ч,	g,	pos. 87,	AAG (wt)
IBC 176291	Bactec 460	× 1	× 1	× 1	¥,	s	katG, S31511;	× 1	× 1	× 1	× 1	H526D	X (s S	Ζœ	D S	uə -	<i>bB</i> codon 306,	ц	g,	s S	^	s s	rpsL pos. 4	2, AAG (wt);
TRC 177836	MGIT 960 Bacter 460	× ¤	× 2	2 2	_ ¤	s P	<i>inhA</i> , wt <i>batG</i> \$315T1.	х ¤	х 2	× ¤	× ¤	S5311	х ¤	2 C 2 C	nΖ	2 0 F	U "	wt (ATG) <i>bB</i> codon 306	× 2	s a	s c	su	n v	pos. 87,	AAG (wt) 2 AAG (wt):
	MGIT 960	4 22	4 22	4 22	4 22	4 22	inhA. wt	4 22	4 22	4 22	4 22		4 22	R N	200	2 2	D	wt (ATG)	4 22	4 22	4 22	00	000	nos. 87.	Z, 727 (mt), AGG (Are)
TBC 181783	Bactec 460	Ч	Ч	Ч	S	s	katG, wt; inhA,	Я	Ч	Ч	Ч	S531L	Ч	R	Z	D	en	<i>bB</i> codon 306,	Ч	Ð	S	s	S	rpsL pos. 4	2, AAG (wt);
	MGIT 960	ц	ц	ц	I d	s	C15T	ц	ц	ц	ц		ц	R R	22	s o		GTG (valine)	ц	ч,	ΩΩ,	I	g,	pos. 87,	AGG (Arg)
NZM 186069	Bactec 460 MGIT 960	× ¤	хc	× ¤	n n	n u	katG, S31511, inh 4 mt	× 2	× 2	× 2	× ¤	SS3IL	× ¤	X -	ζυ	n z D	u u	<i>bB</i> codon 306, ATA <i>(isolancina)</i>	× 2	n v	nu	n v	nu	rpsL pos. 4	2, AAG (wt); A AG (wt)
TBC 6890	Bactec 460	2	4 24	4 22	ŝ	s s	katG, S315T1;	4 22	4 22	4 22	4 22	S531L	30	s S S	'nΖ	D S	en en	$bB \mod 306$.	ź	Ę	² ND	, g	, g	ND	(111) 0171
	MGIT 960	К	К	К	I	s	<i>inhA</i> , wt	R	Z	ND	g		S	s S	S	Z	D	wt (ATG)	Q	g	ND	ŊŊ	Q		
TBC 7392	Bactec 460		2		s F	s s	katG, S315T1;	ŝ	s	s	s	wt	Ś	s c s c	Ζŭ	s z Q	u cu	bB codon 306,	g	22	g	g	g	QN	
TBC 295	Bactec 460	2 22	2 22	¥ 22	- 22	0 P	<i>mnA</i> , wt <i>katG</i> S315T1·	0 P2	z z	n N	d Z Z	S5311	0 CZ	n v n v	nΖ	s s E	n n	wt (AIG) bB codon 306	N N N		N N N		N N N	rnsI. nos 4	2 AGG (Aro)
	MGIT 960	2	4 24	4 22	2	2	inhA. wt	4 22	źŻ	: Z	: £		4 22	IS	ŝ	n Z 1	D	ATA (isoleucine)	: đ	; Ð	a	â	: B	pos. 87.	AAG (wt)
TBC 179882	Bactec 460	Ч	Я	Я	Я	s	katG, S315T1;	S	S	S	S	wt	S	s S	Z	DS	en	<i>bB</i> codon 306,	Q	Ð	ND	Ð	R	<u>A</u> D	
	MGIT 960	Ч	Ч	Ч	К	s	inhA, wt	S	Ħ	DN C	g		S	s S	S	2	́ Д	wt (ATG)	g	Ð	QN	QZ	g		
NZM 186073	Bactec 460	Ч	К	Ч	К	s	katG, S315T1;	S	S	S	S	DN	S	s S	Z	D	en	tbB codon 306,	Ð	Ð	Q	g	g	ND	
	MGIT 960	Ч	К	Ч	К	I	inhA, wt	S	Z	g	g		S	S	S	2	́ Д	wt (ATG)	Ð	Ð	QN	g	g		
NZM 186044	Bactec 460	2	2	2	2	с К	katG, S315T1;	29	Ë.	Z	g	ND	2	s c s	Ζc	S 2	un cu	$bB \mod 306$,	g	2	g	g	g	QN	
A DOOL FARM	MGI1 960	× c	¥ r	¥ 0	× c	х c	mhA, CISI	Z,	z.	z.	z.	1	¥ 0	<u>^</u>	nž	ہ ہے ج	n L	A I A (isoleucine)	2	2		2	2		
4000/1 INTN	MGIT 960	2 2	2 2	2 2	2 2	- 0 0	Katu, 5515111; inh 4 wt	2 E	٥Ż	۵Ż	۶ E	١M	nu	n v n v	ζυ	o ≥ ⊇	u U	wt (ATG)		E		E	2 E	UN	
		:	-	:	:		1W (17)10								,	1		(011)		2					

" ND, not done. bwt, wild type. c box, position. d Corresponds to an AGC to ACC substitution.

BACTEC460 BACTEC460 BACTEC460 BACTEC460 BACTEC460 0.1 mg/l 0.4 mg/l 1.0 mg/l 3.0 mg/l 10.0 mg/l R s R S R S R S R S R 18 R R 10 4 29 R 18 1 MGIT960 10.0 mg/l R _ **MGIT960 MGIT960 MGIT960** MGIT960 1.0 mg/l 0.4 mg/l 3.0 mg/l 0.1 mg/l _ T 8 T. 2 1 2 5 Т 1 3 s s 3 s 9 s S _ _ 11 21 Rifampicin ≥ 1.0 R (=11) BACTEC460 BACTEC460 BACTEC460 10.0 mg/l .0 mg/ 50.0 mg/l R S R S R S R 11 R 10 R 10 MGIT960 MGIT960 MGIT960 mg/l 50.0 mg/l 1.0 mg/l _ _ Т _ _ 10.01 s S 1 s 1 _ Ethambutol \geq 2.5 R (=12) BACTEC460 BACTEC460 BACTEC460 BACTEC460 2.5 mg/l 12.5 mg/ 50.0 mg/l 5.0 mg/l R s R s R s R s 7 R 12 _ R _ R 1 _ R 1 MGIT960 12.5 mg/l MGIT960 25.0 mg/l **MGIT960 MGIT960** 2.5 mg/l 5.0 mg/l 2 2 _ 1 I 1 1 _ _ s s 8 s S _ _ 2 _ 11 Streptomycin ≥ 1.0 R (=10) BACTEC460 BACTEC460 BACTEC460 BACTEC460 1.0 mg/l 10.0 mg/l 20.0 mg/l 50.0 mg/l R R R R S S S S 10 MGIT960 10.0 mg/l 5 MGIT960 20.0 mg/l 4 4 R R R **MGIT960** R NGIT960 _ 20.0 mg/l -1.0 mg/l _ -1 1 -3 S S 5 S 5 Ethionamide \geq 2.5 R (=19) BACTEC460 BACTEC460 2.5 mg/l 12.5 mg/l R R S S 2 6 R 17 MGIT960 R 4 **MGIT960** 12.5 mg/l 2.5 mg/l 2 _ L

Isoniazid ≥ 0.1 R (=29)

FIG. 2. Correlation of Bactec 460 and MGIT 960 susceptibility testing results.

isolate at a concentration of 50 mg/liter in MGIT 960 gave a susceptible test result, indicating a MIC for the isolate between 25 and 50 mg/liter. Seven of the 12 isolates with resistance levels of ≥ 2.5 mg/liter showed mutations in *embB* codon 306, while 5 isolates showed a wild-type (ATG) codon. No resistance of \geq 50 mg/liter was found.

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Ten isolates with streptomycin resistance levels of $\geq 1 \text{ mg/}$ liter were included in our study. Resistance at >1 mg/liter was consistently detected with both systems (Table 3; Fig. 2). All isolates (n = 4) with resistance levels of ≥ 50 mg/liter (Bactec 460) or \geq 20 mg/liter (MGIT 960) harbored an AAG \rightarrow AGG mutation at codon 42 of the rpsL gene. Of the strains with lower resistance levels (n = 6), two harbored AAG \rightarrow AGG mutations at codon 87 of the *rpsL* gene; in four strains with a low-level-resistant phenotype ($\leq 4 \text{ mg/liter}$), no *rpsL* mutation was found.

To further evaluate the capacity of the MGIT 960 system for quantitative DST of second-line agents, we determined the resistance profiles of *M. tuberculosis* isolates for ethionamide, amikacin, ofloxacin, capreomycin, and linezolid.

Nineteen isolates showed ethionamide resistance of ≥ 2.5 mg/liter in at least one of the two test systems. Of the 19 isolates tested, 2 strains showed deviations at a test concentration of 2.5 mg/liter (Tables 2 and 4): the Bactec 460 system gave a susceptible test result, while resistance was found using the MGIT 960 system. The two strains consistently showed a resistant phenotype at a test concentration of 1.25 mg/liter and susceptibility at a test concentration of 12.5 mg/liter for both test systems. At high drug concentrations, the MGIT 960 system tended to yield more resistant results than the Bactec 460 system, leading to four discrepancies at a test concentration of 12.5 mg/liter (Fig. 2).

TABLE 4.	Clinical M.	tuberculosis	strains	with	drug	resistance:	second line d	rugs
	Chine an in	1000100000	ou cuino			100100000000000000000000000000000000000	beechid mile d.	

				Et	hiona	mide					Amik	acin		Susc	eptib	ility t	the co	follo ncn (wing mg/lit	drug er):	at the	indic	ated
Strain	Method	Si	iscep follov (n	tibilit wing o ng/lite	y at th concn r):	he	Genetic resistance	S	uscep follo (n	tibilit wing ng/lite	y at t concr er):	he	Genetic resistance		Oflo	xacin		L	inezo	lid	Cap	reom	ycin
		1.25	2.5	5	12.5	25		1	4	10	20	50		2	10	20	50	0.4	1	4	1.25	5	25
NZM 130	Bactec 460	R	R	ND	S	S	inhA, wt ^b	S	ND	S	ND	S	ND	S	S	S	ND	S	ND	S	ND	ND	ND
	MGIT 960	R	R	R	R	S		S	S	ND	S	ND		S	S	ND	S	ND	S	S	I	S	S
NZM 179	Bactec 460	K D	R	R	R	ND	inhA, CIST	S	ND	S	ND	S	ND	S	S	S	ND	S	ND	S	ND	S	ND
NZM 186006	MGIT 900	K ND	K ND	K ND	K ND	I ND	inh 1 wit	5	S ND	ND S	S ND	ND S	ND	5	5	ND S	S ND	D	S ND	5	I ND	5	S ND
(NZM 154/04)	MGIT 960	ND	ND	ND	ND	ND	<i>uuu</i> /1, wt	S	S	ND	S	ND	ND	s	s	ND	S	ND	S	s	I	s	S
NZM 186137	Bactec 460	R	R	ND	S	ND	inhA wt	R	ND	R	ND	R	rm A1408G	S	S	S	ND	R	ND	S	ND	S	ND
(NZM 152/04)	MGIT 960	R	R	R	R	R	<i>uuu</i> 1, we	R	R	ND	R	ND	<i>m</i> , m , m	š	š	ND	S	ND	S	š	R	ND	S
TBC 6893	Bactec 460	S	S	S	S	S	inhA. wt	S	ND	S	ND	S	ND	Š	Š	S	ND	S	ND	Š	ND	ND	ND
	MGIT 960	ND	Š	ŝ	ŝ	Š		ŝ	S	ND	S	ND		ŝ	ŝ	ND	S	ND	S	Š	R	S	S
TBC 3106	Bactec 460	S	S	ND	S	S	inhA, wt	S	ND	S	ND	S	ND	S	S	S	ND	S	ND	S	ND	S	ND
	MGIT 960	ND	S	ND	S	S		S	S	ND	S	ND		S	S	ND	S	ND	S	S	S	S	S
NZM 186038	Bactec 460	R	R	ND	S	S	<i>inhA</i> , wt	S	ND	S	ND	S	ND	S	S	S	ND	R	ND	S	ND	S	ND
(NZM 3/05)	MGIT 960	R	R	R	Ι	S		S	S	ND	S	ND		S	S	ND	S	ND	S	S	S	S	S
TBC 176291	Bactec 460	ND	ND	ND	ND	ND	<i>inhA</i> , wt	S	ND	S	ND	S	ND	S	S	S	ND	ND	ND	ND	ND	ND	ND
	MGIT 960	ND	ND	ND	ND	ND		S	S	ND	S	ND		S	S	ND	S	ND	S	S	S	S	S
TBC 177836	Bactec 460	R	R	ND	S	S	<i>inhA</i> : wt	S	ND	S	ND	S	ND	S	S	S	ND	S	ND	S	ND	S	ND
TDC 101702	MGIT 960	R	R	R	R	R	: 1.4 C15T	S	S	ND	S	ND	ND	S	S	ND	S	ND	S	S	S	S	S
IBC 181/83	Bactec 460	K D	K D	K D	K D	ND D	inhA: C151	5	ND	S ND	ND	S ND	ND	K D	5	S ND	ND	K ND	ND	5	ND	5	ND
N7M 186060	MGIT 900	K ND	K ND	K ND	K ND	K ND	inh A. wt	5	S ND	ND S	S ND	ND S		к с	5	ND S	S ND	D	S ND	S C	S ND	5	S ND
INZINI 100009	MGIT 960	ND	ND	ND	ND	ND	<i>uuu2</i> 1. wt	s	S	ND	S	ND		s	s	ND	S	ND	S	s	P	s	S
TBC 6890	Bactec 460	S	S	S	S	S	$inhA \cdot wt$	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
100 0000	MGIT 960	ND	š	š	š	š	<i>uuu</i> 1. we	ND	ND	ND	ND	ND	T(D)	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
TBC 7392	Bactec 460	ND	ND	ND	ND	ND	inhA: wt	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
	MGIT 960	ND	ND	ND	ND	ND		ND	ND	ND	ND	ND		ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
TBC 295	Bactec 460	R	R	R	R	ND	inhA: wt	S	ND	S	ND	S	ND	S	S	S	S	S	ND	S	ND	ND	ND
	MGIT 960	R	R	R	R	R		S	S	ND	S	ND		S	S	S	S	ND	S	S	S	S	S
TBC 179882	Bactec 460	ND	ND	ND	ND	ND	inhA: wt	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
	MGIT 960	ND	ND	ND	ND	ND		ND	ND	ND	ND	ND		ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
NZM 186073	Bactec 460	S	S	S	S	ND	<i>inhA</i> : wt	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
N77 4 40 60 4 4	MGIT 960	S	S	S	S	ND		ND	ND	ND	ND	ND	NID	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
NZM 186044	Bactec 460	K D	5	5	S	ND	inhA: CIST	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
NZM 178264	Rector 460	K ND	K ND	K ND	3 ND	э ND	inh A. wt	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
INZIVI 1/0304	MGIT 960	ND	ND	ND	ND	ND	uuu/1. wi	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
	WIGH 900	ц	ц	ц	T D	TAD.		TAD.	ц	цр	ц	ц		чD	чD	ц	TAD.	ц	ц	цр	ц	чD	

^a ND, not done.

^b wt, wild type.

Of 12 strains tested, one isolate showed high-level resistance to amikacin (\geq 50 mg/liter). This isolate harbored the typical 16S rRNA mutation A1408G (23). Resistance at all test concentrations for this isolate, as well as susceptibility for the other strains investigated, was consistently determined with both test systems (Table 4).

Twelve strains were available for the comparison of testing for susceptibility to ofloxacin. Both systems consistently gave susceptible test results at a concentration of 2 mg/liter for 11 isolates. One isolate showed resistance at 2 mg/liter and sensitivity at 10 mg/liter (Table 4).

For linezolid and capreomycin, no resistant isolates with MICs above 4 mg/liter and 5 mg/liter, respectively, were present in the collection of strains under investigation. For the isolates studied, both systems consistently revealed sensitive test results at these drug concentrations (Table 4).

DISCUSSION

With the global rise of MDR strains, there is an increasing need to determine susceptibility to first and second-line anti-TB agents exactly. Treatment of patients with drug-resistant TB should be based on reliable and quantitative measures of susceptibility testing, a cornerstone for preventing further amplification of resistance (18) and for optimally exploiting available compounds. Detailed knowledge on quantitative drug resistance may guide empirical treatment of drug-resistant TB, e.g., addressing whether and when to add second-line drugs. A fundamental drawback to this strategy is that even in industrialized countries, only a limited panel of anti-TB drug concentrations is tested, leaving the exact resistance level of clinical *M. tuberculosis* isolates vestigial. In principle, automatic systems have the potential to meet the challenge of precise determinations of drug resistance levels with reasonable labor input. The MGIT 960 platform has been extensively validated for testing susceptibility to first-line anti-TB agents at critical drug concentrations (2, 24).

Here we systematically evaluated the performance of the MGIT 960 instrumentation for quantitative susceptibility testing of drug-resistant clinical isolates of *M. tuberculosis*. To investigate the underlying molecular resistance mechanisms, we used direct PCR-mediated sequence analysis of nucleic acids or the GenoType MTBDRplus assay. Resistance to isoniazid or ethionamide is due mainly to alterations in proteins that metabolize the prodrug into its active component, such as KatG or EthA. In contrast, resistance to rifampin, ethambutol,

Drug	Recommendation	Comment
Isoniazid	0.1 mg/liter is used for screening Low-level-resistant isolates are R at 0.1 mg/liter and S at 1.0 mg/liter R at 10.0 mg/liter indicates that isoniazid is of no clinical use Test results of R at 1.0 mg/liter and S at 10.0 mg/liter need further clinical data for interpretation 1.0 mg/liter is presumably the most important concn to test	Mutations in <i>inhA</i> typically lead to R at 0.1 mg/ liter and S at 1.0 mg/liter; mutations in $katG$ result in R at 1.0 mg/liter and show varying test results at 10.0 mg/liter
Rifampin	1.0 mg/liter is used for screening 10.0 mg/liter is used to recognize high-level drug resistance	Most, if not all, mutations in <i>rpoB</i> are associated with R at >50.0 mg/liter; good correlation between genotype and phenotype
Ethambutol	5.0 mg/liter is used for screening 25.0 mg/liter is used to recognize high-level drug resistance	Isolates with mutations in <i>embB</i> are mostly, if not always, susceptible or intermediate at 25.0 mg/ liter
Streptomycin	 1.0 mg/liter is used for screening 20.0 mg/liter is used to recognize high-level drug resistance 4.0 mg/liter is used to separate low-level (R at 1.0 mg/liter; S at 4.0 mg/liter) from intermediate (R at 1.0 mg/liter; I or R at 4.0 mg/liter; S at 20.0 mg/liter) resistance 	Good correlation between genotype and phenotype
Amikacin	 1.0 mg/liter is used for screening 20.0 mg/liter is used to recognize high-level drug resistance 4.0 mg/liter is used to separate low-level (R at 1.0 mg/liter; S at 4.0 mg/liter) from intermediate (R at 1.0 mg/liter; I or R at 4.0 mg/liter; S at 20.0 mg/liter) resistance 	
Ethionamide	2.5 mg/liter is used for screening Additional testing at 12.5 mg/liter	Mutations in <i>inhA</i> (C15T) may show phenotypic variability: unanimously R at 2.5 and 5.0 mg/ liter; I or R at 12.5 mg/liter; S, I, or R at 25.0 mg/liter
Ofloxacin	2.0 mg/liter is used for screening Additional testing at 10.0 and 50.0 mg/liter	
Capreomycin	5.0 mg/liter is used for screening Additional testing at 25.0 mg/liter	
Linezolid	1.0 mg/liter is used for screening Additional testing at 4.0 mg/liter	

TABLE 5. Recommendations for quantitative DST on the basis of the EpiCenter-equipped MGIT 960 system

streptomycin, or quinolones is due to mutational alterations in genes encoding the drug target itself, such as *rpoB*, *rpsL*, *rrs*, or *gyrA* (21, 31).

For the first-line drugs, the quantitative resistance levels determined by the Bactec 460 and MGIT 960 systems were in good agreement. Introduction of the parameter "intermediate," however, allowed a more-precise determination of resistance in the MGIT 960 system. It turned out that susceptibility testing of isolates with MICs near the test concentration may result in intermediate test results in the MGIT 960 system. The most prominent isoniazid resistance-conferring *katG* mutation (S315T) is associated with a heterogeneous resistance phenotype, with MICs ranging between 1 and 10 mg/liter.

The need for international standardization of testing of susceptibility to second-line anti-TB agents still remains. We have evaluated the performance of the MGIT 960 system for quantitative testing of susceptibility to the second-line drugs ethionamide, amikacin, ofloxacin, linezolid, and capreomycin. The results with the two test systems show good concordance. However, the conclusions are limited due to the small number of resistant isolates available to us. The drug most frequently affected by Bactec 460–MGIT 960 discrepancies was ethionamide. This is in accordance with previous studies (12, 25), which revealed more resistant results with the MGIT 960 instrumentation than with comparative test systems. Promoter alterations of *inhA* resulting in ethionamide resistance were observed in 14 of our ethionamide-resistant isolates (Tables 2 and 4). However, ethionamide resistance may also be due to *ethA* mutations not included in the GenoType MTBDRplus assay.

In general, the MGIT 960 system tended to give resistant test results more frequently than the Bactec 460 system. Medium differences and technical differences in the procedure used for growth measurement may account for this finding. Both instruments determine growth as a measure of metabolic activity. The Bactec 460 instrument measures volatile radioactive CO_2 as metabolized from the radioactive precursor palmitic acid; for this purpose, most of the gaseous phase in the vial is removed and replaced. In contrast, the MGIT 960 instrument uses a fluorescence-quenching-based oxygen sensor for growth detection without manipulation of the closed vial. Under conditions of exceedingly slow growth, e.g., near the MIC, the Bactec 460 system would be expected to result in "false" susceptible test results, since it constantly depletes the vial of the little CO_2 produced.

Existing procedures for DST of mycobacteria are adequate for screening but require complementation with quantitative DST measures, in particular for those drugs where heterogeneity in phenotypic resistance is present. We have established the conditions for quantitative DST using the MGIT 960 system in combination with EpiCenter software equipped with the TBeXiST module, thus providing a fully automated walk-away system for quantitative DST of M. tuberculosis. This platform allows electronic data management and is compatible with expert systems for interpretation. The MGIT 960 platform in conjunction with the EpiCenter software shows high consistency with Bactec 460 test results over a wide range of concentrations tested for first- and second-line anti-TB drugs. While we note that further studies are needed to address the correlation of phenotypic resistance levels and treatment outcome, we have summarized our recommendations for quantitative DST of M. tuberculosis in Table 5. Widespread implementation of MGIT 960 protocols for quantitative DST should provide standardized data to enable the correlation of results from quantitative DST with clinical outcomes by high-throughput statistical analysis in order to address the issue of phenotypic drug resistance levels and treatment failure. In addition, data sets obtained by an automated standardized procedure based on agreed guidelines provide optimal input for monitoring the epidemiology of resistance at a supranational level.

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