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Unexpectedly High Proportion of Ancestral Manu Genotype *Mycobacterium tuberculosis* Strains Cultured from Tuberculosis Patients in Egypt[∇]

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Tuberculosis is one of the important public health problems in Egypt. However, limited information on the *Mycobacterium tuberculosis* genotypes circulating in Egypt is available. A total of 151 *M. tuberculosis* strains were characterized by spoligotyping. The results revealed that 74.8% of *M. tuberculosis* isolates grouped into 13 different clusters, while 25.2% had unique spoligotype patterns. Comparison with an international spoligotyping database (the SITVIT2 database) showed that types SIT53 (T1 variant) and SIT54 (Manu2 variant) were the most common types between cluster groups. In addition, new shared types SIT2977, SIT2978, and SIT2979 were observed. The results identified for the first time an unusually high proportion of ancestral Manu strains of *M. tuberculosis* from patients in Egypt. The percentage of the Manu clade in this study (27.15%) was significantly higher than its overall representation of 0.4% in the SITVIT2 database. We show that in Egypt tuberculosis is caused by a predominant *M. tuberculosis* genotype belonging to the ancestral Manu lineage which could be a missing link in the split between ancestral and modern tubercle bacilli during the evolution of *M. tuberculosis*.

Despite the availability of antituberculosis (anti-TB) drugs, the disease burden of human TB remains a very serious and widespread public health problem. Many of the 22 countries most affected by human TB identified by the World Health Organization (WHO) are developing countries (11). Although Egypt is not on the WHO list of 22 high-TB-burden countries, it is considered one of the high-burden countries in WHO's Eastern Mediterranean region, where TB constitutes the second most important public health problem after schistosomiasis (21). Every year, the National Tuberculosis Control Program of the Ministry of Health and Population (MOHP) registers over 12,000 new TB patients; more than 50% of the cases are sputum smear-positive pulmonary TB. On the basis of an annual risk of infection of 0.32%, it is estimated that about 8,000 people receive a diagnosis of TB at facilities other than those of MOHP (25).

In the context described above, it is important to have a precise picture of the predominant and emerging *Mycobacterium tuberculosis* clones in Egypt, in order to have a first snapshot of the epidemiology of TB. The advent of molecular typing methods has given a new boost to epidemiological studies of TB in recent years (3); the most commonly used methods

are IS6110-based restriction fragment length polymorphism (RFLP) (IS6110-RFLP) and spoligotyping (12, 22). As opposed to IS6110-RFLP, which was considered the “gold standard” for the genotyping of *M. tuberculosis* in the early 1990s (22) and which is time-consuming and labor-intensive and requires large amounts of highly purified DNA, PCR-based spoligotyping avoids the problems associated with the slow growth of these bacteria, thereby offering the possibility of the rapid recognition of outbreaks (2, 4, 12). The clinical usefulness of spoligotyping is determined by its rapidity both in detecting the causative bacteria and in providing epidemiological information on strain identities (12). Spoligotyping, which allows determination of the phylogeographical specificity of circulating clades of tubercle bacilli, is also useful for population-based studies (4, 10). Recent data from international spoligotyping studies have identified a growing number of important clades or genogroups, further facilitating the study of the spread of the disease due to human migratory movements (4, 8, 9, 17). To date, there is very limited genotyping information on the *M. tuberculosis* strains circulating in Egypt. The SITVIT2 database, which is an updated version of the previously released SpolDB4 database (4; available online at <http://www.pasteur-guadeloupe.fr:8081/SITVITDemo>), showed very little information on the isolates circulating in Egypt, with only 79 of 69,000 clinical isolates, i.e., 0.1% of the total recruitment. The aim of this study was to identify the predominant international spoligotypes responsible for the transmission and prevalence of TB in Egypt. The spoligotypes obtained were further compared with those in an updated version of the international database named SITVIT2.

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MATERIALS AND METHODS

Patients and specimens. A total of 151 *Mycobacterium tuberculosis* strains from as many patients collected over a period of 1 year (2005) at three different chest hospitals in Egypt were included in this study. The isolates were recovered from sputum specimens from patients diagnosed with pulmonary TB on the basis of clinical symptoms, chest X ray, and bacteriological examination. Despite an attempt to ensure that the sample is an accurate representation of the pulmonary TB patients attending the three chest hospitals over a 1-year period, it instead represents a coincidental group (convenience sample) of patients originating from 14 different governorates across Egypt. Demographic and epidemiological data were obtained from the medical records of each patient. Three sputum specimens were collected in the early morning on consecutive days and transported to the Tuberculosis Center Laboratory, Faculty of Medicine, Cairo University, Cairo, Egypt. Samples were decontaminated with *N*-acetyl-L-cysteine and sodium hydroxide, as described previously (13). The decontaminated specimens were concentrated by centrifugation at $3,000 \times g$ for 20 min. All handling and manipulations of the clinical specimens were done in a certified biosafety level II cabinet. The processed specimens were subjected to Ziehl-Neelsen staining, followed by culture onto Lowenstein-Jensen medium. The cultures were identified as *M. tuberculosis* complex on the basis of the growth rate, colony morphology, and cultural and biochemical properties by standard microbiological procedures. The phenotypic identification of the *M. tuberculosis* complex was further confirmed by the amplification of a 540-bp DNA fragment specific for the IS6110 gene (14). For this purpose, bacterial DNA was extracted by using a Qiagen DNA isolation kit, according to the manufacturer's protocol.

Spoligotyping. Spoligotyping was carried out as described previously (12). Briefly, the direct repeat (DR) region of the TB genome was amplified with primers DRa and DRb, and the amplified biotinylated products were hybridized to a set of 43 oligonucleotides covalently bound to a membrane (Isogen Life Science B.V., Utrecht, The Netherlands). The hybridized PCR products were then incubated with streptavidin-peroxidase conjugate, and the membrane was exposed to chemoluminescence (Amersham, Little Chalfont, England) and X-ray film (Hyperfilm ECL; Amersham), according to the manufacturer's instructions. The X-ray film was developed by standard photochemical procedures after overnight exposure. To avoid any possibility that nonspecific hybridization spots would appear on the commercial membranes, appropriate controls that included DNA from the known strain *M. tuberculosis* H37Rv and deionized autoclaved water were used in each experiment. The reproducibility of spoligotyping was confirmed by repeating the test. Spoligotyping in a binary format was converted to an octal code (7; the conversion tool is available at <http://www.pasteur-guadeloupe.fr:8081/SITVIT2Demo/outilsConversion.jsp>) for comparison with the codes in the SITVIT2 proprietary database of the Pasteur Institute of Guadeloupe.

Comparison of spoligotyping results with database types. Spoligotypes in binary format were entered into the SITVIT2 database (Pasteur Institute of Guadeloupe), which is an updated version of the previously released SpolDB4 database (4; available at <http://www.pasteur-guadeloupe.fr:8081/SITVIT2Demo>). At the time of the present study, SITVIT2 contained genotyping information on 69,000 *M. tuberculosis* clinical isolates from 160 countries of origin. In this database, the Spoligotype International Type (SIT) designates spoligotypes shared by two or more patient isolates, whereas "orphan" designates the patterns reported for a single isolate. Major phylogenetic clades were assigned according to the signatures provided in the SpolDB4 database, which defined 62 genetic lineages and sublineages (4). These include specific signatures for various *M. tuberculosis* complex members, as well as rules defining the major lineages and sublineages for *M. tuberculosis* sensu stricto. These include specific signatures for various *M. tuberculosis* complex species, such as *M. bovis*, *M. microti*, *M. caprae*, *M. pinipedi*, and *M. africanum*, as well as rules defining the major lineages and sublineages of *M. tuberculosis* sensu stricto. These include the Beijing clade, the Central Asian (CAS) clade and 2 sublineages, the East African-Indian (EAI) clade and 9 sublineages, the Haarlem clade and 3 sublineages, the Latin American-Mediterranean (LAM) clade and 12 sublineages, the Manu family and 3 sublineages, the S clade, the IS6110-low-banding X clade and 4 sublineages, and an ill-defined T clade with 5 sublineages.

Phylogenetic analysis. The evolutionary relationships among all the spoligotype patterns in our sample were studied by drawing a minimum-spanning tree (MST) with Bionumerics software (version 3.5, Applied Maths, Sint-Marteen-Latem, Belgium). MST is an undirected network in which all the samples are linked together with the smallest possible linkages between nearest neighbors. By this approach, one considers that all intermediate stages are present within the sample analyzed, by including first the individual that shows the most possible linkages to other individuals in the population studied.

RESULTS AND DISCUSSION

TB remains a great public health concern in Egypt; however, little information on the prevailing genotypes of the tubercle bacilli and their spread is available (1, 6). In a country where TB is endemic, it is critical to identify the genotypes of the predominant strains in order to study the transmission patterns and the epidemiological features of the disease. The present study is a first attempt to provide insight into the population structure of *M. tuberculosis* in Egypt. We selected spoligotyping to create this snapshot, as it has been shown to be a valuable method for determination of the clonal and phylogenetic relationships of *M. tuberculosis* strains, providing in parallel useful information about TB in a given population, in a given country, among countries, and throughout the world (4, 5, 9, 10, 12, 16, 17, 19).

This study was performed over a period of 1 year (2005) with a total of 151 *M. tuberculosis* complex strains from as many patients from Egypt. The demographic information for the patients (Table 1) underlined that within our sample the disease predominated in male patients (male-to-female sex ratios, 3.52 for the cluster group and 3.75 for the unclustered group) and individuals aged 15 to 45 years (70.7% of the cases grouped in clusters and 68.4% of the unclustered group). Of all cases, 118/151 (78.1%) patients had newly diagnosed TB, whereas 33/151 (21.9%) patients were previously treated. Lastly, 135/151 (90%) patients were native and 15/151 (10%) were immigrants (the origin was unreported for 1 patient). However, none of the parameters shown in Table 1 varied significantly among any of the groups (by Fisher's exact test with an online statistical calculator [<http://www.graphpad.com/quickcalcs/contingency1.cfm>]).

Detailed spoligotyping results are summarized in Tables 2 and 3, while the evolutionary relationships of the strains are illustrated in an MST shown in Fig. 1. Starting with our sample of 151 clinical isolates, spoligotyping produced a total of 51 different patterns. A total of 12 patterns corresponding to 13 strains were not present in the database and could be termed orphans (Table 2); nonetheless, 2 strains (1 from a patient born in Egypt and another from a patient born in Sudan) matched each other, thereby creating a new shared type, SIT2977. Two more orphans matched one strain each from Austria and Taiwan, thereby creating SIT2978 and SIT2979, respectively. As illustrated in Table 3, 138/151 strains from this study belonged to a total of 39 SITs (not including the 3 newly created SITs shown in Table 2). The binary and octal descriptions, the total number of strains, and the percentage of strains in the present study compared to the same data in the SITVIT2 database at the time of comparison and the corresponding genetic lineages and sublineages are summarized for each of the shared types shown.

Irrespective of the database comparison, a total of 113/151 (74.8%) clinical isolates were clustered in the present study (13 clusters containing 2 to 51 isolates per cluster). Twelve clusters are shown in Table 3, and 1 additional cluster of two strains is represented by the newly created shared type (SIT2977; Table 2). Thus, 38/151 (25.2%) of the strains in this study were unclustered. SIT53, which belongs to the ill-defined T clade (the T family was defined by default [4]), predominated in our setting and contained 51/151 (33.8%) of the isolates. That was

TABLE 1. Clinical and epidemiological characteristics of patients harboring clustered versus nonclustered strains

Parameter (<i>n</i> = 151)	Cluster groups (<i>n</i> = 113)				Noncluster group (<i>n</i> = 38)
	All clusters (<i>n</i> = 113)	SIT53 (<i>n</i> = 51)	SIT54 (<i>n</i> = 33)	Other cluster groups (<i>n</i> = 29)	
No. (%) of patients ages (yr):					
15–45	80 (70.7)	39 (76.6)	22 (66.6)	19 (65.5)	26 (68.4)
≥46	33 (29.3)	12 (23.3)	11 (33.4)	10 (34.5)	12 (31.6)
No. (%) of patients of the following sex:					
Male	88 (77.8)	39 (76.6)	28 (84.8)	21 (72.4)	30 (78.9)
Female	25 (22.2)	12 (23.3)	5 (15.2)	8 (27.6)	8 (21.1)
Sex ratio ^a	3.52	3.25	5.6	2.6	3.75
No. (%) of patients with history of TB					
No previous therapy	86 (76)	42 (82.3)	21 (63.6)	23 (79.3)	32 (84.2)
Previously treated	27 (24)	9 (17.7)	12 (36.4)	6 (20.7)	6 (15.8)
No. (%) of patients with the following country of origin:					
Egyptian	101 (89.3)	46 (90.1)	27 (81.8)	28 (96.5)	34 (89.4)
Foreign born	12 (10.7)	5 (9.9)	6 (18.2)	1 (3.5)	3 (10.6)

^a The sex ratio is the number of males/number of females.

followed by SIT54, which belongs to the Manu2 clade and which contained 33/151 (21.8%) of strains, followed by SIT25, a CAS1-Delhi variant (4/151 [2.7%]); SIT41 of the LAM7-TUR lineage (4/151 [2.7%]); and SIT291 of the ill-defined T1 variant (4/151 [2.7%]). Overall, 51 strains were identified in SIT53, making up 33.8% of the strains studied. These were from different location in Egypt: 36 from Cairo; 7 from Qalyubia; 4 from Giza; and 1 each from Monufia, Sohag, BeniSuef, and Asyut. The second largest cluster, SIT54 (21.8%), contained 21 strains from Cairo; 4 from Giza; 3 from Qalyubia; and 1 from each Sohag, Helwan, Monufia, Sinai, and Kafr-elSheikh.

The overall classification of the strains according to the major *M. tuberculosis* genotypic lineages (and not the shared types *stricto sensu*) showed that the most predominant group was the ill-defined T superfamily (81/151 [53.6%]). However, the T genotype does not represent a clade in a strict evolutionary sense, since it was defined by default to include strains that cannot be classified in one of the established genotypic lineages, such as the Haarlem, LAM, CAS, and EAI lineages, with well-established phylogeographical specificity (4). In this sense, the most predominant clade in our setting was the ancestral Manu lineage, which comprised not only the 33 isolates of SIT54 (Manu2_var) but also a group of other Manu strains (Tables 2 and 3): 2 Manu2_var orphans; 1 SIT523 strain classified as a Manu ancestor; 3 Manu2_var strains belonging to SIT1088, SIT1196, and SIT1690, respectively; a single SIT1378 (Manu3) strain; and 1 SIT2781 strain classified as Manu ancestor var_Δ33 (lacking a single spacer at position 33). Thus, by taking all the Manu isolates together (shown in boldface in Tables 2 and 3), they represented 41 of 151 (27.15%) of the strains in the present study. This is one of the highest distribution patterns ever observed for the ancestral Manu lineage in any single setting. The predominance of the Manu lineage was followed by the LAM (7/151 [4.6%]), CAS1-Delhi (6/151 [4%]), Haarlem (3/151 [2%]), and S (2/151 [1.3%]) lineages.

Before the entry of our data, spoligotypes from 79 strains were recorded in the SITVIT2 database. These strains were isolated from patients either in Egypt or from elsewhere in the world from a patient of Egyptian origin (56 strains classified

within 30 different SITs and 23 orphans). Some previously found SITs belong to predominant lineages in Egypt, particularly SIT53 (T1 prototype), SIT54 (Manu2 prototype), SIT25 (CAS1-Delhi_var), and SIT41 (LAM7-TUR), while others (SIT34, SIT37, SIT42) are found in smaller numbers (Table 2). On the other hand, many of the SITs found in patients of Egyptian origin but isolated in distant countries were not found in the present study performed in Egypt. Such examples include SIT35 (Haarlem3), SIT47 (Haarlem1), SIT52 (T2), SIT390 (Haarlem3), SIT460 (EAI5), SIT717 (T1), SIT732 (T1), SIT798 (T), and SIT878 (X1).

The unusually high proportion of strains belonging to the Manu clade (27.15%) in the present study is really exceptional. This percentage is significantly higher than its overall representation (0.4%) in the SITVIT2 database (*n* = 261). The Manu lineage was initially described as a new family from India in 2004 (16), and later, similar strains in small numbers were reported in a study from Madagascar (8). Soon afterwards, it was tentatively subdivided into the Manu1 (deletion of spacer 34), Manu2 (deletion of spacers 33 and 34), and Manu3 (deletion of spacers 34 to 36) sublineages; and it was suggested that this lineage could represent an ancestral clone of principal genetic group 1 strains (4). More recently, Manu lineage strains were reported from Saudi Arabia (2) and Tunisia (15).

Interestingly, the Manu2 strains (prototype SIT54) have been reported only scarcely in the SITVIT2 database (*n* = 130, not counting this study). In the updated database, SIT54 strains are present in the highest numbers in India (*n* = 21), followed by Saudi Arabia (*n* = 20), the United States (*n* = 20), South Africa (*n* = 17), Russia (*n* = 9), Brazil (*n* = 6), Great Britain (*n* = 3), Madagascar (*n* = 3), and Poland (*n* = 3). Smaller numbers of SIT54 strains (two or less) have been reported from Armenia, Austria, Bangladesh, China, Cuba, Italy, France, Georgia, Greece, Guinea-Bissau, Iran, Latvia, Panama, Turkey, Senegal, Sweden, and Sudan. Although the origin of the patient was not always indicated in the SITVIT2 database, many of these patients were foreign born; e.g., the two patients from Saudi Arabia were born in Chad, while one patient was from Indonesia. The single case from Australia was from Eritrea, and the two cases from South Africa were from

ISO no. ^d	Spoilgotype description	Ociel no.	Country of origin ^e	City of isolation	MATCH with another orphan (new SIT) ^f	Claude ^c
EGGY0120053	███████	70337777763771	EGY	Cairo		Manu2_yar
EGGY0120056	███████	7573777377763771	EGY	Aswan		Manu2_yar
EGGY01200512	███████	7761775777760771	EGY	Cairo		Unk
EGGY01200516	███████	7777374677760771	EGY	Giza		Unk
EGGY01200526	███████	5777774377760771	EGY	Cairo		Unk
EGGY01200590	███████	4377770377760771	EGY	Qalyubia		Unk
EGGY010000174	███████	203067777670671	EGY	Monufia		Unk
EGGY010000179	███████	7533477777760771	EGY	Cairo		Unk
EGGY01200576	███████	7023777777760771	SDN	Cairo		Unk
EGGY0120054	███████	777777377360771	EGY	Qalyubia	This study (SIT2977)	T4_var
EGGY01200552	███████	777777377360771	SDN	Cairo	This study (SIT2977)	Unk
EGGY01200549	███████	777777777060771	EGY	Giza	AUT (SIT2978)	Unk
EGGY012005105	███████	0000000000000171	EGY	Cairo	TWN (SIT2979)	Beijing_var

^a EGY, Egypt; SDN, Sudan.
^b Concerns only new SITs created after a match with another orphan strain in the SITVIT2 database. AUT, Austria; TWN, Taiwan.
^c Clade designations according to SITVIT2 by use of revised SpoIDB4 rules. Unk, unknown patterns within any of the major clades described in SITVIT2. The Manu lineage strains are indicated in boldface.
^d ISO, International Organization for Standardization.

^b Concerns only new SITs created after a match with another orphan strain in the SITVIT2 database. AUT, Austria; TWN, Taiwan

Clade designations according to SIVIT2 by use of revised SpolDB4 rules. Unk, unknown patterns within any of the major clades described in SIVIT2. The Manu lineage strains are indicated in boldface.

^d ISO, International Organization for Standardization

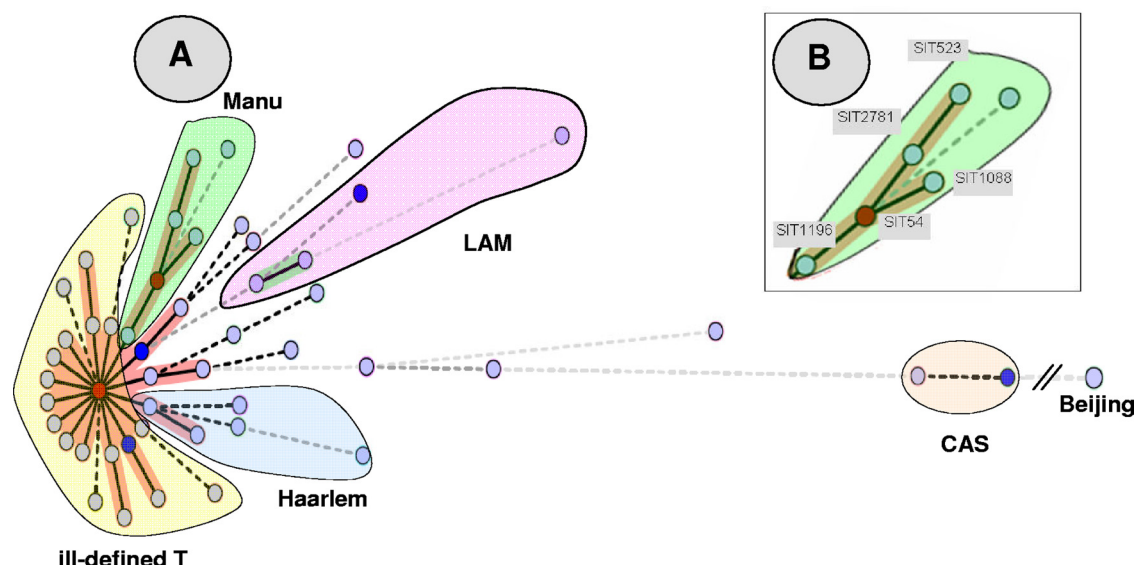


FIG. 1. MST of potential evolutionary relationships of *M. tuberculosis* spoligotypes from Egypt built with Bionumerics software. Solid lines, single spacer change; dotted lines, two spacer changes (black) or more spacer changes (gray). The color of the circles is proportional to the number of clinical isolates in our study, illustrating unique isolates (sky blue) versus clustered isolates (deep blue, two to five strains; red, six strains and more). (A). As opposed to the clades belonging to the principal genetic group 1 (such as Beijing and CAS) in the MST, the Manu strains are the ones situated closest to the principal genetic group 2 or 3 lineages, such as Haarlem, LAM, and T. The central node of this unrooted tree is represented by SIT53, which is the prototype of the T1 lineage. (B). A zoom of the Manu lineage representing SIT523 (Manu ancestor), SIT2781 (Manu ancestor var_Δ33), SIT1088 (Manu2_var), SIT54 (Manu2), and SIT1196 (Manu2_var). In this scenario, the evolution is supposed to happen starting from SIT523 (Manu ancestor), which is the most conserved spoligotype in the SITVIT2 database (with positive hybridization for all 43 spacers of the DR locus) to SIT1196 (Manu2_var, with absence of consecutive spacers 33 to 35).

that these ancient strains could represent an intermediate clone between an *M. africanum* lineage and the Manu lineage. In conclusion, our study shows that in Egypt TB is caused by a predominant genotype that belongs to the ancestral Manu lineage and that could be a missing link of the split between ancestral and modern tubercle bacilli during the evolution of *M. tuberculosis*. We believe that detailed investigations of the Manu lineage strains by 24-locus mycobacterial interspersed repetitive unit analysis, extended spoligotyping, analysis of regions of deletions, and single-nucleotide polymorphism analysis would shed light on the central role of isolates from Egypt and Mesopotamia in the origin, evolution, and global spread of TB.

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