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Classified under the phylum Actinobacteria, the mycobacteria consist of over 150 different species that include certain pathogens known to cause severe diseases in mammals. The *Mycobacterium tuberculosis* complex (MTBC) involving *M. tuberculosis*, *M. bovis*, *M. bovis BCG*, *M. africanum*, *M. microti*, *M. cannetii*, *M. caprae* and *M. pinnipedii* are the cause for human and animal tuberculosis. The causative organism for tuberculosis (TB) is the *M. tuberculosis* (*Mtb*) which is one of the most widely spread disease causing mycobacteria in humans. The MTBC species are closely related taxonomic group of bacteria contributing to about 100 per cent chromosomal homology between each other¹. The *Mtb* genome is about 4.4 Mbp and contains 0.01-0.3 per cent synonymic nucleotide polymorphisms². Apart from the chromosomal homology, the similarity in the clinical presentation and treatment of these infections has made the study of these organisms more difficult³. These findings suggest that the MTBC may have evolved from a common progenitor⁴.

TB is a growing public health problem among the HIV infected individuals. Around 7 to 8 million new cases of TB are recorded annually with 1.5 to 2 million deaths occurring in about one third of the infected population⁵. It is said that only 10 per cent of the individuals who are diagnosed with TB present a disease pattern that is heterogeneous, suggesting that host factors play a major role in disease vulnerability and natural history⁶.

An understanding about the transmission of TB demands the analysis of distinct epidemiological populations based on universal molecular epidemiological techniques and long term surveillance programmes. The molecular epidemiology (ME) is a combination of both molecular biology and epidemiology, which involves the study

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of distribution of the diseases in human populations, identified at the molecular level⁷. It is a powerful technique for monitoring infectious diseases such as TB, where patients infected with a given strain may undergo relapse due to reactivation of the same strain or a different strain after cure⁸. Not only does ME help in improving our understanding of the pathogenesis of the disease, but also provides unique insights into the international dissemination of tuberculosis by the geographic comparison and evolutionary analysis of highly widespread pathogen populations⁹. In addition, cross-contamination in the laboratories and the risk factors associated with the TB transmission can also be conveniently traced⁷.

There are several available applications of ME in control of TB¹⁰. The first important role of genotyping studies is on intensive use of clustering rate (%) to trace outbreaks¹⁰⁻¹⁴. Modern methods for molecular epidemiological typing of mycobacteria are usually based on the revelation and comparative characteristic of repetitive mycobacterial genome sequences evolving quickly¹⁵. Due to the high resolving capacity and the sensitivity of these methods, the differences and similarities between various strains can be easily revealed, even if the epidemiological information is completely absent. This review article throws light on the ME of *Mtb* with insights into the next generation sequencing (NGS) platforms to differentiate the strains more efficiently.

Polymorphisms within the mycobacterial genome

Earlier, the *Mtb* genome was considered to be stable that lacked polymorphisms¹⁵. Studies carried out in the mid-1990s¹⁶ showed the discovery of monomorphic and polymorphic sites in the genome of the bacteria that can be possibly divided into three groups namely single nucleotide polymorphisms (SNPs), large sequence polymorphisms (LSPs), and polymorphisms in repetitive sequences. When the latter was subjected to subsequent characterization, two classes of DNA were revealed: (i) transposable DNA elements or scattered repeats such as insertion sequences (IS), and (ii) short tandem repeats^{1,17}. Short tandem repeats are further classified into major polymorphic tandem repeats (MPTR); which are basically multiple repeats of 10bp in size that are separated by interspersed units of 5bp and are found in different positions in the *Mtb*

genome and short tandem repeats (STR) containing 49 repeats of 36 bp each separated by short DNA sequences of 41bp in size called as the spacer DNA¹⁸. When compared to SNPs and LSPs, polymorphisms in repetitive sequences tend to evolve more quickly thereby resulting in analogous patterns in *Mtb* species that are completely unrelated¹⁵. Therefore, research on the polymorphisms in repetitive sequences has led to its usage in epidemiological investigations; the other two polymorphisms have also proven to be a useful marker in ME¹⁹.

Genetic markers and genotyping methods of *M. tuberculosis*

IS6110: IS6110, first described by Thierry *et al*¹⁷ is a mycobacterial insertion element that has been consistently used as a genetic marker for typing of *Mtb* species. Originally discovered in *Escherichia coli* and *Shigella* species, this 1.4kb size element belonging to the IS3 insertion sequence family contains three independent sequences that are virtually identical to each other and differ only in a few base pairs^{17,18}. The number of copies of this widespread insertion element varies from strain to strain indicating that it is highly unstable. This instability induces genetic rearrangements at a very high frequency. It can also be attributed to the presence of innumerable amounts of IS6110 DNA sequences in the *Mtb* genome¹⁹. These properties result in fingerprint alterations after a few generations thereby validating its use in epidemiological studies²⁰. Experimental evidences prove that though IS6110 is an unstable element, its transposition events are very rare²¹. Hence, these elements are useful in fingerprinting various strains with exceptions to those strains that lack this element or with those strains that contain only a fewer copies of such elements²².

IS6110 restriction fragment length polymorphism (RFLP) typing: IS6110-RFLP has been widely used since the early 1990s owing to its ability to differentiate between two unrelated strains²³. The genotyping method is based on the variability of the number of copies of IS6110 and the molecular weights of DNA fragments in which the insertions are found²⁴.

The main advantages of the IS6110-RFLP method are its high discriminatory power and the availability of studies for comparison. The main limitation of this method is the low discriminatory power in isolates

presenting five or fewer IS6110 bands. Studies have demonstrated that the frequency of clinically confirmed epidemiologic links between cases is decreased in clusters formed by isolates with six or fewer IS6110 bands^{7,20,25}. For these isolates, secondary markers such as polymorphic GC-rich repetitive sequence (PGRS) and clustered regularly interspersed short palindromic repeats (CRISPRs) are used to increase the discriminatory power and to determine the epidemiological links among patients²⁶. Furthermore, IS6110-RFLP has significant technical limitations, including the need for 2–3 µg of high quality DNA and, therefore, the need of prior culture of the isolates and the determination of results based on visual inspection of images of band patterns that are difficult to share between laboratories⁷.

PGRS-RFLP: PGRS - RFLP has been used as a secondary typing method in isolates of *Mtb* with five or fewer copies of IS6110²⁷. It was first described by Ross *et al*²⁸. Being similar to IS6110 RFLP analysis, it is also a hybridization technique that utilizes the PGRS specific probe (a 3.4 kb fragment of the PGRS sequence) cloned in plasmid pTBN12²⁷⁻²⁹. This has helped in distinguishing strains from unrelated cases of TB and demonstrated identical banding patterns for isolates from epidemiologically related cases^{27,29}. The isolates clustered by IS6110-based RFLP analysis were further discriminated by PGRS typing³⁰. This is particularly the case when IS6110 low copy number strains are further analyzed by PGRS genotyping²⁹. This method, like IS6110 genotyping, is resource intensive¹⁶, but the disadvantages are similar to the technical limitations of IS6110-RFLP, and PGRS-RFLP produces an image with many more bands of different intensities, complicating its reading and interpretation²⁶. It is no longer in use these days.

Mycobacterial interspersed repetitive units-variable number tandem repeat (MIRU-VNTR): Variable number tandem repeat is used as an invaluable genetic marker that provides data in a simple and format based structure on the number of repetitive sequences in polymorphic micro- and mini-satellite regions³¹. Mycobacterial interspersed repetitive units (MIRUs) are a VNTR introduced for *Mtb* by Supply *et al*³². Of the 41 different MIRU loci, 12 loci were identified as hypervariable repetitive units^{10,31,33}. The repeated units are 52 to 77 nucleotides in length and, therefore, power

of this method may be comparable to that of IS6110-RFLP.

MIRUs help in studying the population structure and also in epidemiologic studies by means of the number of repeats of each loci present in various strains³¹. Most important advantage of this method is that it can be applied to *Mtb* cultures without DNA purification³⁴. On the other hand, the difficulty of this method is associated with accurate sizing of multiple small fragments³⁵.

A study based on a worldwide collection of tubercle bacillus isolates defined an optimized set of 24 MIRU-VNTR loci, including a subset of 15 discriminatory loci proposed to be used as a first-line typing method. These 15 and 24 loci sets reliably improved the discrimination of *Mtb* isolates compared to the original 12-locus set^{10,36}.

MIRU-VNTR typing: MIRU-VNTR typing is based on PCR amplification using primers specific for the flanking regions of the different MIRUs²⁶. This high resolution typing method based on the VNTR of MIRUs has been successfully employed in typing the mycobacterial isolates yielding a resolution power close to IS6110-RFLP⁷. Of the 41 loci identified³¹⁻³³, 12 were selected and used for typing of *Mtb* isolates. In this method, each locus is amplified and the product is visualized in a gel. The size of the PCR product will reflect the number of copies of the repeat unit. The result yielded 12-digit number corresponding to the number of repeats at each MIRU loci, forming the basis of a coding system that facilitates inter-laboratory comparisons⁷. There is a technical difficulty of sizing the multiple small PCR fragments, which is overcome by combining multiplex PCR with a fluorescence-based DNA analyzer^{31,32}. A global epidemiological database is available^{37,38} which has led to insights into the distribution and evolution of *Mtb*, including the identification of clonally related families in specific geographic distributions³⁹.

The discriminatory power of MIRU-VNTR analysis is typically proportional to the number of loci evaluated¹⁶. In general, when only the 12 loci are used, it is more discriminating for isolates with low copy number of IS6110 insertions but less discriminating than IS6110-RFLP genotyping for isolates with high copy number of IS6110^{7,16,26} especially with the Beijing spoligotypes (ST)^{36,40}. When more than 12 loci are used, or MIRU analysis is combined with spoligotyping, the

discriminatory power approximates that of IS6110 RFLP analysis¹⁶. A set of 15 and 24 MIRU loci is currently recommended for molecular epidemiologic phylogenetic studies⁴¹.

It has been shown that 24 MIRU loci sets are specially recommended to discriminate the Beijing isolates⁴² as the set of 15 MIRUs was not sufficient to discriminate them^{40,42}. A recent study done by Allix-Béguec *et al*⁴³ showed that the set of 24 loci MIRU-VNTR lacked the sufficient discriminatory power to differentiate specifically Beijing isolates as these seemed to be hypervariable and were associated with the spread of multi drug resistant (MDR) strains. An additional seven hypervariable MIRU-VNTR loci were identified by this group, which produced a better resolution and reduced the clustering rate for the Beijing strains⁴³. The MIRU-VNTR method is also considered to be the gold standard for typing the *Mtb* isolates⁴⁴.

Direct repeats (DR) or clustered regularly interspersed short palindromic repeats (CRISPR) locus: The DR region in MTBC strains is composed of multiple direct variant repeat sequences (DVRS) each of which is composed of 49 repeats of 36 bp DR and a non-repetitive spacer sequence of 35 - 41 bp in size^{1,7}. This unique cluster of DR is located in a hot spot for integration of IS elements¹. There is extensive polymorphism in the DR region by the variable presence of DVRS which is probably driven by homologous recombination between adjacent or distant chromosomal DRs^{7,16}. As a result of these events, some spacers may be deleted from the genome¹⁶.

Spacer oligonucleotide typing (Spoligotyping): Spoligotyping is the most commonly used PCR-based technique for differentiating *Mtb* strains⁴⁵. This method is based on revealing the presence and order of location of spacers that separate DRs in a specific locus of the *Mtb* genome. Forty three types of mycobacterial spacers have been revealed, of which 37 types are typical *Mtb* and another six types additionally characterize the *M. bovis* BCG strain¹⁵. In practice, membranes are spotted with 43 synthetic oligonucleotides which hybridize the PCR amplified DR locus of the tested strain that is labelled. This results in a pattern that can be detected by chemiluminescence. These patterns reveal absence or presence of the spacers⁷ and are read in the form of a binary that can be easily interpreted and computerized¹⁶.

An edition of the international spoligotyping database namely SpolDB4/SITVIT⁴⁶ was introduced containing 1,939 different spoligotypes (ST) identified. The disadvantage of the database is that the actual version of SITVIT/SpolDB4 does not offer functionality for more complex analysis such as the tree based analysis⁴⁶. Also, some important profiles such as Latin American Mediterranean (LAM), Haarlem (H), *etc.* are still assigned to wrong families even by improved decision rules of SITVIT database⁴⁶.

Apart from being simple, rapid, robust and an economical means for typing MTBC^{34,35,47,48}, this method has other similar advantages as that of the IS6110 RFLP method in that it can be performed with small amount of DNA and a little time after inoculation of bacteria into liquid culture³⁴. Spoligotyping is useful for discrimination between isolates of *Mtb* with a few copy number of IS6110⁴⁹. Although spoligotyping can be a method to study the molecular epidemiology of *Mtb*, but the differentiating power of spoligotyping is less than IS6110 typing when high copy number strains are being analyzed^{7,48,50}.

It has been shown that strains having identical ST patterns with distinct IS6110 fingerprint profiles are often encountered^{16,51}. For example, the W-Beijing family of strains which are large phylogenetically related group of *Mtb* isolates comprising hundreds of similar yet distinct IS6110 variations, has an almost identical spoligopattern lacking spacers 1 through 34⁵². In such cases, spoligotyping may be useful in identifying W-Beijing strains in a population. Kremer *et al*⁴⁸ have shown that spoligotyping together with IS6110 genotyping can provide an accurate and discriminatory genotyping system.

Single nucleotide polymorphisms (SNPs): Research in genetic polymorphisms at the nucleotide level has revealed certain genetic markers that not only help in differentiating various clinical isolates but also in studying the phylogenetic relationship between these strains. SNPs can be categorized into two types namely: nonsynonymous SNP (nsSNP) and synonymous SNP (sSNP)^{16,53}.

In general, nsSNPs cause changes in the amino acids and also in the genetic loci that determine the drug resistance, thereby resulting in changes in the internal or external pressure and phenotypic drug resistance^{15,16}.

These nsSNP genes conferring in drug resistance can aid in understanding the spread and the nature of the drug resistant isolates within the populations.

In contrast, sSNPs are considered functionally neutral as these do not alter the amino acid profile. These neutral alterations, when in structural or housekeeping genes, can provide the basis to study genetic drift and evolutionary relationships among mycobacterial strains¹⁶. Apart from helping in the study of phylogenetic relatedness of clinical isolates, sSNP also serves to determine the different epidemiologies in a given population.

SNP analyses are amenable to targeting multiple polymorphisms that are informative in phylogenetic grouping, drug resistance, virulence, and other epidemiologically instructive markers, but their low discriminatory power limits their use¹⁶. In a study done by Homolka *et al*⁵³, 26 different genes were sequenced to find 161 polymorphisms, of which, 59 were genotype specific and 13 polymorphisms defined deeper phylogenetic branches. Study of the most variable set of 11 genes in a set of population from Germany, validated the SNP analysis with high accuracy.

Large sequence polymorphisms (LSPs): Comparative genomic analysis of strains H37Rv and CDC1551 has revealed LSPs in addition to SNPs⁵⁴. LSPs are thought to mainly occur as a result of genomic deletions and rearrangements rather than through recombination following horizontal transfer^{16,55}. In the absence of horizontal gene transfer, deletions are irreversible and, therefore, these have been proposed for genotyping as well as for constructing phylogenies^{16,56}.

A deletion occurring in the progenitor strain can serve as a genetic marker to differentiate various strains and to aid in the epidemiological analysis. Some chromosomal deletions are associated with IS transpositions and the deletions occurring in these regions are called as the regions of difference (RD). There are two types of deletions namely: ancient and recent. The ancient deletions occur at different stages in the speciation process and are widespread whereas the recent deletions have a more restricted distribution⁵⁷. For instance, IS6110 mediated deletion of the 7 kb locus RvD2 in *Mtb* H37Rv is still present in the closely related avirulent derivative H37Ra⁵⁸. This region undergoes great variability in clinical isolates of *Mtb* and seems to

represent a hot-spot for IS6110 transposition events⁵⁹. Other RD loci such as RD9, RD105, RD 207 and RD239 have also been studied⁴¹. It has been revealed that RD105 is a genetic marker to identify Beijing strains⁶⁰. But, this was proved otherwise in another study that RD105 was a marker for East Asian lineage⁴¹ whereas RD207 was the true marker for Beijing strains. Another interesting study done by Fenner *et al*⁶¹, showed certain pseudo Beijing characteristics in three different strains that were previously classified as Beijing strains. Three different genetic markers were used to analyse them and the study concluded that phylogenetically robust markers should be used to differentiate clinical or experimental phenotypes. From close inspection of the DNA sequences bordering these RD regions it is apparent that deletions occur within coding regions also.

Another example of LSPs is the deletion of TbD1, a 2,153bp *Mtb* specific deletion 1 fragment which was identified in all *Mtb* strains but was absent in the ancestral strains⁶². It is found that these deletions tend to aggregate and are not always randomly distributed in the chromosome⁶³. The correlations between IS elements and the deletion regions such as TbD1 have not yet been determined¹⁶. Using these deleted fragments as genetic markers, this analysis can be performed by simple PCR-based methods⁶³. A recent study performed in Northeast Thailand by Faksri *et al*⁶⁴ showed the classification of *Mtb* based on LSP using multiplex real-time PCR which provided a simple, rapid and high performance tool for characterizing *Mtb* based on LSPs.

Deletion analysis: Deletion analysis can be very efficiently used for epidemiological investigations especially when the presence of a specific deletion (LSPs) associated with a single strain has been predetermined. In these cases, a single PCR may suffice to track down the spread of a single strain^{65,66}. However, in studies where no particular clone or strain has been predetermined, simultaneous analysis of multiple deletion regions is required⁶⁷.

A high-throughput method for detecting large polymorphic deletions was developed⁶⁸. In this method, 43 genomic regions for large scale deligotyping analysis were selected and PCR was performed. The PCR products generated from these 43 deligotes were hybridized to a membrane containing the target

Table I. Evaluation of various genotyping methods used in molecular epidemiology (ME) of tuberculosis

Marker	Principle	Advantages	Disadvantages
IS6110-RFLP typing ⁷	Based on the number of copies of IS6110 elements present and the molecular weights of the DNA fragments in which the elements are found.	High discriminatory power for isolates having higher IS6110 copies, widely used till date.	Low discriminatory power in isolates containing five or fewer IS6110 copies, lengthy process, inter-laboratory comparisons are difficult.
PGRS-RFLP ¹⁶	Based on the number and the location of the PGRS regions within the genome.	Discriminates isolates having ≤ 5 copies of IS6110 elements, resource intensive.	Analysis is difficult, lengthy process, limited data available using this technique.
Spoligotyping ³⁴	Based on the spacers present between the 36bp direct repeat sequences in the genome.	Simple, rapid and robust, highly reproducible PCR based method, requires low quantities of DNA, data available in exchangeable format.	Limited discriminatory power.
MIRU-VNTR ⁴¹	Based on the polymorphisms of MIRU loci within the genome.	Highly discriminatory, data available in exchangeable format, 24 loci used for ME and phylogenetic studies.	Analysis of the DNA by electrophoresis is less reproducible than sequencer based method. Sequencer based method is expensive.
Deletion analysis ^{7,16,68}	Based on the detection of deletions in the selected 43 genomic regions.	High throughput with microarray analysis, data available in exchangeable format, irreversible genetic marker used.	The target deletions need to be pre-determined, technique has to be evaluated in different settings.

RFLP, restriction fragment length polymorphism; PGRS-RFLP, polymorphic GC-rich repetitive sequence-restriction fragment length polymorphism; MIRU-VNTR, mycobacterium interspersed repetitive units-variable number of tandem repeats.

sequences of the 43 loci. Amplification of the selected deletion region is possible by using the flanking regions and usage of the flanking regions has shown to increase the discriminatory power of the technique⁶⁸. This approach proved to be highly sensitive and efficient for the rapid screening of clinical isolates¹⁶.

The most recent method to differentiate the strains based on the deletions within the genome is the deletion microarray. In this method, the microarray used to compare the genome of a strain is compared against that of a sequenced reference strain⁷. This helps to identify the deletion that has occurred in the sample genome. The number and distribution of these deletions help in phylogenetic/evolutionary studies, facilitation of genome structure-function studies, host-pathogen interactions based on specific genomic deletions, in molecular epidemiology both on phylogenetic relationships and information about the various phenotypes^{7,16}. Table I evaluates the various genotyping methods used in epidemiologic studies.

Whole genome sequencing (WGS)

The genotyping helps to determine the patients involved in the part of chain of transmission but does not allow to distinguish the chain of transmission of events²⁶. One of the important breakthroughs in the field of ME is the WGS technology which involves the shearing of the DNA to several distinct sizes that are sub-cloned in plasmids. The sub-clones are then over sampled to generate sequencing reads which provide the necessary information to perform whole genome assembly algorithms. This method is relatively faster and affordable⁶⁹.

One of the widely used WGS methods is the Sanger's method of sequencing. Sanger sequencing is method of sequencing DNA which was developed by Frederick Sanger and his co-workers⁷⁰. It uses the ABI 3730xL platform for sequencing and has been employed in sequencing large scale projects⁷¹. The main disadvantage of this method is that it includes poor quality in the first 15-40 bases of the sequence

Table II. Comparing performance characteristics of various next generation sequencing (NGS) platforms

Properties	Roche (454)	Illumina	ABI SOLiD	ION PGM	Heliscope	Pacific Biosciences
Sequencing chemistry	Pyrosequencing	Sequencing by synthesis	Ligation based sequencing	Semiconductor sequencing	Single molecule approach	Real time single molecule approach
Accuracy (%)	99	99.9	99.94	99	99.5	99.999
Millions of reads/run	1	3.4	>700	0.10	800	0.01
Run time ^a	10 h	26 h	~ 1-2 wk	~2 h	~1 day	~2 h

^a Run time depends on the length of the genome to be sequenced. Here, it is considered to be the length of the human genome.
Source: Refs 74-78

due to primer binding and deteriorates the quality of sequencing traces after 700-900 bases.

Niemann *et al*⁷² performed a study using WGS on two *Mtb* Beijing family clinical isolates that had matching IS6110 RFLP pattern, ST pattern and a similar MIRU-VNTR profile. With the help of WGS results, the authors were able to discriminate the two isolates that differed in 130 SNPs and one large deletion thereby suggesting that the epidemiological link between the two isolates may have been remote⁷².

WGS is considered as an important tool to determine sequence variation at a real epidemiological scale, to determine the evolutionary relationship of the strains and also to determine the source of infection and the transmission of the disease between various patients²⁶. WGS may become the gold standard for

typing various strains for ME in the near future^{26, 73} but, there are certain limitations such as the need for specialized software to analyze the various sequence reads produced and the incomplete understanding of the various polymorphisms such as the SNPs and LSPs²⁶. Hence, the Next Generation Sequencing (NGS) technologies are introduced which produce millions of short reads of the entire genome that are then analyzed by specialized softwares. These do not require any cloning of the template DNA into the bacterial vectors and are optimally suited for re-sequencing⁷³.

Next generation sequencing (NGS) technologies

Several NGS technologies have emerged which generate a magnitude of 3 to 4 times more sequence and are considerably less expensive than the Sanger's method of sequencing⁷⁰. Various platforms of NGS for

Table III. Advantages and disadvantages of the various next generation sequencing (NGS) platforms

NGS platform	Primary advantages	Primary disadvantages
Roche (454)	Produces maximum read length.	Very expensive – high cost per Mb
Illumina	Versatile instrument and scalable in future.	Relatively few reads and higher cost / Mb
ABI SOLiD TM	Low cost of the instrument and high accuracy.	Takes a long time for sequencing and cost per read is high.
ION PGM	Simple and a low cost instrument can be easily upgraded.	New platform with low accuracy.
Heliscope TM	Single molecule approach produces large number of reads.	Accuracy and longevity of this approach remain questionable.
Pacific Biosciences	Real time sequencing through single molecule approach produces large number of reads; the run time is very low thereby enabling sequencing of large number of samples.	High capital cost.

Mb, mega base pair
Source: Refs 74,75,78

the production of massively parallel DNA reads are being used widely [Roche/(454) FLX, Illumina/ Solexa Genome Analyzer, Applied Biosystems SOLiD™ System, Helicos Heliscope™, Ion Personal Genome Machine (Ion PGM), Pacific Biosciences Single Molecule Real Time (SMRT) and nanopore sequencing instruments]. These instruments allow simple and stepwise preparation of the sample prior to DNA sequencing thereby saving time⁶⁹. The performance characteristics and the primary advantages and the disadvantages of the various NGS platforms are described in Tables II and III, respectively.

The strains can be easily distinguished from each other due to small variations in sequence of the DNA. Moreover, more complex determination of SNPs and LSPs can be effortlessly identified. In southern India, NGS has paved way in identifying new *Mtb* isolates having completely different DNA sequences from the ones that have already been characterized. For example, WGS of clinical isolates of *Mtb* from Kerala, Andhra Pradesh and other parts of south India were characterized⁷⁹⁻⁸¹.

Apart from these, NGS is also used in the field of comparative genomic studies and in transmission dynamics. Ever since the sequencing of the entire mycobacterial genome is done, investigators have constantly used low and high resolution comparative genomic techniques to identify small differences between various strains⁸². With the evolution of NGS, this application has made differentiation between various strains much easier. This has been proved in a recent study where NGS platform was used to reveal the genetic heterogeneity of *Mtb* in extra pulmonary TB patients⁸³. Transmission dynamics has become an important aspect to understand the pathogenesis of the disease. A retrospective observational study was done using WGS to establish the relapse and the reinfection of TB in patients. This study concluded that WGS has more resolving power than any other genotyping methods and also may define end points for clinical studies⁸⁴. Complex evolutionary patterns of MDR *Mtb* Beijing strains in patients have been revealed using WGS⁸⁵.

Roche/454 FLX pyrosequencer: The 454 GS20 pyrosequencing platform has been replaced by GS FLX platform in 2005. It is the first generation high

throughput sequencing technology⁷³. The Roche 454 sequencer (Roche, USA) is based on the principle of pyrosequencing which involves the release of the pyrophosphates due to the incorporation of nucleotides during the PCR reaction. The pyrophosphate so released sets out a series of downstream reactions that lead to the production of light by means of the enzyme luciferase⁷³.

In this method, single stranded DNA library is constructed by fragmentation of the genomic DNA and ligation of the fragments to the adaptor sequences. The ligated products are selected based on avidin-biotin purification and the library is created. This step takes approximately 4.5 h and does not involve the cloning of the genomic DNA into plasmids. The fragments are then mixed with agarose beads whose surfaces carry oligonucleotides complementary to the adaptor sequence present in the fragment library. PCR amplification is carried out by subjecting individual fragment : bead mixture in oil : water emulsion containing all the PCR reactants. The emulsion PCR step takes about eight hours to complete. The fragments are amplified as a whole on a picotitre plate where each bead is placed in a single well. With the addition of nucleotides during PCR amplification, the pyrophosphates are released and the light is produced⁶⁹. The produced light is recorded as an image for analysis⁷³.

The Roche/454 FLX instrument provides 100 flows of each nucleotide during an 8 h run, which incorporates an average of 2.5 bases per flow thereby producing an average read length of 400 nucleotides. These raw reads analyzed by the software and are screened to remove poor quality sequences. Finally, 100 MB of quality data on average are obtained⁶⁹. The main disadvantage of this method is that it incorporates more than one nucleotide to the complementary strand in one cycle and lacks reversible nucleotide terminator to stop the addition of nucleotide. It is also said to have certain problems in resolving homopolymeric stretches of sequences⁷³.

Illumina/solexa Genome analyzer: Illumina genome analyzer (Illumina, USA) uses sequencing by synthesis approach that involves amplification of a single nucleotide. In this method, the single stranded genomic DNA library is created where the fragments are ligated with the adaptor sequences. The creation of the library takes place on the oligo derivatized solid support of a flow cell which is done automatically by a device called

as a cluster station⁶⁹. The flow cell is an 8-channel sealed glass device where the DNA polymerase enzyme allows the generation of *in situ* copies of DNA molecule on the oligo decorated surface (Bridge Amplification). Fluorescent labelled dNTPs are added where the 3'OH group is blocked such that each addition occurs as a unique event⁶⁹. An image is taken after every addition of the base to the growing strand. This is followed by the chemical removal of 3' blocking group which prepares the strand for the next base incorporation.

The sequencing is carried for approximately four days and the computed data filters to remove the poor quality reads. Around 40-50 million reads are produced for every run⁷¹. The amplification step used in this method primarily reduces the background noise and increases the levels of the signals produced. On the other hand, inaccuracies can be caused if the simultaneous stepwise sequencing of the molecules is disturbed due to non-synchronous behaviour of individual molecules⁷³.

Applied Biosystems SOLiD system: The SOLiD platform uses adaptor ligated fragment library and an emulsion PCR technique similar to other NGS platforms⁶⁹. The DNA amplified on the beads undergoes two cycles. The first cycle involves the deposition of the beads on to a glass slide which is eventually subjected to hybridization with random fluorescent labelled oligonucleotides containing known 3' dinucleotide. The synthesized DNA is removed from the template after five cycles and the process is repeated. The second cycle starts at the upstream region on the template where the synthesis began in the previous cycle. This type of repeated cycle process tends to reduce the error rates and allows for multi-colour analyses of the bases in the DNA sequence⁷³. Complicated DNA variations and deletions such as SNPs and LSPs can be easily detected. AB SOLiD system can detect read lengths between 25-50 nucleotides and the run yields a data of 2-4GB^{69,73}.

The advantage of SOLiD system over the other platforms is that it has slightly better performance in terms of accuracy as the bases are sequenced twice by dinucleotide detection⁸⁶. Apart from having similar limitations as that of the Illumina platform⁷³, a complicated algorithm is required to interpret the raw data⁸⁶.

Helicos Heliscope: Helicos Heliscope (Applied Biosystems, USA) uses single molecule approaches.

This kind of approach avoids amplification and errors and bias and other intensity and phasing related problems⁷³. Similar to the above mentioned platforms, Heliscope sequencing also involves the creation of adaptor ligated DNA fragment library. These fragments remain unamplified and are attached to a solid substrate. Fluorescent labelled oligonucleotides are added to the unamplified DNA fragment by the DNA polymerase enzyme to create a second strand DNA. A virtual terminator helps in preventing the addition of extra nucleotides in a cycle. Image is taken with the amount of fluorescence sensed and the nucleotide from each DNA sequence can be determined. The fluorescent molecule is then cut away, and the process is repeated until the fragments have been completely sequenced⁸⁷. The background noise is greatly reduced due to the use of fluorophore at the start of each cycle. Currently, reads have a final length of 35 nucleotides⁷³.

Ion Personal Genome Machine (Ion PGM): One of the recently introduced NGS systems includes the Ion PGM (Life Technologies, USA). This method is based on the principle of release of hydrogen ions from nucleotides which is sensed by an array of semiconductor chips that are capable of sensing minor changes in pH⁸⁶. Generally, when a nucleotide is added to the growing DNA template by the polymerase enzyme, a proton is released. The PGM recognizes the addition of nucleotide by the release of the proton which induces a pH change. When there is an incorrect base added, there is no voltage difference observed whereas when two nucleotides are added at the same time, a double voltage is detected⁸⁸.

This platform is the first NGS technology that does not require fluorescence and imaging techniques resulting in higher speed, lower cost, and smaller instrument size. The preparation of the sample takes about six hours for eight samples in parallel and the read length produced is about 200bp^{86,88}.

Pacific Biosciences Single Molecule Real Time (SMRT): The Pacific Biosciences SMRT platform (USA) also uses single molecule approach and is considered to be the most revolutionary technique⁷³. In this method, the DNA polymerase is immobilized in a zeptolitre wells (10^{-21} wells)⁷³. The single stranded DNA is added to the wells containing the immobilized enzymes along with the phospho-linked nucleotides and other PCR reactants. The complementary strand

is synthesized by the addition of nucleotides. Since the wells are only nanometers in size, they accumulate the nucleotides leading to a highly focused and continuous detection process.

This platform gives various advantages in that the newly introduced base can be read in a short period of time, has high accuracy and low background noise due to the use of phospho-linked nucleotides⁷³, and large inserts can be used⁸⁶. This method detects the addition of nucleotides in real time thereby paving way for novel applications⁸⁶.

Computational resources for data analysis

The computational softwares necessary to analyse the data vary tremendously for every platform. High end softwares are required to analyse the short reads and most of the platforms produce millions of short reads. To analyse these short reads, generally high end computers are used. Some softwares can work only on LINUX based systems whereas a few others can work in Windows or in Mac systems. This variability poses a great disadvantage for the users to analyse the short reads. For the analysis of high amounts of data, high performance computational clusters produced by Illumina can be used which is less expensive and can be operated on any computer⁷⁴.

Nanopore sequencing

Nanopore sequencing is an upcoming sequencer and is classified under third generation sequencing. The biological role of a nanopore is to facilitate the ion exchange through the protein channels which are embedded on the lipid layer⁸⁹. The basic principle of nanopore sequencing method is to insert a thread of single stranded DNA through the nanopore such as α -haemolysin (α HL) which is isolated from *Staphylococcus aureus*⁹⁰. This 33kDa protein undergoes self-assembly to form heptameric ion channel⁷⁴ which can tolerate a large voltage of up to 100mV⁹⁰. By the exploitation of this unique property, external current is applied constantly and the disruption of the current is detected by electrophysiological technique. The results are based on the size difference between deoxyribonucleoside monophosphates (dNMP). For every size of dNMP, there is a modulation in the current. The ionic current is resumed when the trapped nucleoside comes out of the pore⁸⁸.

Another biological nanopore being investigated for DNA sequencing is the *Mycobacterium smegmatis* porin A (MspA). This particular molecule has been identified as an alternative to α HL molecule due to its structure⁹¹. Generally it has a negative core that blocks the translocation of single stranded DNA but this has been modified by replacing three negatively charged aspartic acids with neutral asparagines thereby allowing the translocation process to take place⁹². The detection of the nucleotides and the identification of the bases by the application of the electric current across the membrane have shown to be ten folds higher than α HL⁹¹. Further studies are currently carried out in the United States to improve its specificity and also the base recognition of the MspA molecule⁹³.

Nanopore sequencing will enable sequencing of large DNA molecules in minutes without modifying or preparing samples. It is expected to offer solutions to limitations of other NGS technologies⁸⁶.

Current applications of molecular epidemiology

Apart from a few applications described earlier, several other applications for ME are available in the control of TB. Epidemiological data are necessary to understand the mechanism of outbreaks of any disease. But, based on the epidemiological data alone, one cannot have an insight into the transmission of TB. In these cases, ME allows the use of the genotyping studies which are extensively used to predict the clustering rates of species, identification of the risk groups and to monitor the outbreaks^{12-14,23,47}. Evaluating the spread of drug resistant strains in patients can also be studied with the help of ME⁹⁴.

HIV infection exerts immense influence on the natural course of TB disease. Individuals with latent *Mtb* infection who contract HIV are at risk of developing active TB at a rate of 7 to 10 per cent per year, compared to approximately 8 per cent per lifetime for HIV-negative individuals⁹⁵. With the help of ME, diseases involving co-infection with HIV can be improved²⁶.

Finally, with the advent of WGS technologies, various polymorphisms in the genome of strains are detected in no time, thereby making ME an easier method to detect the phylogenetic and evolutionary relationships between strains²⁶. Even though *Mtb* strains lacks genetic diversity due to low mutation rate, it is surprising that in the face of rapid emergence of MDR

and extensively drug resistant (XDR) strains, there is evidence of diversity of *Mtb in vivo*. The WGS analyses are providing insights into the ongoing evolution of *Mtb* during infection, treatment and acquisition of drug resistance⁹⁶. When WGS was done on serial isolates of patient, it has revealed that resistance mutations to the drugs are independently acquired several times by different isolates⁹⁷. By sequencing to a high depth of coverage we could identify mutations at specific loci that are present in a proportion of the bacteria sequenced⁹⁸.

Conclusion

ME has contributed extensively to our current knowledge of TB through various observational studies done since the early 1990s. These methods have provided better accuracy and understanding in the global epidemiology of TB. Furthermore, various characteristics specific to phylogenetic lineages or strains comprising virulence properties, different replication rates and differential pathogenesis, have been widely found and suggested. With the advent and widespread availability of NGS systems, DNA sequencing has become a universal readout for a wide variety of front-end assays. There is a need for the development of novel designs and other mathematical models for analyzing the epidemiologic results obtained. An understanding of host pathogen interactions and identification of host susceptibility genes will be continuously integrated due to the current advances in designing ME studies. This will enhance the value of *Mtb* biology and epidemiology and perhaps unravel the mystery behind this elusive pathogen.

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