Review

Genetic variation and DNA markers in forensic analysis

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In this review, we highlighted the DNA markers and their applications in forensic medicine which included the study of genetic variation or polymorphism, markers for genetic variation, types of variation, diallelic polymorphisms (SNPs and Indels), haplotypes, autosomal short tandem repeat (STRs), Y-chromosome short tandem repeat (Y-STRs) and Y chromosome. This is a useful tool for tracing human evolution, analysis of Y-chromosomal microsatellite haplotypes in globally human populations, organization of the human mitochondrial genome, and mitochondrial coding region as a source for variability and interpretation of sequence data. The light has been focused and directed in this study to establish the basic forensic genetic information, knowledge, data and statistics which might be so ultimately helpful practically in forensic science and criminology and to let evaluate and present the DNA weight evidences in medico-legal institute and courts of law.

Key word: Applications, DNA markers, forensic, genetic variation, review.

INTRODUCTION

Every organism stores its information on identity in its DNA (Dobbs et al., 1994). Separate chromosomes which are available in the nucleus of cells of humans have DNA. Serological and DNA typing are being used in forensic biology to analyze evidence found at crime scenes. The sources biological evidences may be bodily fluids or tissues (Pierce and Wangh, 2007).

Microsatellites are a group of molecular markers chosen for a number of purposes which include forensic individual identification and relatedness testing.
(Yamamoto et al., 1999; Nakamura et al., 2009). Low quantities of template DNA are required (10-100 ng) (Markoulatos et al., 2002), when using microsatellites. There is a high genomic abundance of random distribution throughout the genome. There is also an abundance of polymorphism. A nuclear DNA present in one copy per cell and only in males is called the Y chromosome. It includes the sex determining region and known as a paternal lineage marker (Butler et al., 2002; Carolina et al., 2010; Kuppareddi et al., 2010). The genetic information is inherited from the father to the son, and this information does not change except for mutational events (Hanson and Ballantyne, 2007; Mohammad and Imad, 2013). The individual short tandem repeat (STRs) are inherited as a single unit because of the lack of recombination which is called a haplotype and behaves as single allele per individual (Parson et al., 2003; Kwak et al., 2005).

The Y - chromosome is specific to the male portion of a male-female DNA mixed such as is common in sexual assault cases (Park et al., 2007). These STRs can also be useful in missing persons investigations, historical investigations, some paternity testing scenarios, and genetic genealogy (Park et al., 2007; Andrea et al., 2008). Although they are often used to suggest which haplogroup an individual matches, STR analysis typically provides a person haplotype. Most tests on the Y chromosome examine between 12 and 67 STR markers (Kayser et al., 1997; Imad et al., 2013a). The Y chromosome is less variable than the other chromosomes. Many markers are thus needed to obtain a high degree of discrimination between unrelated males (Kuppareddi et al., 2010; Hanson and Ballantyne, 2007).

The mitochondrial DNA (mtDNA) is a small circular genome located within the mitochondria in the cytoplasm of the cell. The mitochondrial genome can be divided into two sections: a large coding region, which is responsible for the production of various biological molecules involved in the process of energy production in the cell, and a smaller 1.2 kb pair fragment, called the control region. It is found to be highly polymorphic and harbors three hypervariable regions (HV): HV1, HV2 and HV3 (Kraytsberg et al., 2004; Imad et al., 2013b). Mitochondrial DNA Comprising of about 37 genes coding for 22 tRNAs, two rRNAs and 13 mRNAs are a small circle of DNA (Helgason et al., 2003).

Mitochondrial DNA does not recombine and thus there is no change between parent and child, unlike nuclear DNA. mtDNA is only passed on from mother to child and this is an important fact (Ingman et al., 2003; Ukhhee et al., 2005). There is more sequence divergence in mitochondrial than in nuclear DNA (Brown et al., 1993; Giulietta et al., 2000). This may be caused by a faster mutation rate in mtDNA that may result from a lack of repair mechanisms. Sequencing of highly polymorphic segments of the control region of mitochondrial DNA (mtDNA) is today a routine method of analysis of biological traces which are not suitable for STR analysis due to insufficient concentration of nuclear DNA or heavy degradation processes (Holland, Parsons, 1999; Tzen et al., 2001). A promising approach in this context seems to be analysis of selected single nucleotide polymorphisms (SNPs) that are useful for identification purposes.

GENETIC VARIATION OR POLYMORPHISM

The following: effective population size, population history (migration, bottleneck, and recent expansion), population structure and location of diseases genes are determined by the amount and nature of genetic variation in a population. There are three items: (1) changes in nucleotides which could be transition or transversion. In the transition mutation, a pyrimidine (C or T) is substituted by another pyrimidine, or a purine (A or G) is substituted by another purine. The transversion mutation involves the change from a pyrimidine to a purine, or vice versa; (2) insertion or deletion of single nucleotides (indel) and (3) variation in number of repeat of tandemly repeated sequences (microsatellite, minisatellite and satellite) are genetic variation or polymorphism (Roach, 2010; Conrad, 2011).

The study of human genetic variation has both evolutionary significance and medical applications. It can help scientists understand ancient human population migrations as well as how different human groups are biologically related to one another. For medicine, study of human genetic variation may be important because some disease-causing alleles occur more often in people from specific geographic regions (Tishkoff and Verrelli, 2003; Conrad, 2011).

Diallelic polymorphisms (SNPs and INDELS)

Single-nucleotide polymorphisms (SNPs), the occurrence of different nucleotides at a specific place in the genome, are the most common type of human DNA sequence variation, occurring on average 1 per 500 to1000 bp on a randomly selected chromosome. As noted later, this number varies somewhat depending on the population studied. Insertion/deletion polymorphisms (Indels), the occurrence of some more or fewer nucleotides at one position of the genome, are less common but do occur frequently.

The inserted/deleted sequence can be a few nucleotides long (Weber et al., 2002) up to several hundred nucleotides long, as is the case with the transposable Alu elements (Carter et al., 2004). Some very large polymorphic duplications, hundreds of kilobase pairs long, have also been identified (Sebat et al., 2004; Iafrate et al., 2004). Because these polymorphisms are diallelic, they are less informative than microsatellites; however, SNPs are more common in the genome and more
of the RFLPs (restriction fragment length polymorphisms) defined between the late 1970s and the late 1990s were SNPs in a restriction site but some Indels were occurring between restriction sites. That is, they are caused by a single mutational event that occurred once in the history of a species. We do not expect to see recurrent mutations at the same site except as extremely rare events. For the polymorphic Alu elements, we know that the ancestral state is absence and the insertion of the Alu is the derived state. For the other markers, we cannot tell from the human polymorphism which allele is ancestral and which is derived. However, we can determine the ancestral state in almost all of those cases simply by determining the genotype of our nearest relatives, the other great apes, following the logic in most cases. Humans share a single allele with the other apes and, by inference this is the ancestral or original allele in humans and the other allele is the derived allele (Hacia et al., 1999).

Multiallelic polymorphisms (STRPs and VNTRs)

Microsatellites consist of approximately 10-50 tandemly repeated copies of particular DNA sequence motifs ranging from 1 to 10 (most commonly 2-4) nucleotide base pairs. These repeat sequences, discovered in 1989 (Litt and Luty, 1989), occur frequently and randomly across the human genome. When the repeat number is polymorphic, microsatellites are also called short tandem repeat polymorphisms (STRPs) (other acronyms have also been used, for example, SSLP). STRP loci usually have multiple alleles and can have high levels of variation, that is, high heterozygosity. They rapidly replaced RFLP markers in gene mapping (especially disease genes) studies owing to these features and to their ease of typing, including the small amount of template DNA required. Remember that heterozygosity can never exceed 50% for a diallelic marker, whereas STRPs can easily have heterozygositities >75%. Because of this high heterozygosity, STRPs are the markers on which the most detailed human linkage maps are based (Dib et al., 1996; Kong et al., 2002; Jorgenson et al., 2005). STRPs have also become the standard for forensics and paternity testing (Budowle et al., 2001). The larger minisatellite arrays (also referred to as VNTRs, variable number of tandem repeats) are also highly polymorphic and powerful markers in forensic and paternity studies (DNA fingerprinting) (Jeffreys et al., 1985; Armour et al., 1996). However, they are less common than STRPs and are not evenly distributed throughout the genome. Their larger sequence motifs make them less amenable to PCR technology and use in genomic screening analyses. STRPs and VNTRs, on the other hand, tend to be much more dynamic than the diallelic SNPs and Indels. The mutation rates are higher, there are more alleles, and it is usually impossible to determine which is the ancestral allele.

AUTOSOMAL SHORT TANDEM REPEAT (STRs)

Microsatellites refer to DNA with varying numbers of short tandem repeats (Allor et al., 2005; Klintschar et al., 2006) between a unique sequence (Figures 1 and 2). DNA regions with repeat units that are 2 to 7 bp in length or most generally short tandem repeats (STRs) or simple sequence repeats (SSRs) are generally known as microsatellites (Ellegren, 2004). In the core repeated bases, long repeat units may contain several hundred to thousands (Butler and Hill, 2012). Within the DNA there are length and sequence polymorphisms (Silvia et al., 2009). DNA can be used to study human evolution using human genome analysis regions that are not subjected to selection pressure (Mats et al., 2007). Besides, information from DNA typing provides vital information in medico-legal with polymorphisms allowing for more biological studies (Walkinshaw et al., 1996). It has been found that microsatellites are evenly distributed in the genome on all chromosomes and all regions of the chromosome (Ensenberger et al., 2010). They can also be found inside gene coding regions, introns, and in the non-gene sequences. Most microsatellite loci are really small, ranging from a few to a few hundred repeats and this small size of microsatellite loci is important for PCR-facilitated genotyping. Basically microsatellites containing a higher number of repeats are more polymorphic.

STR LOCUS NOMENCLATURE

The nomenclature for DNA markers is fairly straightforward. If a marker is part of a gene or falls within a gene, the gene name is used in the designation. For example, the STR marker TH01 is from the human tyrosine hydroxylase gene located on chromosome 11. The '01' portion of TH01 comes from the fact that the repeat region in question is located within intron 1 of the tyrosine hydroxylase gene. Sometimes the prefix HUM- is included at the beginning of a locus name to indicate that it is from the human genome. Thus, the STR locus TH01 would be correctly listed as HUMTH01 (Butler, 2005). DNA markers that fall outside of gene regions may be designated by their chromosomal position. The STR loci DSS818 and DYS19 are examples of markers that are not found within gene regions. In these cases, the ‘D’ stands for DNA. The next character refers to the chromosome number, 5 for chromosome 5 and Y for the Y chromosome. The ‘S’ refers to the fact that the DNA marker is a single copy sequence. The final number indicates the order in which the marker was discovered and categorized for a particular chromosome. Sequential numbers are used to give uniqueness to each identified
Figure 1. Exact physical location of 13 STR markers called CODIS: D2S1338 (q35), D3S1358 (3p21.31), D5S818 (5q23.2), D7S820 (7q21.11), D7S1517 (7q31.33), D8S1179 (8q24.13), D8S1132 (8q23.1), D12S391 (12p13.2), D13S317 (13q31.1), D16S539 (16q24.1), D18S51 (18q21.33), D19S433 (19q12), D21S11 (21q21.1). (Butler, 2006).

Figure 2. The structure of Short Tandem Repeat (STR) (Butler, 2006).

DNA marker (Butler, 2005). Thus, for the DNA marker D3S1358: D = DNA, 3 = Chromosome 3, S = single copy sequence, 1358 = 1358th locus described on chromosome 3.
Table 1. Comparison of STR loci present in kits used in the United States.

<table>
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<th>Power Plex®21</th>
<th>Power Plex®Fusion</th>
<th>Identifier</th>
<th>Mini Filer</th>
<th>Extended*</th>
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*see Ref. (Lareu et al., 1996) for more information on the proposed U.S. extended core.

**COMMON COMMERCIAL KITS USED IN THE UNITED STATE**

To further assist efforts in forensic DNA testing and human identification, Promega and Applied Biosystems have released a number of new STR typing kits Table 1. The European Standard Set of STR loci added to U.S. core loci used for the Combined DNA Index System are among the most recent STR kits were developed. Some of these kits are capable of simultaneous, multicolor fluorescence detection of 15 STRs and the sex-typing marker amelogenin in a single PCR reaction (Butler et al., 2006).

**AUTOSOMAL STRs LOCUS INFORMATION**

Autosomal STRs locus information were evaluated and selected at The Cooperative Human Linkage Center. http://www.chlc.org evaluates the genetic markers and the loci are selected from there. Additional STR loci, chromosomal location and repeat sequence for each core STR locus are provided in Table 1 and Table 2 (Ruitberg et al., 2001; Klintschar et al., 2005; Klintschar et al., 2006; Imad et al., 2014a). According to the International Society of Forensic Genetics (ISFG) recommendation the repeat sequence motif be defined so that the first 5'-nucleotides on the Gene Bank forward strand define the repeat motif used. So the repeat motif for each STR marker is listed based on this. A significant fact is that STR allele sizes are measured relative to an internal size standard during electrophoresis. This depends on the DNA strand that is labeled using a dye that may have a different apparent measured size.

**Y-CHROMOSOME SHORT TANDEM REPEAT (Y-STRs)**

Chromosome Y microsatellites or STR's seem to be an ideal markers to delineate differences between human populations for several reasons: (i) they are transmitted in uniparental (paternal) fashion without recombination (De Knijff et al., 1997), (ii) they are very sensitive for genetic
Table 2. Information on 21 autosomal STR loci present in The PowerPlex® 21 System kits

<table>
<thead>
<tr>
<th>STR Locus*</th>
<th>Label</th>
<th>Physical position</th>
<th>Chromosomal location¹</th>
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The 13 CODIS core loci are highlighted in bold font. Adapted from Cotton et al., 2000; Wiegand et al., 1993; physical positions are from Schneider et al., 1998.

Drift, and (iii) they allow a simple highly informative haplotype construction (Kayser et al., 1997). Also, for forensic applications, this ability to differentiate distinct Y chromosomes makes Y-STR’s an advantageous addition to the well characterized autosomal STR’s. For a number of forensic applications Y-STR’s could be superior to autosomal STR’s. Especially in rape cases where (i) the differential extraction was unsuccessful, (ii) the number of sperm cells is very low, (iii) due to vasectomy epithelial cells instead of sperm cells from the ejaculate of the perpetrator have to be analyzed, and (iv) the perpetrator, due to a familial relationship shares many autosomal bands with the victim, Y-STR’s could provide crucial evidence. Also, in the case of male-male rape or rape cases with multiple perpetrators Y-STR’s could lead to essential qualitative evidence. In all such cases Y-STR’s facilitates a simple and reliable exclusion of suspects (Redd et al., 1997).

**COMMON COMMERCIAL KITS USED IN THE UNITED STATES**

Two most popular commercial kits in the U.S., are namely PowerPlex®Y System (a,b) and AmpFISTR®Yfiler™ PCR Amplification Kit. There are currently four online searchable Y-STR haplotype databases based in the United States and intended for forensic use. Three are maintained by commercial vendors: Reliagene, Inc., Promega Corporation and Applied Biosystems, Inc. The fourth is maintained by the University of Arizona. The National Center for Forensic Science (NCFS) also maintains a Y-STR database that will soon be available online. However, these databases are somewhat limited in the number of individuals and loci profiled, which sometimes limits their operational usefulness.

**Y CHROMOSOME IS A USEFUL TOOL FOR TRACING HUMAN EVOLUTION**

The Y chromosome is becoming a useful tool for tracing human evolution through male lineages (Jobling, 1994) as well as application in a variety of forensic situations (Kayser et al., 1997) including those involving evidence from sexual assault cases containing a mixture of male and female DNA (Prinz et al., 1997; Prinz et al., 2001). Y-chromosome specific methods can improve the chances of detecting low levels of male DNA in a high background of female DNA.

The European Y chromosome typing community has
established a "minimal haplotype" and an "extended haplotype" for inclusion of common loci into a central DNA database. These haplotypes consist of results from the following Y STR markers: DYS19, DYS389I, DYS389II, DYS390, DYS391, DYS392, DYS393, DYS385, and YCAII (Roewer et al., 2001). The combination of these Y STRs can distinguish approximately 74 to 90% of male individuals in various local European populations (Kayser et al., 1997). The minimal haplotype, the results of which have been accepted for court use in Europe (Pascali et al., 1999), is most commonly used and is typically obtained via two or three separate multiplex amplifications. In this study, a Y STR 20plex capable of the simultaneous amplification of 20 polymorphic Y chromosome-specific PCR products is introduced. This 20plex includes all of the markers in the European extended haplotype (Roewer et al., 2001) and also contains the trinucleotide loci DYS388 (M. Kayser et al., 1997) and DYS426 (Jobling et al., 1996), the tetranucleotide loci DYS437 (Ayub et al., 2000; Grignani et al., 2000), DYS439 (Ayub et al., 2000; Grignani et al., 2000), GATA A7.1 (DYS460) and H4 (White et al., 1999; Gonzalez-Neira et al., 2001), the pentanucleotide loci DYS438 (Ayub et al., 2000; Grignani et al., 2000) and DYS447 (Redd et al., 2002), and the hexanucleotide marker DYS448.

Analysis of Y-chromosome microsatellite haplotypes in globally human population

Variability at microsatellite or short tandem repeat (STR) loci is being used, in various species, for linkage analysis, individual identification (Hammond et al., 1994; Redd et al., 1997), and population-genetic analyses. Autosomal STR loci have also been successfully applied to reconstruct human evolutionary history (Goldstein et al., 1995). The resulting phylogenetic trees reveal evolutionary relationships similar to those based on mtDNA sequence variation, and it has been a long lasting wish to add to these trees the one strictly based on Y chromosome-specific markers. Approximately 5 years ago, the number of published STR loci on the human Y chromosome was <15, and only one Y chromosome-specific minisatellite was known (Roewer et al., 1992). The number of known Y chromosomal single-nucleotide polymorphisms (SNPs) was <10 (Nakahori et al., 1989; Jobling, 1994; Mathias et al., 1994; Seielstad et al., 1994; Hammer, 1995; Whitfield et al., 1995), and only one Alu-insertion polymorphisms had been discovered. This picture, however, has totally changed, owing to the recent introduction of many Y SNPs (Underhill et al., 2000) and STRs (White et al., 1999), and additional new markers are to be expected.

Recently, Y-STR variability has been used both for the dating of SNP mutations, in order to draw conclusions about the origins and history of human populations (Underhill et al., 1996; Zerjal et al., 1997), and for human identification in forensic casework (Kayser et al., 1997; Prinz et al., 1997). Nevertheless, global studies of Y-STR variability are still rare (Seielstad et al., 1999), and most of these analyses have been based on combined single-locus information. The major advantage of analyzing the nonrecombining part of the Y chromosome is that single-locus information can be used to construct compound haplotypes that allow male lineages to be characterized in a much more detailed fashion. It has been questioned whether the tracing of human migration history can be achieved solely on the basis of Y STRs (De Knijff et al., 1997).

Y-STR database

When the DNA profile of a known suspect or victim matches the DNA profile from crime scene evidence, the individual is "included" as a potential source of that evidence. In the U.S., the strength of the match is most often expressed as a statistic that describes the estimated frequency of occurrence of the DNA profile in unrelated individuals within various population groups. Due to the lack of recombination along most of the length of the Y chromosome, Y-STR loci are not statistically independent of one another (unlike standard autosomal STR markers) and are co-inherited as extended haplotypes of linked markers. Therefore, multiplication of single-locus allele frequencies to obtain estimated Y-STR haplotype frequencies is not appropriate. An estimation of the frequency of occurrence of a particular Y-STR haplotype necessitates the use of a counting method, which, with the limited sizes of databases available, produces an estimate that depends entirely upon the size of the database used. Thus, large databases of multi-locus Y-STR haplotypes need to be generated to maximize the probity of Y-STR evidence. A large comprehensive European-based Y-STR database is maintained by the Institute of Legal Medicine, Charité-University Medicine Berlin (www.yhrd.org).

However, although a subset of this database comprises the SWGDAM core loci, it is less useful for frequency estimates from haplotypes that have been generated using the two most popular commercial kits in the U.S., namely the PowerPlex® Y System(a,b) and AmpFiSTR®Yfiler™ PCR Amplification Kit. There are presently four online searchable Y-STR haplotype databases based in the United States and intended for forensic use. Three are maintained by commercial vendors: Reliagene, Inc., Promega Corporation and Applied Biosystems, Inc. The fourth is maintained by the University of Arizona. The National Center for Forensic Science (NCFS) also maintains a Y-STR database that will soon be available online. These databases differ in the number of Y-STR markers and individuals, although all possess the SWGDAM core loci. However, these
Table 3. Current U.S.-Based Y-STR databases.

<table>
<thead>
<tr>
<th>Agency</th>
<th>URL</th>
<th>Number of markers</th>
<th>Number of samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>National Center for Forensic Science</td>
<td>To be determined</td>
<td>76</td>
<td>1.396</td>
</tr>
<tr>
<td>University of Arizona</td>
<td><a href="http://Amadeus.biosci.arizona.edu/kcaldero/str">http://Amadeus.biosci.arizona.edu/kcaldero/str</a></td>
<td>83</td>
<td>2.518</td>
</tr>
<tr>
<td>Applied Biosystems</td>
<td><a href="http://www.Appliedbiosystems.com/yfilerdatabase">www.Appliedbiosystems.com/yfilerdatabase</a></td>
<td>17</td>
<td>3.561</td>
</tr>
<tr>
<td>Promega Corporation</td>
<td><a href="http://www.promega.com/techserv/tools/pplexy/">www.promega.com/techserv/tools/pplexy/</a></td>
<td>12</td>
<td>4.004</td>
</tr>
<tr>
<td>Potential Size of National Y-STR Database</td>
<td></td>
<td>16.102</td>
<td></td>
</tr>
</tbody>
</table>

databases are somewhat limited in the number of individuals and loci profiled, which sometimes limits their operational usefulness. For example, the biggest U.S.-based database comprises haplotypes from 4,623 individuals. By combining data from these U.S. databases, a much larger Y-STR database of approximately 16,000 individuals can be created, resulting in a significant increase in the probative value of Y-STR evidence. Also, merging the NCFS and University of Arizona databases will increase the number of samples with extended Y-STR loci haplotypes, which may be of assistance to those interested in developing the next generation of Y-STR multiplex systems.

Establishing a national database that incorporates data from a multitude of sources requires the implementation of a number of quality indicator metrics. Quality assurance procedures must be developed to govern the suitability and quality of data from diverse sources. For example, it may be necessary for donors of data to establish analytical prowess by testing externally provided proficiency samples. Since each commercial kit or academic multiplex system uses different primer sets, it will also be essential to ensure that allele calls are equivalent regardless of the multiplex system employed.

Importantly, merged data must be purged of duplicate samples that have been submitted by the same donor to multiple databases. To effectively manage the data, a Y-STR Database Consortium comprised of database stakeholders from commercial companies, academia, the FBI and U.S. crime laboratories was formed at the February 2006 AAFS meeting in Seattle (Table 3). It was agreed that NCFS, a program of the National Institute of Justice (NIJ) hosted by the University of Central Florida, would maintain and manage the consolidated Y-STR database on behalf of stakeholders. The National Institute of Justice is funding this effort. As a group, we are working to collate existing Y-STR data from various commercial and academic sources and have enlisted the aid of geographically diverse crime laboratories to furnish additional samples.

MITOCHONDRIA

Structure of mitochondria

The typical human cell has several hundred mitochondria, cytoplasmic organelles that convert energy to forms that can be used to drive cellular reactions. Without them cells would be dependent on anaerobic glycolysis for all their adenosine triphosphate (ATP). The mitochondria have a characteristic double membrane structure, in which the outer membrane contains large channel-forming proteins (called porin) and is permeable to all molecules of 5000 daltons or less, while the inner membrane is impermeable to most small ions and is intricately folded, forming structures called cristae. The large surface area of the inner mitochondrial membrane accommodates respiratory chain and ATP synthase enzymes involved in the process of oxidative phosphorylation (OXPHOS). The mitochondrial matrix contains hundreds of enzymes, including those required for the oxidation of pyruvate and fatty acids and those active in the tricarboxylic acid (TCA) cycle. The matrix also contains several identical copies of the mitochondrial DNA, mitochondrial ribosomes, tRNAs and various enzymes required for the transcription and translation of mitochondrial genes (Alberts et al., 1994).

Organization of the human mitochondrial genome

The human mitochondrial genome is 16,569 base pairs (bp) in length (Anderson et al., 1981), a closed, circular molecule located within the mitochondrial matrix and present in thousands of copies per cell. Mitochondrial DNA has two strands, a guanine rich heavy (H) strand and a cytosine-rich light (L) strand. The heavy strand contains 12 of the 13 polypeptide-encoding genes, 14 of the 22 tRNA-encoding genes and both rRNA-encoding genes. Introns are absent in mtDNA, and all of the coding sequences are contiguous (Anderson et al., 1981; Wallace et al., 1992; Zeviani et al., 1996).

The only non-coding segment of mtDNA is the displacement loop (D-loop), a region of 1121 bp that contains the origin of replication of the H-strand (OH) and the promoters for L and H-strand transcription. The mtDNA is replicated from two origins. DNA replication is initiated at OH using an RNA primer generated from the L-strand transcript. H-strand synthesis proceeds two-thirds of the way around the mtDNA, displacing the parental H strand until it reaches the L-strand origin (OL), situated in a cluster of five tRNA genes. Once exposed on the displaced H-strand, OL folds a stem-loop structure.
Mitochondrial DNA replication

Mitochondrial DNA is replicated by the DNA polymerase gamma complex which is composed of a 140 kDa catalytic DNA polymerase encoded by the POLG gene and a 55 kDa accessory subunit encoded by the POLG2 gene (Van der Tovar, 2005). Replication of mitochondrial DNA begins on only one strand within the non-coding “control” region. As this strand is replicated, the opposite strand of the original DNA duplex is displaced and forms a single-stranded loop hence the name “D-loop” for Displacement loop. The displaced strand is then broken down and the replication process begins again displacing a single strand in the process (Schon et al., 2012).

Reasons for using mitochondrial DNA rather than nuclear DNA

First, multiple copies: Each mitochondrion contains its own DNA, with many copies of the circular mitochondrial DNA in every cell. It is thought that each mitochondrion contains between 1 and 15, with an average of 4 to 5, copies of the DNA (Reynolds et al., 2000) and there are hundreds, sometimes thousands, of mitochondria per cell. The result is that there are many thousands of copies of the mitochondrial DNA in every cell. This compares with only two copies of nuclear DNA. Second, Better protection: The mitochondrion also has a strong protein coat that protects the mitochondrial DNA from degradation by bacterial enzymes. This compares to the nuclear envelope that is relatively weak and liable to degradation.

Third, higher rate of evolution: DNA alterations (mutations) occur in a number of ways. One of the most common ways by which mutations occur is during DNA replication. An incorrect DNA base may be added; for example, a C is added instead of a G. This creates a single base change, or polymorphism, resulting in a new form. These single base mutations are rare, but occur once in every 1,200 bases in the human genome. The result is that the rate of change, or evolutionary rate, of mitochondrial DNA is about five times greater than nuclear DNA (Bar et al., 2000). This is important in species testing, as even species thought to be closely related may in time accumulate differences in the mitochondrial DNA but show little difference in the nuclear DNA. Finally, maternal inheritance: A further reason for the use of mitochondrial DNA in species testing, and in forensic science, is its mode of inheritance. Mitochondria exist within the cytoplasm of cells, including the egg cells. Spermatzoa do not normally pass on mitochondria and only pass on their nuclear DNA. The resulting embryo inherits all its mitochondria from its mother (Tully et al., 2004; Brown et al., 2002; Brown et al., 2000).

The result is that mothers pass on their mitochondrial DNA type to all their offspring, but only the daughters will pass on the mitochondrial DNA to the next generation. Mitochondrial DNA is therefore passed from generation to generation down the maternal line. Mechanisms for this include simple dilution (an egg contains 100,000 to 1,000,000 mtDNA molecules, whereas a sperm contains only 100 to 1000), degradation of sperm mtDNA in the fertilized egg, and, at least in a few organisms, failure of sperm mtDNA to enter the egg. Whatever the mechanism, this single parent (uniparental) pattern of mtDNA inheritance is found in most animals, most plants and in fungi as well. Also, most mitochondria are present at the base of the sperm's tail, which is used for propelling the sperm cells. Sometimes the tail is lost during fertilization. Also, unlike nuclear DNA, where there is a shuffling of the chromosomes at every generation, the mitochondrial DNA does not recombine with any other DNA type and remains intact from generation to generation (Brown et al., 2002; Brown et al., 2000; Pastores et al., 1994; Guntheroth et al., 1990). The role of DNA is to encode protein and RNA molecules, and the mitochondrial DNA is no different. All mammalian mitochondrial DNA is very similar, with the order and position of the genes being the same. The general structure of the mitochondrial DNA is shown in Figure 3. As with nuclear DNA, to indicate the significance of the match, analysts usually estimate the frequency of the sequence in some population. The estimation procedure is actually much simpler with mtDNA. It is not necessary to combine any allele frequencies because the entire mtDNA sequence, whatever its internal structure may be, is inherited as a single unit (a “haplotype”). In other words, the sequence itself is like a single allele, and one can simply see how often it occurs in a sample of unrelated people (Lutz et al., 2003; Alvarez et al., 2007).

This polymorphism allows scientists to compare mtDNA from crime scenes to mtDNA from given individuals to ascertain whether the tested individuals are within the maternal line (or another coincidentally matching maternal line) of people who could have been the source of the trace evidence (Coble et al., 2006).
Mitochondrial coding region as a source for variability

Forensic mtDNA analysis commonly investigates the HVI and HVII regions, but analysis of the coding region is gaining interest as this region harbours highly polymorphic mutations that may increase the discrimination of most common mtDNA types (Parsons and Coble et al., 2002).
2001; Andreasson et al., 2002; Imad et al., 2014b). The development of high throughput capillary sequencing has made whole mitochondrial genome analysis feasible and large collections of coding region data are now progressing (Herrnstadt et al., 2002). Thus, selection of new markers suitable for forensic coding mtDNA analysis is feasible. In a study by Coble et al. (2004) a panel of coding region SNPs were selected to resolve the most common HVI and HVII sequences including the H haplogroup. An SNP assay was chosen as whole mitochondrial genome sequencing is not a practical approach for the analysis of forensic samples. Moreover, coding SNPs were also chosen to resolve four additional commonly observed HVI and HVII haplotypes as the H-defining SNPs may not strained by only choosing silent and non-private polymorphism (variant observed) in more than one individual for ethical and practical reasons, nearly a four-fold increase in discrimination of the five most common haplogroups (H, J, T, V and U) was obtained (Coble et al., 2004).

**INTERPRETATION OF SEQUENCE DATA**

Current Sanger capillary sequencing systems are like the widely used Applied Biosystems (Figure 4), the dNTP/ddNTP mixture causes random, non-reversible termination of the extension reaction, creating from the different copies molecules extended to different lengths. Following denaturation and cleaning up of free nucleotides, primers, and the enzyme, the resulting molecules are sorted by their molecular weight (corresponding to the point of termination) and the label attached to the terminating ddNTPs is read out sequentially in the order created by the sorting step. Additionally, the advent of more sensitive detection systems and several rounds
Figure 5. Shows the sketch of an electropherogram for two D16S539 alleles. One allele has eight repeats of the sequence GATA and the other has five. A small rectangle represents a GATA repeat. For illustration here only one copy of each allele (with a fluorescent molecule, or “tag” attached) is shown. However, PCR generates many more copies from the DNA sample with these alleles at the D16S539 locus. Whilst these copies are drawn through the capillary tube, the tags glow as the STR fragments move pass the laser beam. The colored light from the tags is measured using an electronic camera. A computer is finally used to produce the electropherogram based on the signals received.

of primer extensions (equivalent to a linear amplification) permit smaller amounts of starting DNA to be used for modern sequencing reactions. Using current Sanger sequencing technology, it is technically possible for up to 384 sequences (Emrich et al., 2002) of between 600 and 1,000 nt in length to be sequenced in parallel.

Base designations:

“A” designation—green peaks
“G” designation—black peaks
“T” designation—red peaks
“C” designation—blue peaks
“N” designation—peaks that, for whatever reason, are not clear enough to designate as A, G, T, or C. These bases are generally inconclusive. Often, a position that has an inconclusive (N) base in one direction may appear very clear in the other direction.

TYPES OF POLYMORPHISMS

Purines to pyrimidines - Transversions: (purines to pyrimidines or pyrimidines to purines).

Insertions: an extra base is present when compared to the Anderson reference sequence.
Deletions: a base is missing when compared to the Anderson reference sequence.

SEQUENCER™ 4.7

The validate mitotypes command allows ours to compare the results of separate analysts. Select two contigs which have been assembled to the same reference sequence (SEQUENCER™ 4.7 User Manual for Windows © 1991 - 2007 Gene Codes Corporation, Inc. All rights reserved).

After the raw sequencing data has been collected by the 310 genetic analyzer and analyzed by the Sequencing Analysis program, it must be reviewed manually and interpreted with the aid of the Sequencher™ software (Figure 5). Sometimes the data will present characteristics that make interpretation more complex, though not impossible. Interpretive skills improve with experience, but listed below are suggestions for those less familiar with mtDNA analysis on dealing with some of the more common features that affect interpretation. However, all data (especially indications of mixtures, heteroplasmy, etc.) should be reviewed on a case-by-case basis, and interpretations made
The forensic community generally recognizes the HV1 region as consisting of base positions 16024-16365 and HV2 as base positions 73-340, as numbered according to the Cambridge Reference Sequence (also referred to as the Anderson Sequence) (Bainbridge et al., 2011). This laboratory will attempt to sequence all the bases within these defined regions, as well as an additional 25 base pairs in either direction (that is, 15999-16390 and 48-365). While it is preferable to have confirmation of each sequence by comparing the forward and reverse strands, it is sometimes necessary to use two forward strands or two reverse strands as confirmation of the sequence (Al-Haggar et al., 2013; Pareek et al., 2011). In these cases it is suggested that the sample be cycle sequenced a second time to obtain the confirmatory strand in the same direction. In addition to case samples, a positive and negative control must be sequenced. The purpose of controls is to show that each stage of the analysis is working properly. Therefore, only one positive and negative control is necessary for each step of the procedure (for example, if samples from three amplification runs are combined into one cycle sequencing run, only one positive and one negative must be carried through to show the sequencing reagents and thermal cycler have performed as expected). If the results obtained for the controls are not as anticipated, the evaluation of the results will be determined on a case-by-case basis (Goldberg et al., 2006).

Conflict of Interests

The author(s) have not declared any conflict of interests.

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