Dental DNA Fingerprinting

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Abstract

DNA fingerprinting has revolutionized the concept of identification. Although clinical observation of available medical and dental patient records remain the gold standard for forensic pathology, dental DNA fingerprinting is booming in forensic science. The evolution of DNA fingerprinting techniques is discussed in this mini-review with a special emphasis to forensic odontology.

Keywords: DNA Fingerprinting; Molecular Level; Consanguinity; Paternity; Forensic Odontology; Polymerase Chain Reaction; Radiological Identification; High Molecular Weight; Single Locus Probing; Multiple-Locus Probing

Introduction

Molecular biology developed when it was realized that DNA lies behind all the cell’s activities. The development of methods and techniques to study processes at the molecular level has led to new and powerful ways of isolating, manipulating, and exploiting nucleic acids. DNA fingerprinting is the result of such an endeavor. This technique is mostly known by its application in forensic medicine, but is also used in transplant medicine, in the search for hereditary disorders, consanguinity, paternity, and in anthropology [1,2]. The role of dental restorations, prosthesis, and radiological identification as the main strain of forensic odontology has declined lately, whereas molecular biology and laboratory procedures are rapidly increasing in efficiency and availability [3]. The tooth is the most valuable source to extract DNA since it is a sealed box preserving DNA from extreme environmental conditions, except its apical entrance. Teeth have been the subject of DNA studies as the dental hard tissue physically encloses the pulp and offers an anatomical configuration of great durability [4]. Moreover, even a single tooth provides valuable information regarding the individual to whom the tooth belongs [5-7]. In this mini-review dental DNA fingerprinting and its applications will be discussed. The applications in health and medicine and in anthropology are beyond the scope of this mini-review.

History

Jeffery described in 1985 hypervariable regions of human DNA using multilocus probes and the applicability of these DNA polymorphisms to the individualization of human blood and tissues [8]. The potential forensic application was immediately recognized [9]. Polymerase chain reaction which was originally described by Saiki et al. and subsequently automated by Mullis and Fakona, has emerged as a powerful tool in forensics for the exponential in vitro amplification of specific sequences of interest from minute quantities of DNA or RNA and was applied to forensic odontology [10,11]. Schwartz isolated high molecular weight (HMW) DNA from the teeth under different environmental conditions such as varying pH, humidity, temperature, storage, etc. It was determined that the environmental conditions did not affect the ability to obtain HMW human DNA from dental pulp [12]. Pötsch performed genomic dot blot hybridization for sex determination using the botinylated repetitive DNA probe pHY2.1 and sex was correctly classified in all cases using 50-100 ng target DNA from dental pulp [3]. Sweet and Sweet, a case of human remains identification by a preserved unerupted third molar which enabled 1,35ug DNA from the pulp [13,14].

How does it work?

a) Making DNA fingerprints is a laboratory procedure that requires 6 steps [1]. Isolation of DNA; DNA must be removed from the cells or tissues of the body. Only a small amount of tissue e.g. dental pulp is needed.

b) Cutting, sizing, and sorting. Special enzymes called restriction enzymes are used to cut the DNA at specific places. For example, an enzyme called EcoR1, found in E coli bacteria, will cut DNA only when the sequence GAATTC occurs...
(guanine, adenine, adenine, thymine, thymine, cytosine) The DNA pieces are sorted according to size by a sieving technique called electrophoresis. The DNA pieces are passed through a gel made from seaweed agarose (Southern blot). This technique is the biotechnology equivalent of screening sand through progressively finer mesh screens to determine particle sizes.

c) Transfer of DNA to nylon; The distribution of DNA pieces is transferred to a nylon sheet by placing the sheet on the gel and soaking them overnight.

d) Probing; Adding radioactive or colored probes to the nylon sheet produces a pattern called the DNA fingerprint. Each probe typically sticks in only one or two specific places on the nylon sheet.

e) DNA fingerprint, The final DNA fingerprint is built by using several probes (5-10 or more) simultaneously. It resembles the bar codes used by grocery scanners; The final product of a DNA fingerprint is an autoradiograph that contains at least five essential lanes. The markers are standardized DNA fragments of known size, which have been radioactively labeled. They help determine the size of the various fragments. The “control” is DNA from a source known to react positively and reliably with the DNA probes and shows whether the test has worked as expected. In crime cases the experimental lanes have samples from the victim, the defendant, and the crime scene.

Two variants of DNA fingerprinting have been used: single locus probing (SLP) and multiple-locus probing (MLP). In SLP, each probe is specific for a single site, that is a single locus in the genome. Because humans are diploid, each SLP probe normally gives rise to two bands from each person, provided that the chosen locus shows substantial allelic variation. Occasionally persons will be homozygous and show only a single band. For full identification using SLPs it is necessary to run several reactions, each using a different SLP probe. Multilocus probes bind to multiple sites in the genome and consequently generate multiple bands from each individual. Because it is not known which particular band comes from which particular locus, interpretation is difficult. Furthermore statistical analysis is impractical, and data cannot be reliably standardized for reference. In practice, fingerprints generated by MLP must be directly compared with others run on the same gel.

Historically MLP was used before SLP. However, SLPs uses smaller amounts of material and is easier to interpret and compare. Statistical analysis and population frequencies are possible using SLP data. Consequently, MLP methods have largely been displaced by SLP analysis [15]. In addition to preparing DNA for fingerprinting, PCR may be used to generate DNA segments for direct comparison by DNA sequencing or hybridization. PCR is especially useful for amplifying small regions of DNA with high -person to person variability. If the sequences of two samples match in several highly variable regions, they are probably from the same person. [15]. In this approach DNA from a forensic sample is amplified by PCR and compared with DNA from the suspect. For hybridization, spots of DNA samples are bound to a membrane and tested for binding a DNA probe that is tagged with a fluorescent dye (radioactive probes were used in early work). In such dot blots, the probe either binds or doesn’t bind, so any spot is either positive or negative. Alternatively, segments of DNA that have been amplified can be fully sequenced. DNA fingerprinting relies on the unique pattern found in different individuals when a series of DNA fragments is separated according to length. The polymerase chain reaction (PCR) is now used to generate the DNA fragments for DNA fingerprinting [15].

Teeth are important evidentiary material in forensic cases since they are more resistant to postmortem degradation as well as extremes of environmental conditions. Teeth are also easy to transport and serve as a good source of DNA. Comparisons of antemortem dental records with skeletal remains provide useful means to identify individuals; even in a mass grave. In affluent societies, dental records may be decisive in determining the identity of individual victims. However, in less affluent communities, and these are more likely to be involved in human rights abuses leading to mass murder, dental records are unlikely to be available. In this situation the only option for identification is DNA analysis [16]. It is possible to discriminate one individual from all others with a high level of confidence by starting with only 1 ng or less of target DNA, whereas the amount of DNA that can be recovered from molar teeth with pulp volumes of 0.023-0.031 cc is nearly 15-20 mg. In a study by Pötsch et al. the total production of genomic DNA obtained from a dental sample ranged from 6-50 µg DNA. The results were obtained from DNA extracted from the dental pulp and did not show any difference when compared to the patterns obtained from DNA extracted from blood samples or available lung tissues [3].

Remualdo evaluated the PCR amplification of DNA retrieved from teeth subjected to heat (200 degrees Celsius, 400 and 600 degrees C) during 60 minutes, testing three different extraction methods (organic, ammonia acetate/isopropanol and silica). Using the organic method for genomic DNA extraction 50% of samples subjected to burning were amplified, but only at lower temperatures (200 degrees C and 400 degrees C). At higher temperatures (500 and 600 degrees C) the isopropanol/ammonia acetate extraction yielded better results, mainly for extraction of mitochondrial DNA (mtDNA) [17]. Mitochondrial DNA can be sourced from dentine powder and also via dentine in the case of root-filled tooth [18].

**Methods of Dental DNA Fingerprinting**

**RFLP**

Restriction fragment length polymorphism (RFLP) was the first DNA profiling technique inexpensive enough to see widespread application. RFLP analysis was an important test in genome mapping, localization of genes for genetic disorders, determination of risk of disease and paternity testing. Due to the rise of inexpensive DNA sequencing technologies RFLP is now becoming largely obsolete, except in developing countries. As mentioned...
above,special enzymes,so called restrictive endonucleases deaves
the DNA like scissors at specific sites,each recognizing a particular
sequence.These DNA scissors are specifically chosen to cut DNA at
sites which are not found within the sequence of “tandem repeats”
rather than conserved less variable regions. The cut fragments will
contain variable number of tandem repeats (VNTRs) of varying
lengths,there by producing DNA fragments of various sizes. The
VNTR testing,which may present short repeated sequences of
intermediate size,is rarely used in forensic analysis due to the poor
quality DNA provided with this method.

PCR methods

PCR is used to amplify the amount of DNA material available,as
mentioned before [15]. To carry out the reaction special enzymes
and DNA primers are required.These primers are like known
constant sections of DNA but not labeled.They are designed to
know constant sections of DNA at the ends of variable regions to
be amplified. The principle of PCR is that the DNA is capable of
duplicating itself. This is done by unwinding the strands of DNA
and each strand acts as a template for synthesis of a new strand. By
PCR technique one can amplify specific DNA segments dependent
on the primer deployed. The standard PCR reaction runs through
30 cycles in a couple of hours which result in amplification of the
original by over 10^9 times [19]. The DNA found can be genomic
(found in the nucleus) and mtDNA(in the mitochondria). The
teeth are an excellent source of genomic and mtDNA because PCR
analyses allow comparing the collected postmortem samples to
known antemortem samples or parental DNA [18]. Main advantage
of mtDNA is the high number of copies per cell (from thousands
to thousands of organelles).

STR Analysis

Short Tandem Repeat (STR) analysis is one of the most useful
methods in molecular biology, which is used to compare specific
loci of DNA from two or more samples. A short tandem repeat is
a microsatellite consisting of a unit of two to thirteen molecules
repeated hundreds of times in a row on the DNA strand [20]. It is a
frequent and routinely used marker in forensics. STRs have a high
power of individual discernment because of their high standards
of polymorphic informative content. The nonoverlapping size of
the alleles from different contributors serves to distinguish them.
Currently they are detected by fluorescent detection methods. STR
serves as the standard for the combined DNA index system ( CODIS).
The FBI (Federal Bureau of Investigation) has chosen 13 definite
STR loci which are together referred to as CODIS markers and the
sex amelogenin marker. It is said that the likelihood that any two
individuals would have identical patterns with this system is 1
in 250.000.000.000.000 [21]. STR was used on 45 DNA samples
from teeth obtained from unidentified bodies buried in 1995
and exhumed in 2000,and dental pulp showed strongest PCR
amplification signals [22].

Mitochondria DNA analysis

When the sample lacks nucleus DNA is extracted from mitochondria. Silva and Passos in 2002 stated that mtDNA analysis
is used for ancient tissues like bone,hair and teeth, where
analysis of nuclear DNA cannot be done [23]. High molecular
weight mtDNA are obtained from teeth, especially in degraded
remains [24]. Every child has the same mtDNA as its mother because
mitochondrion of the embryo is from the mother’s egg while genomic
DNA is from father’s sperm. It is thus a valuable test in identifying
missing persons by comparing mtDNA of unidentified remains with
that of a possible maternal relative [25]. The technique is expensive
as it is performed by direct sequencing of nitrogenous bases and
provides only limited information as it is primarily matrilineal.

Analysis of Y- chromosome

It involves targeting of the polymorphic region of the Y
chromosome (Y-STR) using primers. As the Y chromosome is passed
to the son from his father, analysis of markers on the chromosome
helps in sketching relationships among males [26].

SNP: SNP, or single nucleotide polymorphism is a detection
technique to scan for new polymorphisms and to determine the
allele(s) of a known polymorphism in target sequences [27]. SNP
techniques provide valuable information on descent, sex, evolution
and is highly automated [28, 29]. The Indian Ocean tsunami of 26
december 2004 created major challenges for forensic identification
of dead bodies. DNA profiling were useful in identification of those
bodies where other dental procedures have been unsuccessful [30].

Recent technologies in genetic identification

Microarray techniques: The nucleic acids of the target
are hybridized to high-density microarrays containing several
thousand oligonucleotides immobilized on chips or beads [29].
Many commercial platforms are available for SNP analysis. E.g.,
Illumina and Affymetrix. DNA analysis using this technology is used
in forensic testing for sequencing and resequencing, paternity
testing, SNP genotyping and identification of the individual [31].

Next generation genome sequencing: Next generation
genome sequencing permits analysis of several hundred loci or even
the entire genome by producing enormously parallel sequencing
[32-35]. Amplification or cloning of the sequenced DNA fragments
is automated and provided with a reading process. Next generation
genome sequencing platforms available are Illumina genomic
analyzer, Roche 454 genome sequencer and ABI Sequencing by
Oligonucleotide Ligation and Detection NGS permits analysis of
copy number variants (CNVs) and other structural rearrangements.
Next generation sequencing can be used for both genome and
transcription analysis. In genome analysis it permits the high-quality
variant calling for SNPs, insertions and deletions, and allows the
analysis of CNVs and other structural rearrangements.

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Conclusion

DNA fingerprinting has revolutionized the concept of identification. Meanwhile, clinical observation of available medical and dental patient records remains the gold standard for forensic pathology. DNA analysis is now the greatest forensic tool used in forensics. Of the three main kinds of DNA fingerprints, RFLP, VNTR and STR, the most commonly used is STR. RFLP and VNTR require a lot of DNA, which is usually very difficult to find at the scene and often the DNA fragments being analyzed are too large to amplify via a PCR. STR, on the other hand, uses shorter sections of DNA, which are often the DNA fragments being analyzed are too long to amplify via a PCR. Additionally, STR analysis does not require the hybridization to a DNA probe. New techniques as the microarray techniques [36] and next generation genome sequencing [37] will stimulate dental forensics and identification further.

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