Forensic Application of DNA Methylation in Age-Prediction

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Abstract
Advances in Epigenetics suggested that DNA methylation profiling, plays an important role in various biological process such as silencing, maintaining or expression of genomic stability. DNA methylation is the most comprehensively studied epigenetic mark, and it involves the addition of a methyl group to the 5' cytosine of CG dinucleotide, referred to as CpGs. DNA methylation is related to gene regulation and cell differentiation by affecting transcription factor binding sites, insulator components, and chromosome morphology. Recent advances suggested that DNA methylation markers not only can provide information about tissue of origin of evidence sample, but also can rely information about the age and lifestyle or environmental exposure of an individual, for an unknown suspect. In this review, we will be discussing age-associated DNA methylation in various tissues, blood and semen and the applicability of the DNA methylation-based age prediction method to the forensic investigations.

Introduction
DNA methylation defined as the addition of methyl group to the 5’ cytosine of CG dinucleotide (CpGs). As CpG sites across wide range of aging research found to be highly associated with age, multiple age predictive models using DNA methylation at CpG site have been made. DNA methylation considered the most accurate method for age prediction [1]. Variable DNA methylation can be determined by cell differentiation, aging as well as environmental exposure. The established DNA methylation pattern during early years of development are maintained and remain relatively stable throughout life, decrease or increase in different loci [2]. Age-related methylation changes throughout the life of an individual concluded in two phenomena: the epigenetic clock and epigenetic drift. Epigenetic in general is defined as the modification of DNA and DNA packaging without any change in DNA sequence. Epigenetic clock is the age associated DNA methylation changes that are common across individuals of the same age. However, epigenetic drift is the accumulation of small changes or errors that is determined by the environment in which the person ages and it influences [3,4]. In epigenetic more specific age-associated DNA methylation changes at certain loci or genes were easier to be detected by the emergence of microarray and massive parallel sequencing technologies (MPS). Although age prediction using DNA methylation is not quite parallel to the chronological age [4]. DNA methylation measures at multiple CpG sites provide close prediction of a chronological age, with a narrow range that varies and specify the accuracy of each method introduced. Thereby, choosing the most appropriate analytical method for forensic application is more difficult than in clinical application. This review will improve the understandings about DNA methylation markers and their potential to be used as biomarkers in the forensic field, with a future plan to apply the previously identified markers on different tissues and test accuracy as well as identify more DNA methylation markers for various tissues, blood and semen.

Tissues
The age associated CpG sites have been found within a specific tissue and across tissues, because DNA methylation profile are diverse across different type of tissues. Earlier studies,
by Bocklandt [5] identified three age associated CpG sites in saliva at the promotor region of EDARADD, TOM1L1 and NPTX2, using the 27K Bead Chip array. The regression model resulted and built predicted the age of an individual with an average accuracy of 5.2 years. Following that Koch [6] using five data sets were able to identify 19 CpG hypermethylated aging sites of 20-30 sample each from epidermis, dermis, monocytes, T-lymphocyte and uterine cervical smear using Illumina 27K. Four of these in the GRIA2, TRIM58, KCNQ1DN and NPTX2 genes, and a hypomethylation site in the BRC4BP gene. Age prediction was implemented and the predictive model showed accuracy with mean absolute deviation (MAD) from chronological age of 12.7 years in eight validation sets. Breast tissue repeatedly showed a lower DNA methylation than other cell types at the CG site of the BIRC4BP gene, as it was also described by Horvath [7]. The prediction accuracy increased to 11.4 years of MAD when age prediction was focused on three CG sites in the NPTX2, GRIA2 and KCNQ1DN [8]. Followed his peers path using 450K Bead Chip array, was able to demonstrate a high association between the age and DNA methylation in three CpG sites of the genes ELOVL2, FHL2 and PENK in blood. A quantitative model was first built by Hannum [9], using 71 CpG sites of the 450K array and resulted in a very high prediction accuracy of age with an error of ~3.9 years in blood. Afterward Horvath [7] developed a highly accurate model, that can be implemented on multiple tissue for age prediction by allowing an accurate estimation of DNA methylation age of multiple cells and tissues. The analysis results of 7,840 non-cancer samples from 82 data sets produced using Illumina 27K or Illumina’s Human Methylation450 Bead Chip array (Illumina 450K), which encompass 51 different tissues and cell types. The 353CpGs that were automatically selected from the elastic net allowed highly accurate age prediction with an error of 3.6 years in the test data.

Blood based samples

Blood was also used in DNA methylation age-prediction. Pyrosequencing of blood at three CpGs at the genes ASPA, PDE4C and ITGA2B for age prediction was described by Weidner [10]. The model demonstrated high accuracy with a mean absolute deviation (MAD) from chronological age of 4.3 years. This was followed by Piekarska [11] who reported an age predictive model for blood using two CpGs in the ELOVL2 gene. Zbiec model had a prediction error of 6.85 years and accuracy of 5.03 years MAD from chronological age. Another model of five CpG sites of the genes TRIM59, C1orf132, FHL2, ELOVL2 and KLF14 was established and had an accuracy with a mean absolute deviation (MAD) from chronological age of 3.9 years [12]. Lately Park [13] reported an age predictive model for blood using three CpG sites of the genes CCDC102B, ZNF423, and ELOVL2 by evaluating of 760 blood samples based on a pyrosequencing platform. The model was highly accurate with a MAD from the chronological age of 3.4 years. Of all the genes tested, DNA methylation at FHL2 and KLF14 were significantly associated with age; yet, the primer design for pyrosequencing of these genes failed. As a consequence of the high association between the age and DNA methylation at the genes FHL2, KLF14, and ELOVL2, it has been repeatedly observed in many studies with blood, they are considered the most promising age-predictive markers for blood.

Semen

Lately, Lee [14] identified a new age associated CpG sites found in semen samples, using the 450K array and subsequent methylation Snapshot analyses. Semen is a particularly important body fluid in the forensic analyses. Although, Horvath model (we discussed earlier) is applicable for age-prediction in several types of cells and tissues, the age prediction values for sperm were much lower than the chronological age of the donors [7]. The model by Lee et al. showed a high correlation between the chronological and the predicted age and it consisted of three CpG sites (cg12837463, cg06979108 in the NOX4 gene, and cg06304190 in the TTC7B gene) a MAD from chronological age of ~5 years. Furthermore, the area around the TTC7B gene showed an alteration in DNA methylation in the sperm methylome of two samples collected 9-19 years apart from each individual [15]. Thus, indicating TTC7B as one of the most promising age predictive marker for semen. Also, Bekaert et al. [16] developed a model with a four age-associated markers suggested for blood PED4C, EDARADD, ELOVL2 and ASPA. The model proved to be highly accurate for age predictions using teeth samples with a MAD from chronological age of 4.9 years. Giuliani [17] also proved that the previously reported age-associated markers for blood, i.e., CpG sites located in the PENK, ELOVL2 and FHL2 genes [8], could be a great tool to predict the age in teeth; although, the mean absolute deviation (MAD) from chronological age, depending on the part of the tooth from which the sample taken, varied between 1.2-7.1 years.

Conclusion

DNA methylation and age-prediction models can be applied across broad spectrum of tissues, but the accuracy varies depending on the tissue type. In the field of forensics, the prediction accuracy of a model considered high if the MAD is less than 5 years, models for age prediction in blood, saliva, semen have been reported using only a few age-associated CpGs, that measure the biological age and provide information about life expectancy as well as unknown sample donor’s appearance [18]. We believe, in the future, DNA methylation markers will play an important role in the forensic investigations as well as in clinical research.

References