Autosomal SNP typing of forensic samples with the GenPlex(TM) HID System: Results of a collaborative study


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ABSTRACT

The GenPlex™ HID System (Applied Biosystems – AB) offers typing of 48 of the 52 SNPforID SNPs and amelogenin. Previous studies have shown a high reproducibility of the GenPlex™ HID System using 250–500 pg DNA of good quality. An international exercise was performed by 14 laboratories (9 in Europe and 5 in the US) in order to test the robustness and reliability of the GenPlex™ HID System on forensic samples. Three samples with partly degraded DNA and 10 samples with low amounts of DNA were analyzed in duplicates using various amounts of DNA. In order to compare the performance of the GenPlex™ HID System with the most commonly used STR kits, 500 pg of partly degraded DNA from three samples was typed by the laboratories using one or more STR kits. The median SNP typing success rate was 92.3% with 500 pg of partly degraded DNA. Three of the fourteen laboratories counted for more than two thirds of the locus dropouts. The median percentage of discrepant results was 0.2% with 500 pg degraded DNA. An increasing percentage of locus dropouts and discrepant results were observed when lower amounts of DNA were used. Different success rates were observed for the various SNP typing kits. The rs763869 SNP was the least successful. With the exception of the Minifiler™ kit (AB), GenPlex™ HID performed better than five other tested STR kits. When partly degraded DNA was analyzed, GenPlex™ HID showed a very low mean match probability, while all STR kits except Minifiler™ had very limited discriminatory power.

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1. Introduction

The usefulness of SNP typing in forensic genetics has previously been pointed out in several publications [1–3]. Short DNA fragments with SNPs can be amplified, which makes it possible to analyze SNPs in partly degraded DNA samples [1,4]. Moreover, the low mutation rate of SNPs is an important advantage in kinship analysis [3]. The SNPforID consortium [5] selected 52 biallelic SNP markers with high levels of polymorphism in the major population groups [1]. Short amplicons (up to 115 bp) were used to analyze the 52 SNPs and the mean match probability was at least $5.0 \times 10^{-19}$. Several platforms can be used to analyze SNP typing kits [6–10]. The most widely used assay is the single base extension (SBE) assay using the SnAShot™ kit.
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GENPlex™ HID System (AB) and the SBE-based 49plex SNP assay [12]. Concordant results were obtained from at least two of the investigations. The results obtained by the organizing laboratory were used as reference. The protocols were approved by the Danish ethical committee (KF-01-037/03).

2.2. DNA extraction and quantification

DNA from the three samples with partly degraded DNA was extracted using a standard phenol–chloroform procedure. DNA from blood samples from 10 healthy donors was extracted from 15 µL of blood using the MagAttract DNA blood midi M48 kit (Qiagen GmbH, Hilden, Germany) and a Tecan Freedom EVO® robot (Tecan trading AG, Switzerland) [17]. The DNA samples were quantified with the Quantifiler® Human DNA Quantitation kit (AB) using an AB 7900 (AB) real time PCR analyzer according to the manufacturer’s recommendations.

2.3. Shipment of samples and reagents

DNA extracts and the critical reagents needed for SNP typing with the GenPlex™ HID System (AB) were sent on dry ice by courier to the participating laboratories.

2.4. GenPlex™ HID SNP typing

The participating laboratories typed the degraded DNA samples in duplicate with the GenPlex™ HID System using 500 pg, 250 pg, 100 pg, 50 pg and 20 pg of DNA in the PCR reaction. Ten blood DNA extracts were SNP typed in duplicate by the participating laboratories with the GenPlex™ HID System adding 50 pg, 25 pg, 10 pg, 5 pg and 2 pg of DNA to the PCR reaction. The DNA titrations were prepared by each participating laboratory.

Details on the GenPlex™ HID Procedure are published elsewhere [15]. Briefly, a multiplex PCR reaction was performed using 2 µL of DNA with amounts indicated above. All laboratories performed the PCR reaction in a GeneAmp® PCR System 9700 thermal cycler (AB) according to the manufacturer’s recommendations. Leftover primers and unused DNTPs were removed from the PCR products using 2 µL of ExoSAP-IT® (USB Corp., Cleveland, OH, USA). An oligo ligation assay (OLA) was performed using the PCR products as templates. The OLA assay required the phosphorylation of the oligos. This may be performed at the same time as the OLA assay or as a step preceding the OLA assay. Twelve of the fourteen participating laboratories phosphorylated the oligos prior to the OLA assay, while two of them performed the phosphorylation reaction and the OLA assay in a single step (Table 1). All the

Table 1

Details of the GenPlex™ HID protocol.

<table>
<thead>
<tr>
<th>Lab #</th>
<th>Previous experience with GenPlex</th>
<th>ABI sequencer</th>
<th>No. of steps in the OLA reaction</th>
<th>OLA product (µL)</th>
<th>Sample loading reagent (µL)</th>
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<tr>
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<td>Yes</td>
<td>3130xl</td>
<td>2</td>
<td>10</td>
<td>17.5</td>
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<tr>
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<td>17.5</td>
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<tr>
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<tr>
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<td>3130xl</td>
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<td>5</td>
<td>10</td>
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<tr>
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<td>17.5</td>
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<td>10</td>
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<td>Yes</td>
<td>3130xl</td>
<td>2</td>
<td>10</td>
<td>17.5</td>
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</tbody>
</table>
participating laboratories performed the OLA assay in a GeneAmp® PCR System 9700 thermal cycler (AB) according to the manufacturer’s recommendations. Five or ten microliters of biotinylated OLA products (Table 1) were added to a streptavidin coated capture microtiter plate (AB). Hybridization of ZipChute® probes was performed on the captured OLA products. A volume of 10 μL or 17.5 μL of sample loading reagent (AB) was used to elute the ZipChute® probes (Table 1). Twelve of the fourteen laboratories used an ABI 3130xl Genetic Analyzer (AB) to separate the eluted ZipChute® probes. One laboratory used an ABI 3130 (AB) and one laboratory used an ABI 3730xl (AB) (Table 1). All laboratories used POP-7™ polymer (AB), and all laboratories used 36 cm capillaries except for one that used 50 cm capillaries. Capillaries that had not previously been used were pre-conditioned using a pre-conditioning kit (AB).

2.5. STR typing

In order to compare the efficiencies of the GenPlex™ HID System and STR kits on degraded DNA samples, all laboratories typed 500 pg of partly degraded DNA samples in duplicates with one or more of the following STR kits: Identifier™ (AB) [6 laboratories], SGM Plus™ (AB) [5 laboratories], SEfiler Plus™ (AB) [3 laboratories], MiniFiler™ (AB) [2 laboratories], PowerPlex 16® (Promega) [2 laboratories], PowerPlex ES® (PPES) (Promega) [1 laboratory], according to the usual procedures of the laboratories.

2.6. Data collection

All the results were collected and analyzed by the organizing laboratory, GeneMapper™ 4.0 (AB) was used to call peaks higher than 10 relative fluorescent units (RFUs) and the results were further analyzed using Excel© (Microsoft). In house criteria for allele calling were established. Peak heights lower than 10 RFUs were classified as “not amplified”. Peak heights below 100 RFUs for presumed homozygous loci and below 50 RFUs for presumed heterozygous loci were classified as “weak”. All systems that were classified as “not amplified” or “weak” were considered as locus dropouts. Peak height ratios were calculated for each genotype by dividing the higher peak height by the lower peak height. Unbalanced peak height ratios (ratios between 7.0 and 12.0) were classified as “uncertain assignments”. Heterozygotes were identified when the peak height ratio was less than 7:1. Homozygotes were identified when the peak height ratio was greater than 12:1.

The success rate was defined by the proportion of reported SNP types that were identical to the reference genotypes. Discrepant results were defined by the number of SNP types that were not identical to the reference genotypes. Three kinds of discrepancies were observed: (i) allele dropout (heterozygous loci that only showed one of the two expected peaks), (ii) allele drop-in (homozygous loci that showed two peaks; one corresponding to the real allele and a spurious peak), and (iii) allele dropout plus allele drop-in (apparent homozygous loci with one peak that did not correspond to the real allele).

The data were grouped and analyzed in two ways. First, all the data generated by each participating laboratory were collected, analyzed and compared to the reference genotypes (i.e. 69,180 SNP types corresponding to 49 SNPs from 13 samples analyzed in duplicates by 14 laboratories using 5 DNA dilutions). Second, consensus SNP types were obtained from the duplicated investigations reported by each laboratory (i.e. 44,590 intra-laboratory consensus SNP types). The intra-laboratory consensus SNP types were compared with the reference genotypes.

SNP and STR profiles obtained from 500 pg of degraded DNA were compared by calculating percentages of full profiles, i.e. number of alleles observed in relation to the maximum number of alleles that are present in a full profile.

2.7. Statistical analysis

The results were further analyzed using box and whisker plots with the SPSS statistical package (SPSS Inc., Chicago, IL, USA). The box and whisker plot summarizes the overall range of the data graphically. Five values are used to construct a box and whisker plot: (1) the mean value (represented by an horizontal line that divides the box), (2) the first quartile (the bottom of the box), (3) the third quartile (the top of the box); the length of the box (distance between the first and the third quartiles) is the inter-quartile range (IQR), (4) the ‘maximum value’, up to 1.5 IQRs from the top of the box (represented by a whisker) and (5) the ‘minimum value’, up to 1.5 IQRs from the bottom of the box (represented by a whisker). Values that were between 1.5 and 3 IQRs from the end of

Fig. 1. Box-plot of the success rates of the GenPlex™ HID System typing results reported by 14 laboratories. Degraded DNA (a) and DNA from blood samples (b).
a box are called extreme values and were represented by circles. Outliers were defined by the values that were more than 3 IQRs from the end of the box and were represented by asterisks.

In order to compare the results obtained with the GenPlex™ HID System and the various STR kits, a median Polish analysis [18] was performed using the statistical software Minitab™ 15.

3. Results and discussion

3.1. The reported results: success rates and discrepant results

The 14 laboratories reported 3377 SNP types (82%) with 500 pg of partly degraded DNA from the three samples. The success rate ranged from 15.6% to 98.6% (median: 92.3%) among the 14 laboratories (Supplementary material 1). Laboratories # 3, 4 and 9 showed success rates clearly lower than the average of the other 11 laboratories. If the results from laboratories 3, 4 and 9 were not taken into account, an overall percentage of 94.6% of the expected SNP types were reported with an overall success rate of 94.2% (Supplementary material 1). The low success rate reported by laboratories 3, 4 and 9 can mainly be explained by technical problems experienced by the three laboratories during the analysis of the samples (e.g. evaporation of the PCR products and prolonged waiting time between the various protocol steps). Although the GenPlex™ HID protocol was not considered complicated by the participating laboratories, multiple steps made the process laborious and time consuming. All laboratories performed the protocol manually. As previously discussed [14,16], the use of a robot decreases the laboriousness of the protocol and would most likely increase the success rate. The success rate decreased when less than 500 pg of DNA was used in the PCR reaction (Fig. 1a). With the exception of laboratory # 9, low percentages of discrepant

![Fig. 2. Box-plot of the discrepant results observed in the results reported by the 14 laboratories typing various amounts of degraded DNA (a) and DNA from blood samples (b) with the GenPlex™ HID System.](image)

![Fig. 3. Box-plot of the percentage of intra-laboratory results with no consensus SNP types reported by the 14 laboratories typing various amounts of degraded DNA (a) and DNA from blood samples (b) with the GenPlex™ HID System.](image)
results were reported when 500 pg of partly degraded DNA was analyzed (median: 0.2%, Supplementary material 1). After a closer analysis of the results reported by laboratory 9, we suspect that this laboratory switched two of the DNA samples (degraded samples 2 and 3). Low numbers of discrepant results were observed with 500 pg and 250 pg DNA, but it increased with 100 pg and less DNA reaching a median value of 3.4% with 20 pg of DNA (Fig. 2a).

For the 10 blood DNA samples, 10,370 SNP types (75.6%) were reported by the 14 laboratories for 50 pg of DNA (Supplementary material 2). The success rate decreased with smaller amounts of DNA (Fig. 1b) not only due to an increasing number of locus dropouts, but also due to an increasing proportion of discrepant SNP types (Fig. 2b). Thus, the use of small amounts of DNA strongly decreased the reliability of the results. When 5 pg DNA was used, 5.3% of the results were discrepant. This value decreased when 2 pg of DNA was used due to the few data reported for 2 pg of DNA.

3.2. The unreported results: locus dropouts and uncertain assignments

When 500 pg of partly degraded DNA was analyzed, 739 SNP types (18%) were not reported. Uncertain SNP typing results were observed in 1.6% of the results and 16.4% of the SNP loci dropped out (Supplementary material 1). The number of uncertain assignments could most likely be decreased by adjusting the SNP calling criteria. Nevertheless, this is only possible for laboratories that routinely use the GenPlex™ HID System. Three of the fourteen laboratories (labs # 3, 4 and 9) counted for more than two thirds of the locus dropouts. If the results of these three laboratories were removed, the overall number of locus dropouts decreased from 16.4% to 3.8% in the remaining 11 laboratories (Supplementary material 1). In these 11 laboratories, the percentage of locus dropouts observed with 500 pg of DNA ranged from 0% (observed three times) to 11.9% (median: 1%). The

![Fig. 4: Locus dropouts, discrepancies and uncertain assignments observed in the 48 SNPs and amelogenin included in the GenPlex™ HID System. Global results correspond to the analysis of 500 pg of degraded DNA by 11 of the 14 participating laboratories. Results reported by laboratories 3, 4 and 9 were not included.](image-url)
number of unreported results was increased to 2024 (49.2%) with 20 pg DNA.

For the 10 blood DNA samples, the total number of unreported results varied from 24.4% with 50 pg DNA (Supplementary material 2) to 85.7% with 2 pg DNA. With 50 pg DNA, locus dropouts were observed in 22.3% of the results, while uncertain assignments represented 2.1% of the results. There was a large variation in the amount of locus dropouts reported by each laboratory (0.1–62.3%; median: 16.3%; Supplementary material 2), most likely due to the stochastic effect of the use of small amounts of DNA in the PCR reaction [19].

3.3. Analysis of consensus SNP types

We tested the intra-laboratory reproducibility of the results by comparing the duplicated runs reported by each participating laboratory. Fig. 3 represents the percentage of intra-laboratory results with no consensus SNP types. In most part of the cases, an intra-laboratory consensus was not reached due to the lack of results in one or two runs. Less often, it was due to discrepant results. A maximum number of 2058 consensus SNP types were expected with 500 pg of degraded DNA and a total number of 1557 consensus SNP types were obtained (75.7%). If the results of laboratories 3, 4 and 9 were removed, intra-laboratory consensus SNP types were reached in 91.2% of the SNP types. A total of 20 inter-laboratory discrepancies (1%) were observed if all 14 laboratories were considered, but only two (0.1%) if the 11 laboratories were taken into account. With the 10 blood DNA samples, 4366 consensus SNP types (63.6%) were obtained with 50 pg DNA. Intra-laboratory discrepancies increased with lower amounts of DNA in the same way as did the inter-laboratories discrepancies.

3.4. Performance of the various SNPs in the GenPlex™ HID System

The performance of the various SNP loci was analyzed using the results that showed the highest success rates (results from 500 pg degraded DNA). The results reported by laboratories 3, 4 and 9 were not taken into account due to their poor quality. Fig. 4 shows the percentage of locus dropouts, uncertain assignments and discrepant results observed for each SNP. Twenty out of the 49 loci in the GenPlex™ HID System showed success rates of 100% and the lowest success rate was 69.7% (rs763869). Locus dropout was the main cause of unsuccessful results but, in 7 of the SNPs, the percentage of uncertain results and discrepancies was higher than that of locus dropouts. Discrepant results were observed in six of the 49 SNP loci (rs1886510, rs2107612, rs2831700, rs719366, rs717302, and rs1335873), and in all but one (rs1335873) the discrepancies were due to allele drop-ins. The SNP typing of some of the loci (especially rs1886510, rs2107612 and rs2831700) was less reliable due to the presence of a spurious peak that did not correspond to the real allele. As mentioned above, the SNP assignment of some of the loci could probably be improved by a re-adjustment of the allele calling criteria, but not all uncertain and discrepant results could be corrected in this way. A previous study [16] showed a poorer performance of some of the SNPs in the GenPlex™ HID System, especially rs907100, and an improvement of the GenPlex™ HID protocol was suggested. Based on the results of the present study, an improvement in the protocol is still needed, mainly in order to balance the reactions and avoid locus dropouts and spurious peaks that may simulate real alleles.

3.5. GenPlex™ HID results vs. STR results

The 14 participating laboratories analyzed 500 pg of three partly degraded DNA samples in duplicates using one or more STR kits in order to compare the performance of the GenPlex™ HID System with the most commonly used STR kits. The percentages of consensus profiles were calculated (Fig. 5) and the median polish results were used to compare the success rates of the various systems. Table 2 shows the results of the median polish analysis for each kit (common effect of samples and laboratories). With the exception of MiniFiler™ (AB), GenPlex™ HID performed better than the other five STR kits showing a median percentage profile of 91.3%. Although the MiniFiler™ kit showed good results, the partial profiles obtained with the GenPlex™ HID System were more informative than the results obtained with MiniFiler™ (Table 2). The SEfiler Plus™ kit (AB)
The authors showed the best results after MiniFilerTM and the GenPlexTM HID System.

4. Conclusions

Partial but reproducible SNP profiles were obtained from 250 pg to 500 pg naturally degraded DNA using the GenPlex™ HID System in 11 of 14 laboratories. Three of the 14 participating laboratories encountered technical problems during the typing process. Due to the limited amount of samples and reagents available for this study, it was not possible for the laboratories to repeat the investigations. The quality of the results reported by the laboratories was not related to their experience with the GenPlex™ HID protocol or to small alterations in the protocol (e.g. the phosphorylation step). With the exception of the MiniFiler™ kit, the GenPlex™ HID System performed better than the tested STR kits, although the lower match probability was obtained with the GenPlex™ HID System. The GenPlex™ HID protocol is a laborious and time consuming method but it is not difficult to implement in practice since most standard forensic genetic laboratories already have the great majority of the necessary equipment. The results obtained in this collaborative exercise shows that the GenPlex™ HID method is a reliable method for SNP typing also when partly degraded samples are analyzed. Nevertheless, still some optimization is required in order to avoid locus dropouts, allele drop-ins and allele dropouts. Most likely an improvement of the PCR and/or OLA reactions will be needed to obtain a better balance of the heights of the peaks of the various SNP markers. As previously suggested, an improvement in the SNP allele calling method implemented in the GeneMapper™ software would also facilitate the SNP typing.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.fsigen.2010.06.007.

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