

Development of forensic genomics research toolkits by the use of Massively Parallel Sequencing

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Chapter 2

TSSV: a tool for characterization of complex allelic variants in pure and mixed genomes

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Abstract

Motivation: Advances in sequencing technologies and computational algorithms have enabled the study of genomic variants to dissect their functional consequence. Despite this unprecedented progress, current tools fail to reliably detect and characterize more complex allelic variants, such as short tandem repeats (STRs). We developed TSSV as an efficient and sensitive tool to specifically profile all allelic variants present in targeted loci. Based on its design, requiring only two short flanking sequences, TSSV can work without the use of a complete reference sequence to reliably profile highly polymorphic, repetitive or uncharacterized regions.

Results: We show that TSSV can accurately determine allelic STR structures in mixtures with 10% representation of minor alleles or complex mixtures in which a single STR allele is shared. Furthermore, we show the universal utility of TSSV in two other independent studies: characterizing de novo mutations introduced by transcription activator-like effector nucleases (TALENs) and profiling the noise and systematic errors in an Ion Torrent sequencing experiment. TSSV complements the existing tools by aiding the study of highly polymorphic and complex regions and provides a high-resolution map that can be used in a wide range of applications, from personal genomics to forensic analysis and clinical diagnostics.

Availability and implementation: We have implemented TSSV as a Python package that can be installed through the command-line using pip install TSSV command. Its source code and documentation are available at https://pypi.python.org/pypi/tssv and http://www.lgtc.nl/tssv.

Introduction

As a consequence of various mechanisms such as DNA recombination, replication and repair-associated processes, the spectrum of human genetic variation ranges from single nucleotide differences to large chromosomal events. Among the different types of genetic changes, repetitive DNA sequences show more polymorphism than single nucleotide variants (Conrad et al., 2010; Hinds et al., 2006; lafrate et al., 2004; Kidd et al., 2008; Redon et al., 2006; Sebat et al., 2004; Tuzun et al., 2005), and they are important in human diseases (Conrad et al., 2010; de Cid et al., 2009; Girirajan et al., 2011; Hollox et al., 2008; McCarroll et al., 2009; Pinto et al., 2010), complex traits and evolution (Mills et al., 2011; Stephens et al., 2011; Sudmant et al., 2010). In particular, microsatellite variants, also known as short tandem repeats (STR), and their expansion/shortening have been linked to a variety of human genetic disorders (Mirkin, 2007; Pearson et al., 2005; Sutherland and Richards, 1995), and have been used in genotyping (Kimura et al., 2009; Weber and May, 1989) and forensic DNA fingerprinting studies (Kayser and de Knijff, 2011; Moretti et al., 2001).

Because of the repetitive nature of STRs and often the low level of complexity of the DNA sequences in which they occur (Treangen and Salzberg, 2012), characterization of STR variability and understanding of their functional consequences are challenging (Weischenfeldt et al., 2013), So far, sequencing-based strategies have focused on reads mapped to the reference genome and subsequent identification of discordant signatures and classification of associated STRs (Medvedev et al., 2009; Mills et al., 2011). Yet, the mainstream aligners, such as BWA (Li and Durbin, 2009) or Bowtie (Langmead and Salzberg, 2012), do not tolerate repeats or insertions and deletions (indels) as a trade-off of run time (Li and Homer, 2010). This limitation leads to ambiguities in the alignment or assembly of repeats which, in turn, can obscure the interpretation of results (Treangen and Salzberg, 2012). Moreover, the current human genome reference still remains incomplete and provides only limited information on expected and potentially uncharacterized STRs in different individuals (Alkan et al., 2011; lafrate et al., 2004; Kidd et al., 2008; Sebat et al., 2004). Consequently, STRs are not routinely analyzed in wholegenome or whole-exome sequencing studies, despite their obvious applications and their role in human diseases, complex traits and evolution.

Here, we present a method for targeted profiling of STRs that reports a full spectrum of all observed genomic variants along with their respective abundance. Our tool, TSSV, can accurately profile and characterize STRs without the use of a complete reference genome, and therefore minimizes biases introduced during the alignment and downstream analysis. TSSV scans sequencing data for reads that fully or partially encompass loci of interest based on the detection of unique flanking sequences. Subsequently, TSSV characterizes the sequence between a pair of non-repetitive flanking regions and reports statistics on known and novel alleles for each locus of interest. We show the performance of TSSV on robust characterization of all allelic variants in a given targeted locus by its application in several case studies: forensic DNA fingerprinting of mixed samples by STR profiling, characterization of variants introduced by transcription activator-like effector nucleases (TALENs) in embryonic stem (ES) cells and detailed characterization of errors derived from a next-generation sequencing (NGS) experiment.

Material and Methods

TSSV algorithm

The algorithm expects a FASTA file containing sequencing data and a library containing a list of loci of interest that are described by two unique sequences flanking a target locus in the form of a simple regular expression. The description of targeted loci consists of a series of triplets (i.e. CTTA 2 5), each containing a sequence followed by two integers that denote the minimum and maximum number of times the preceding sequence is expected. The notation of expected alleles is then compiled into a regular expression that is used to distinguish between known and new alleles. It is important that a library that contains a description of loci of interest according to the aforementioned instruction should be customized and provided. TSSV reports an overview of marker pair alignments and a detailed description of the identified alleles and their respective frequency per strand. TSSV also provides supporting reads of each locus of interest in separate FASTA files.

TSSV is an open source Python package that can be easily incorporated in any standard NGS pipeline. In addition, we have made the Python package fastools available at https://pypi.python.org/pypi/fastools. fastools offers a series of functions to manipulate, characterize, sanitize and convert FASTQ/FASTA files to other formats. Therefore, it can be used to convert FASTQ files to TSSV desired format (FASTA). For further information on usage and generated data see Supplementary Table S1.

Marker alignment

Each pair of markers (unique flanking sequences) is aligned to the reads by using a semi-global pairwise alignment, a modified version of the Smith–Waterman algorithm (Smith and Waterman, 1981). The alignment matrix is initialized with penalties only for the aligned sequence and not for the reference sequence. By using this approach, we can use the alignment matrix to calculate the edit distance between the aligned sequence and all substrings of the reference sequence. Finally, TSSV uses the alignment matrix to select the rightmost alignment with a minimum edit distance. To guarantee symmetry with regard to reverse complement sequences, TSSV aligns the reverse complement of the right marker to the reverse complement of the reference sequence.

Allele identification

Once TSSV successfully aligns a marker pair to either the forward or the reverse complement of the reference sequence, the region of interest is selected by extracting the sequence between the alignment coordinates, which is then converted to the forward orientation. The target variable sequence is then matched to the regular expression of the corresponding marker pair for classification as either a known or a new allele. In case of partial identification of markers (i.e. only the left or right marker of the pair is identified), the input sequence is flagged as having either no beginning or no end. The assessment of required runtime for TSSV to identify alleles in datasets with different sequencing depth is provided in Supplementary Figure S1. Each dataset is profiled to characterize 16 allelic STR structures. It should be noted that currently TSSV uses a single processor for the analysis.

Annotations

Once a list of new alleles is constructed, TSSV uses a revised version of the Mutalyzer online service (Wildeman et al., 2008; https://mutalyzer.nl) to describe all observed variants compared with the reference sequence. Mutalyzer provides a description of observed variants according to the Human Genome Variation Society format for sequence variant description. This can be used to provide an overview of most frequent mutations that are observed within each locus of interest.

Interpretation guidelines

TSSV provides the frequency in which each allelic structure is observed on plus and minus strand. Based on the experimental design, the frequencies of allelic variants and the balance between supporting reads on the plus and minus strand can aid the identification of potential sequencing biases. Moreover, based on the choice of sequencing technology, homopolymers are prone to introducing artificial allelic structures, so it is advised, when possible, to allow for a tolerance of a few base difference in the homopolymer length while describing targeted loci. The estimation of a lower boundary for the identification of variant alleles is subject to the experimental design. Thus, sequencing of control samples, if possible, can aid a more reliable analysis by ruling out potential slippage and background noise. nce a list of new alleles is constructed, TSSV uses a revised version of the Mutalyzer

Availability

TSSV is available at http://www.lgtc.nl/tssv and https://pypi.python.org/pypi/tssv. It can also be installed through the command line: pip install tssv. All original datasets and the analysis results can be obtained from figshare (http://www.figshare.com): detection of STRs, SNPs and short indels (Anvar, 2013a), determining de novo structural variations (SVs) in TALEN-treated ES cells (Anvar, 2013b), characterization of STRs (Anvar, 2013c) and detection of systematic errors in PGM (Anvar, 2013c).

Library preparations and sequencing

STR PCR products for sequencing were generated using the Powerplex[®] 16-kit from Promega (commercial assay designed and optimized for fluorescent dye-based fragment analysis of STR loci) and were purified with Ampure XP beads according to manufacturer's protocol. Library preparation was performed using the Rapid Library Preparation Kit (Roche). Emulsion PCR and sequencing were performed on the FLX Genome Sequencer (454/Roche) according to the protocol provided by the manufacturer.

PCR products for sequencing of all other samples on the Personal Genome Machine (PGM, Ion Torrent) were prepared using the Ion Plus Fragment Library Kit or amplicon fusion primers. Emulsion PCR was performed using the OneTouch (OT I, Ion Torrent). Sequencing was performed according to LifeTech protocol using the Ion PGM™ 200 Sequencing Kit. PCR reaction was done in 10 µl containing I × FastStart High Fidelity reaction buffer (Roche), I.8 mM MgCl2, 2% DMSO, 200 µM dNTPs, 0.5 U FastStart High Fidelity Enzyme Blend (Roche), 20 ng DNA, 300 nM universal barcoding primer, 300 nM reverse target primer and 30 nM forward target primer. After I0 min of initial denaturation at 95°C, 30 PCR cycles were performed at 20s 95°C, 30s 60°C and 40s 72°C. Primer sequences are provided in Supplementary Table S2.

TALEN design and transfection

The TALENs -pair targeting intron 52 of the human DMD gene (hDMD) was designed using the TALEN toolbox described by Cermak et al. (2011), Next, hDMD/ mdx ES cells (t Hoen et al., 2008; Veltrop et al., 2013) were transfected with the TALENs plasmids without any homologous recombination vector. ES cells were routinely cultured on murine embryonic fibroblast (MEF) feeder cells in knockout DMEM supplemented with 2 mM L-glutamine, I mM sodium pyruvate, non-essential amino acids, 50 units of penicillin as well as streptomycin, 1000 units of leukemia inhibitory factor and 10% fetal bovine serum (FBS Gold, all from Life Technologies Ltd). Per TALEN, total of 750 ng in 1.5 µg of DNA was used to transfect 1 000 000 hDMD/mdx ES cells using Lipofectamin 2000 (Invitrogen), DNA-Lipofectamin 2000 suspension was prepared in serum and antibiotic-free medium according to the supplier's manual Cells were incubated for 30 min in suspension with the DNA-Lipofectamine mixture and then plated in two 9 cm culture dishes coated with MEF in regular ES culture medium. ES cells were cultured for a week, and DNA was isolated from a pool of 1500 ES clones. This DNA was then prepared for sequencing using Ion Torrent PGM according to the instrument guidelines.

Results

Characterization of STRs

We tested the performance of TSSV in characterizing known STRs from Roche/454 targeted sequencing data of 16 STR loci, amplified in a multiplex reaction. To demonstrate the added value of TSSV over mainstream aligners, we generated four sequencing libraries of which two consisted of pure individual samples and two mixtures in the ratios of 50:50 and 90:10 with comparable depth of coverage (Supplementary Table S3). A full spectrum of STR structures and their abundance was generated after a semi-global alignment of the 25 bp flanking regions adjacent to the STR structure, with tolerance of up to three mismatches (Fig. 1A). On average, 8% of reads remained uncharacterized, mostly because the sequences did not cover both flanking reference sequences or that sequences contained too many mismatches for regions that are required for identification of unique flanking reference sequences (Supplementary Table S3). The PCR product used for preparing the sequencing libraries were generated using the Powerplex 16-kit from Promega, which is an assay designed and optimized for fluorescent dye-based fragment analysis of STR loci. This resulted in a strong imbalance in sequencing yield between STR markers with different dyes in the fragment analysis (Supplementary Table S3). Thus, we restricted the analysis to the three markers with highest coverage (D3S1358,TH01 and D13S317). Frequencies of the observed alleles were interpreted to distinguish actual alleles from slippage artifacts (Supplementary Tables S4–S6).

For D3S1358 (TCTA1 TCTG1-3 TCTA12–13), TSSV robustly identified the STR structure associated with each of the samples, with >91% of reads supporting the presence of two STR alleles (Fig. 1B). In addition, TSSV could pick up a minor frequency (7.25%) for alternative STR structures, in which the DNA amplicons show false STR structures because of DNA polymerase slippage during the amplification (Ellegren, 2004; Hauge and Litt, 1993). Despite the presence of PCR amplification artifacts, the major and minor STR structures in balanced and more extreme mixtures (50:50 and 90:10, respectively) could accurately be identified by TSSV (Fig. 1B and Supplementary Table S4).

We next explored whether TSSV can correctly detect alleles of more complex cases differing based on STR length (ATCT12ATCA2 and ATCT11ATCA3) or composition (CATT9 and CATT3CAT1CATT6) as well as mixtures that shared one allelic STR structure (CATT3CAT1CATT6). Markedly, TSSV could correctly detect, characterize and quantify reads supporting all STR alleles, including mixtures with only 10% representation of the minor alleles (i.e. D3S1358 markers) and more complex

mixtures (TH01 markers) where a single STR allele is shared (Fig. 1C, Supplementary Tables S4–S6 and Supplementary Figs S2–S4). Results of the remaining STR markers are provided in supplementary materials, Supplementary Tables S7–S17.

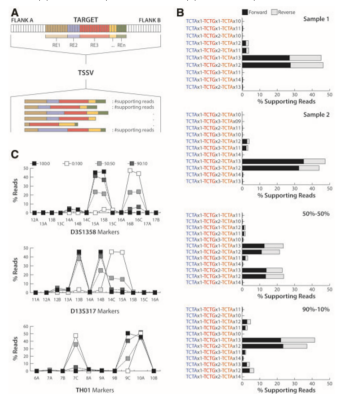


Fig. 1. Characterization of allelic STR structures in samples and their mixtures of differing ratios (A) Schematic representation of STR structure identification and quantification. After proper alignment of two flanking sequences, TSSV performs a strand-specific classification and quantification of repetitive elements (RE) that constructs a given STR-structure. (B) The number of sequencing reads that support the presence of different allelic D3S1358 STR structures on both strands. Pure samples and their mixtures in two different ratios are presented separately. (C) The proportion of reads that support different allelic STR structures for three most abundant markers (D3S1358, D13S317 and THO1). STR markers differ in complexity based on STR length or composition as well as mixtures in which one allelic STR structure in shared.

Determining de novo structural variations in TALEN-treated cells

TALENs have shown promising potential in site-specific genome editing (Boch, 2011; Cermak et al., 2011; Miller et al., 2011; Zhang et al., 2011). Their modular structure enables simple construction of TALENs that can specifically recognize virtually any DNA sequence of interest. On delivery of a TALENs-pair, a double strand break is introduced that is repaired by non-homologous end-joining, introducing a large variety of mutations (Supplementary Fig. S5). Because the method lacks a positive selection procedure, the applicability depends largely on its efficacy. We used TSSV to estimate the efficiency of genome editing in ES cells from a mouse model with the hDMD, stably integrated in the mouse genome (t Hoen et al., 2008), and determine the utility of an assembled TALEN pair (Supplementary Table S18) in introducing mutations within targeted intron 52 of the hDMD (Supplementary Fig. S6).

For 100,000 TALENs-transfected and non-transfected (control) ES cells, a 135 bp fragment encompassing the entire targeted locus was PCR amplified and sequenced using the Ion Torrent PGM (Supplementary Table S19). The targeted locus was covered over 450 000 times, which allows for precise detection and characterization of any variant present. From the control ES cells, we determined a background of 3.1% of reads that contain at least one mismatch, derived from sequencing errors and potential spontaneous mutations (Fig. 2A and Supplementary Table S19). In TALENs-treated ES cells, the rate of sequencing reads that contain at least one mismatch was 11.4%, almost 4-fold higher than controls (Fig. 2A). The majority of mutations introduced by TALENs pair were small insertions and deletions (75.6%; excluding duplications) (Fig. 2B), which is consistent with the expected type of variants introduced by TALENs (Cermak et al., 2011). The frequency in which individual variants occurred was specific to TALENtreated ES cells, even for those that were observed with very low frequency (Fig. 2C and D). However, we observed a few mutations that were not specific to TALEN-treated ES cells (Fig. 2D). These were mainly duplications that arose from inaccurate detection of homopolymer stretches. Overall, TSSV results indicate significant enrichment (P = 2.85E-09; Kolmogorov-Smirnov test) of variants in TALEN-treated ES cells as compared with controls (Fig. 2E). Furthermore, TSSV reported a list of the most frequent variants and cleavage sites, majority of which were either exclusive to TALEN-treated ES cells or with over 3-fold higher frequency in TALEN-treated ES cells than controls (Supplementary Fig. S7). The Ion Torrent variant caller (version 3.2) did not report any variant because of the nature and frequency of variants introduced by TALENs.

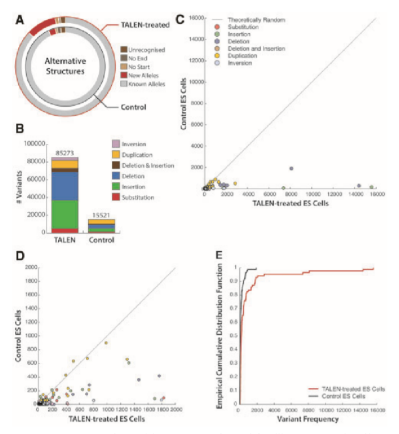


Fig. 2. Variant characterization and quantification of TALEN-treated and Control ES Cells. (A) Basic statistics of the TSSV analysis. Pie charts show the proportion of sequencing reads that support the presence of new alleles in TALEN-treated (outer circle) or control (inner circle) ES Cells. No Start and No End fragments represent reads in which one of the flanking sequences was not recognized. (B) Total number of variants in TALEN-treated ES Cells and Control, grouped by type. (C) Comparative analysis of the number of occurrences for individual variants in both samples. Data points are colored based on the type of variation. (D) Zoomed in scatter plot for variants with frequencies lower than 2000. (E) Empirical cumulative distribution of variant frequencies for TALEN-treated (red) and Control ES Cells (black). Kolmogorov–Smirnov test was performed to assess if two distributions are significantly different.

Detection of systematic errors in PGM Ion Torrent

During the targeted Ion Torrent resequencing of exon 19 of the DMD gene (X-chromosome) in five male patients and a female carrier, we observed a number of shared and unexplained heterozygous variants given that male patients have only one X-chromosome and DMD gene does not locate within pseudo-autosomal regions. We used TSSV to provide a high-resolution map of all sequence variants as a way to understand the origin of these artifacts (Fig. 3A). To assess the reproducibility of our findings, we performed two independent Ion Torrent PGM sequencing runs (PG090 and PG109). Two different versions of the Ion Torrent base-calling algorithm were used

for PG090 (versions 2.2 and 3.0) while PG109 was only processed by version 3.0 (sequencing run was carried out after the upgrade of the lon Torrent Suit). The three datasets enabled us to investigate potential artifacts derived from sequencing and/or different base-calling algorithms. Our first observation indicated a significant decrease in the total number of reads (average of 11.3 and 13.3% in respective to different runs and base-calling algorithms) that were recognized per individual (Supplementary Table S20). We also noticed a significant difference in the fraction of reads per dataset (44.3, 40.3 and 48.7%) that were reported as new alleles, having at least one mismatch with the reference sequence (Fig. 3B).

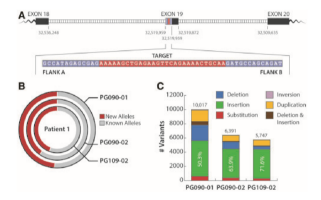


Fig. 3. Identification of mutations within exon 19 of DMD gene. (A) Schematic representation of the locus of interest for resequencing, the design of unique flanking sequences (blue), and the targeted region (red) to be profiled using TSSV. (B) Pie charts show the proportion of reads that support the presence of new alleles (red) in sequencing library of patient 1. Pie charts represent different sequencing runs (PG090 or PG109) or the base-calling algorithm used during the primary analysis (01 or 02). The two most outer pie charts are sequencing reads from the same PGM Ion Torrent run processed using two different versions of base-calling algorithm. The most inner pie chart represents an independent run of the same library. (C) Number of observed variants separated by variation type. Percentages show the proportion of insertion events from the total number of variants in each set.

We observed a significant reduction of variants (36.2%) after adoption of the version 3.0 base-caller, mainly affecting the level of deletions and duplications calls (Fig. 3C). This prominent decrease (68.3 and 48.9%) arises from improvement of the algorithm in determining the length of homopolymer stretches. Notably, the majority of other variants were single nucleotide insertions (excluding duplications and indels) that remained at a comparable rate across different datasets (Fig. 3C). Next, we assessed the strand specificity of the variants based on the sequencing direction. Interestingly, while the majority of variants showed a similar frequency in both directions, the most frequent variants showed a clear imbalance between forward and reverse strand (Fig. 4A and B). The observed strand-specific bias was reproducible and was not influenced by the software version, as it was observed in all three datasets (Fig. 4C and D and Supplementary Figs S8–S11).

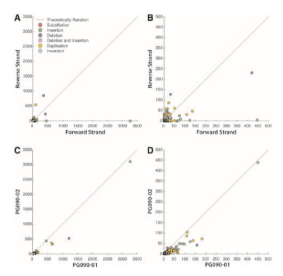


Fig.4. Comparative analysis of observed mutations. (A) The total number of occurrences for variants to be observed on plus or minus strands is compared. Data points are colored based on the variation type. (B) Zoomed in scatter plot for variants with strand-specific frequency <500. (C) Sequencing data from the same sequencing run (PG090) are assessed for frequency of observed variants after the use of two different versions of base-calling algorithm. (D) Zoomed in scatter plot for variants with frequency <500 compared between two datasets generated using different base-caller algorithms

To study the possible nucleotide-specific biases, we quantified the frequency of all calls that predominantly occurred on one strand. Despite slight variation, substitutions were observed on both strands at a comparable rate. However, in each dataset, the majority of substituted bases were 'A's (59.2, 61.3 and 64.9%) and 'T's (28.1, 23.0 and 20.5%) that were predominantly substituted to 'G' and 'C', respectively (Fig. 5A). Insertions were primarily observed on the forward strand (94%, on average) while 'A' remained as the most affected base (77.7%, on average) across all samples (Fig. 5B and Supplementary Figs S9–S11).

We also observed a slight enrichment of deletions and duplications on the reverse strand that were more pronounced in PG109-02 (Fig. 5C and D). Consistently, the most affected base was 'A', which was mainly the result of under- or over-calling of 'A' homopolymers. We used TSSV to report a list of most occurring variants across different samples. A single 'A' nucleotide insertion at cycle 52 was by far the most predominant variant that occurred exclusively on the forward strand (Fig. 6A). In fact, irrespective of co-occurrence of this insertion with any other variants, the new observed sequence remains strand specific (Fig. 6A). This cannot be explained from a biological standpoint and can only arise from a sequencing error. Moreover, we did not observe any variation after sequencing the same library with Sanger sequencing (Fig. 6B), ruling out the possibility of artifacts introduced by sample preparation and PCR amplification.

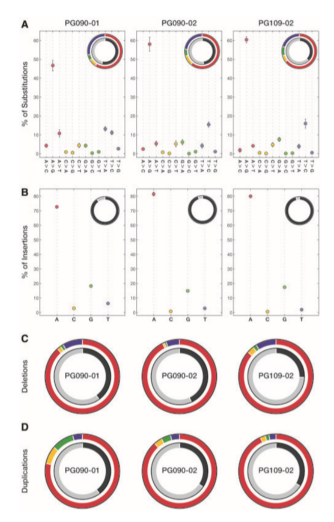


Fig. 5. Strand-specific and nucleotide-dependent variation patterns in targeted sequencing data. (A) Breakdown of substitution type frequencies across all samples. Pie charts depict substitution events per nucleotide (outer circle) and strand (inner circle) for each dataset. The outer pie charts illustrate the proportion of substitutions based on the preferred nucleotide to which substitutions are made. A, C, G and T nucleotides are reflected in red, yellow, green and blue colors, respectively. Black and gray colors represent the plus and minus strands, respectively (B) Breakdown of insertion frequencies per nucleotide across all samples. Pie charts represent the proportion of insertion events per nucleotide (outer circle) and strand (inner circle) for each dataset. (C) The fraction of deleted bases is shown in pie charts based on the nucleotide (outer circle) and strand (inner circle) for each dataset. (D) The amount of duplications per nucleotide (outer circle) and strand (inner circle) is presented for three datasets used in this study.

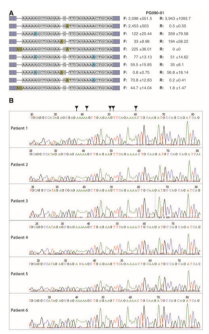


Fig. 6. Most abundant sequences and validation by Sanger sequencing. (A) A list of most abundant sequences across different samples. The observed occurrence of each new allele is quantified per strand along with corresponding standard deviation. (B) The result of Sanger sequencing of the target locus is presented per sample. Arrows mark the location in which mutations are reported based on the PGM Ion Torrent sequencing data.

Comparative analysis of TSSV performance

To our knowledge, lobSTR (Gymrek et al., 2012), STRait Razor (Warshauer et al., 2013) and RepeatSeq (Highnam et al., 2013) are currently the most recent and frequently used STR profiling tools. STRait Razor has limited functionality and only provides an estimated copy number of major STR units. Therefore, we could only compare the performance of TSSV only with lobSTR and RepeatSeq. As lobSTR relies on alignment of sequencing reads to the predefined and indexed STR reference sequences, lobSTR outperformed TSSV in recognizing partial reads, containing only one of the two flanking sites required by TSSV (Supplementary Table S21). Concordantly, lobSTR accepted 1288 reads for the D3S1357 STR locus that were not reported by TSSV. However, TSSV performed significantly better on more complex STR loci (Supplementary Tables S21–S22). Across all four datasets, TSSV identified on average 2471 and 2353 reads in excess of what was recognized by lobSTR for the D13S317 and TH01 STR loci, respectively. This difference is mainly derived from increasingly problematic alignments in lobSTR that is also reflected in inaccurate estimation of STR copy number for TH01 and D13S317 markers in pure samples (Supplementary

Table S22). In addition, lobSTR does not provide information on allelic STR structure, as it only reports the copy number of the major and uninterrupted STR unit and ignores the information from other variable elements or variants outside the STR itself. Consequently, lobSTR failed to accurately detect the presence of mixed simples even in cases in which samples were mixed 50:50 (Supplementary Table S22). Although the information on strand specificity of the aligned reads is present in the alignment file, unlike TSSV, lobSTR does not provide the frequency in which each STR structure is observed. This is an important measure to detect inconsistencies and to rule out potential artifacts. RepeatSeg requires aligned data and uses predefined regions to characterize observed STR alleles. Thus, reads were mapped to the reference genome (hg19) using GS mapper, specifically designed for 454 sequencing data. RepeatSeq reported results for only one STR locus (D8S1179), despite sufficient coverage for a number of STR loci in the BAM file. After manipulating the region descriptions, we could not improve the efficiency of RepeatSeg in identifying the targeted STR loci. Thus, the result of RepeatSeq could not be used for a conclusive comparison with TSSV.

Discussion

Iln the past decade, advances in sequencing technologies as well as computational analysis tools have enabled the study of genomic variations to dissect the mechanisms by which they exert their function in the case of human diseases, evolution and other complex traits. Despite this unprecedented progress, structural variations and repetitive DNA sequences (such as STRs) or coupling of de novo mutations present major obstacles for accurate and reliable allelic analysis (Alkan et al., 2011; Gymrek et al., 2012; Kidd et al., 2008; Treangen and Salzberg, 2012; Weischenfeldt et al., 2013). In particular, most computational tools are not ideal to identify STRs because of biases introduced during alignment as well as strong reliance of algorithms on coverage depth or the presence of split-reads. Here, we present a method (TSSV) that provides a highresolution map of allele-specific genomic variants within targeted loci of interest. Our approach does not rely on the use of a complete reference sequence to reliably profile highly polymorphic sequences (such as STRs) or uncharacterized variants at a singlenucleotide resolution. However, it does require two unique flanking sequences that harbor the region of interest to identify supporting reads. We assess the performance of TSSV on profiling known allelic STR structures across pure samples from a single individual as well as mixed samples with variable abundance. Of I 6 allelic STR structures that were targeted for sequencing, six STR loci were sufficiently covered so that the associated allelic STR structures could be reliably resolved. The strong imbalance between yield of STR markers is because of the assay (designed and optimized for fluorescent dye-based fragment analysis of STR loci) used for preparing the sequencing library. We show that sensitivity of TSSV in determining allelic STR structures exceeds mixtures with only 10% representation of minor alleles and more complex mixtures in which a single STR allele is shared. The lower boundary of detecting minor allele frequencies is subject to experimental design and the complexity of the targeted locus that may result in variable rate of slippage and background noise. Our detailed analysis of three STR loci provides significant insights into forensic DNA fingerprinting of mixed samples while it confirms the feasibility of TSSV to profile causal allelic expansion of triplet, tetranucleotide or more complex repeat structures in variety of human disorders (Brook et al., 1992; Dere et al., 2004; Kremer et al., 1991; Mahadevan et al., 1992; Mirkin, 2007; Pearson et al., 2002; Verkerk et al., 1991).

Second, we sought to profile and annotate the full spectrum of de novo mutations introduced by TALENs that specifically target intron 52 of hDMD in mouse ES cells. The applicability of designed TALENs to introduce mutations in a targeted locus largely depends on its efficacy because this method lacks a selection procedure. Detected TALEN-specific editing events were almost exclusively insertions and deletions that fit the expected mutation profile of TALENs (Cermak et al., 2011). Although it has recently been reported that TALENs induce insertions at a much lower frequency than deletions (Kim et al., 2013), we have observed an extremely balanced rate of insertion and deletion events (37.26% versus 37.20%, respectively), Nevertheless, TALENsinduced deletions tend to affect more bases than insertions. We show that TSSV can resolve difficult-to-call editing events that affect the length of homopolymers based on the variant frequency in TALEN-treated ES cells versus controls. Moreover, the result of TSSV analysis of TALEN-treated and control ES cells suggests that observed de novo structural variants are predominantly caused by initiation of a double-strand break that is repaired by non-homologous end-joining mechanism and are not the result of sequencing errors. Notably, the Ion Torrent variant caller failed to identify any of the observed variants because of their complexity, and therefore does not provide any information on de novo allelic structures that were introduced.

As laboratories begin to generate deep coverage sequencing data to identify low frequent mutations (i.e. cancer genomics), the robustness and accuracy of NGS technology and library preparation methods has become vital (Costello et al., 2013). After running TSSV on a third dataset to identify potential causal mutations in samples from five DMD patients and one female carrier, we observed numerous systematic errors introduced by the Ion Torrent PGM sequencer or the base-calling algorithms. The number of sequencing reads that support the presence of a new allele was in excess of 45% while no mutation was found after Sanger sequencing of the same libraries. Moreover, the amount of allelic discordant reads were unexpected and could not be biologically explained as five out of six samples were derived from male

patients who are expected to have only one copy of the X-chromosome. Across all samples, the majority of detected variations were single nucleotide insertions (~62%), excluding duplications, that were mostly the result of a single 'A' insertion (78%). Surprisingly, insertions were predominantly specific to the plus strand (94%) that can be the result of flow order in specific sequence contexts. Although the second base-caller improved the deletions and duplications rates that were derived from over- or under-calling of homopolymers, the insertion rates remained unchanged. We further observed a preference for erroneous substitution events that were more pronounced in the second base-caller. However, we were unable to identify motifs that may be associated with observed biases. We argue that the result of TSSV analysis and its ability to provide a high-resolution map of variants ever more highlights the importance of robust and vigorous assessment of downstream analysis as we generate volumes of sequencing data to identify rare mutations and in the advent of NGS in clinical diagnosis.

To demonstrate the added value of TSSV over mainstream STR profiling tools, we ran lobSTR (Gymrek et al., 2012) and RepeatSeq (Highnam et al., 2013) on four samples used for resolving allelic STR structures. Because RepeatSeq hardly reported any STR markers, the performance of TSSV could only be compared with that of lobSTR. We show that TSSV robustly and accurately resolved allelic STR structures with differing complexity. TSSV outperformed lobSTR in reporting the accurate copy number of major STR unit while it provides additional information on allelic STR structures and their strand-specific frequencies. Notably, TSSV excelled in resolving complex mixtures, whereas lobSTR failed to differentiate STR structures associated with different samples, and therefore produced unreliable and inaccurate estimations. Although lobSTR performs well on genotyping diploid samples, there is a clear need for tools to resolve mixtures with differing level of complexity and abundance.

Currently, the major limitation of TSSV is the sequencing read length because the detectable allelic structures are restricted to those that can entirely be covered by a single read. Thus, we envision that the immediate developmental outlook for TSSV can be the inference of allelic locus structure by local assembly of partial reads (reads with only one recognizable flanking region) combined with the comparative analysis of coverage of targeted loci and flanking regions. Furthermore, the promise of novel sequencing technologies (such as Pacific Biosciences RS II), and therefore significant increase in read length will aid the study of larger structural variations.

Advances in sequencing technologies and computational analysis algorithms in unraveling genetic variations from SNPs and indels to CNVs (Chen et al., 2009; DePristo et al., 2011; Goya et al., 2010; Koboldt et al., 2009; McKenna et al., 2010;

Ye et al., 2009) have facilitated the study of experimental data on an unprecedented scale to better understand the functional consequences of genetic variations. TSSV complements the existing tools by aiding the study of unknown, uncharacterized or highly polymorphic and repetitive short structural variations that can be used in a wide range of applications, from personal genomics to forensic analysis and clinical diagnostics.

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Supplementary materials

Methodology (extended)

We developed a method to characterise short structural variations (TSSV) which is made available online. In this section, we describe the functionality and design of this program. Calibration of the algorithm and the output is done with optional command line arguments. TSSV can be installed via pip install tssv command.

Input – Our method expects two input files: one file containing sequencing data in FASTA format and one file containing the library description. The format of this description is shown in Table S1. The last column of the description is compiled into a regular expression. This regular expression is used to distinguish between known and unknown alleles.

Marker Alignment – Each pair of flanking markers is aligned to each read by using semi-global pairwise alignment, a modified version of the Smith-Waterman algorithm. In this adaptation, the alignment matrix is initialised with penalties for the aligned sequence, but not for the reference sequence. By using this approach, we can use the alignment matrix for the calculation of the edit distance between the aligned sequence and all substrings of the reference sequence. Finally, we use the alignment matrix to select the rightmost alignment with a minimum edit distance. To guarantee that this method is symmetrical with regard to the reverse complement, we align the reverse complement of the right marker to the reverse complement of the reference sequence.

Allele Identification – If a marker pair can be aligned to either the forward or reverse complement of the reference sequence, we can select the area of interest by extracting the sequence between the alignment coordinates and by converting it into the forward orientation. This area of interest can then be matched to the regular expression of that marker pair. Depending on the match, we either classify the area of interest as a known or new allele.

Output – The output of the analysis consists of an overview report that contains general statistics (such as total number of reads, number of matched pairs, number of unique newly identified alleles, etc.), an overview of the marker pair alignment, and per marker a detailed list of identified alleles (both expected and new alleles). If an output directory is selected, a folder is created to store the marker table and, per marker, a subfolder containing the new alleles and split FASTA files for supporting reads.

Table SI - Full description of settings and generated files

Command Line (Optional Arguments
-h	Show a description of the usage
-m	Number of mismatches per nucleotide (default: 0.08)
-r	Name of the report file (default: stdout)
-d	Name of the output directory
-a	Minimum count per allele to be reported (default: 0)
Library Definitio	n
Column 1	Name of the marker pair
Column 2	Sequence of the left flanking marker
Column 3	Sequence of the right flanking marker
Column 4	Description of the expected alleles
General Output	Files per Marker
known.fa	Reads classified as known allele
new.fa	Reads classified as new alleles
unknown.fa	Reads classified as unrecognized
knownalleles.csv	Table of known alleles
newallels.csv	Table of new alleles
Marker Overvie	W
name	Name of the marker pair
fPaired	Number of pair matches in forward orientation
rPaired	Number of pair matches in reverse orientation
f1	Number of left marker matches in forward orientation
r1	Number of left marker matches in reverse orientation
f2	Number of left marker matches in forward orientation
r2	Number of left marker matches in reverse orientation
Allele Overview	
allele	Sequence of the area of interest
total	Number of times the allele was found
forward	Number of times the allele was found in forward orientation
reverse	Number of times the allele was found in reverse orientation

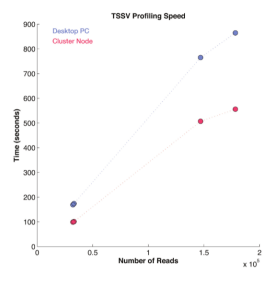


Figure S1 – The speed of TSSV in characterizing known and novel alleles. Four datasets with different number of reads were profiled for 16 STR loci. The analysis was performed on a desktop PC (Intel Core i7 860, 2.80GHz) and a cluster node (Intel Xeon E5-2660, 2.20GHz). ost abundant sequences and validation by Sanger sequencing. (A) A list of most abundant sequences across different samples. The observed

Table S2 - Description of primers used for targeting loci of interest.

Name	Forward Primer	Reverse Primer	
hDMD-int52	gggaaagtgaaagagtaaccaga	ng	ACACACCATCTAATGCTTATGAGG
hDMD-exn19	ccttgtattgaattactcatc		cctaagaagattatctaaatcaactcgt
hDMD-exn21	acgtgttacttactttccatact		caagttagccattttaggctt

Table S3 - Basic statistics of STR datasets sequenced on 454/Roche.

		Reference sample 1	Reference sample 2	Mixture 1 (50%/50%)	Mixure 2 (90%/10%)
Total seque	ences	146968	178282	33127	32201
Sequences	recognized as known alleles	75837	78228	16954	16425
Sequences	recognized as new alleles	15622	17887	4660	3720
Sequences	only recognized for the beginning	35154	106253	16204	16526
Sequences	only recognized for the end	75110	69957	9990	8782
Unknown s	equences	11478	22153	2108	2260
Sequences	s Recognized / locus				
D13S317	13q31.1	15361	11473	3069	3712
D21S11	21q21.1	13290	18477	2544	3218
Amel	X/Y	159	114	37	35
D3S1358	3p21.31	19822	25590	5507	4893
D16S539	16q24.1	82	97	26	19
TPOX	2p25.3	29	80	27	13
vWA	12p13.31	189	215	34	32
D7S820	7q21.11	10998	7339	2626	1975
FGA	4q28	14	28	9	6
TH01	11p15.5	19657	17204	4664	3956
D8S1179	8q24.13	66	80	23	18
CSF1P0	5q33.1	81	291	21	27
D18S51	18q21.33	128	247	38	46
D5S818	5q23.2	11523	14628	2983	2184
PentaD	21q22.3	25	36	6	5
PentaE	15q26.2	35	216	0	6

Table S4 - Sequence numbers and allele structure of D3S1358 Short Tandem Repeat.

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3	Mixture interpretation
Sample 1	12	14	2	16	TCTA(1)	TCTG(1)	TCTA(10)	
	13	49	17	66	TCTA(1)	TCTG(1)	TCTA(11)	
	13	14	16	30	TCTA(1)	TCTG(2)	TCTA(10)	
	14	365	244	609	TCTA(1)	TCTG(1)	TCTA(12)	
	14	403	245	648	TCTA(1)	TCTG(2)	TCTA(11)	
	15	4681	3111	7792	TCTA(1)	TCTG(1)	TCTA(13)	
	15	4774	3211	7985	TCTA(1)	TCTG(2)	TCTA(12)	
	15	12	8	20	TCTA(1)	TCTG(3)	TCTA(11)	
	16	38	19	57	TCTA(1)	TCTG(1)	TCTA(14)	
	16	38	21	59	TCTA(1)	TCTG(2)	TCTA(13)	
Sample 2	13	7	2	9	TCTA(1)	TCTG(2)	TCTA(10)	
	13	9	1	10	TCTA(1)	TCTG(3)	TCTA(9)	
	14	33	13	46	TCTA(1)	TCTG(2)	TCTA(11)	
	14	27	18	45	TCTA(1)	TCTG(3)	TCTA(10)	
	15	592	241	833	TCTA(1)	TCTG(2)	TCTA(12)	
	15	496	214	710	TCTA(1)	TCTG(3)	TCTA(11)	
	16	27	6	33	TCTA(1)	TCTG(1)	TCTA(14)	
	16	7484	2602	10086	TCTA(1)	TCTG(2)	TCTA(13)	
	16	6898	2442	9340	TCTA(1)	TCTG(3)	TCTA(12)	
	17	70	29	99	TCTA(1)	TCTG(2)	TCTA(14)	
	17	35	25	60	TCTA(1)	TCTG(3)	TCTA(13)	
50:50 Mixture	13	3	7	10	TCTA(1)	TCTG(1)	TCTA(11)	<1% of highest allele
	13	4	9	13	TCTA(1)	TCTG(2)	TCTA(10)	-2 stutter Al 15b
	14	52	32	84	TCTA(1)	TCTG(1)	TCTA(12)	-1 stutter Al 15a
	14	51	38	89	TCTA(1)	TCTG(2)	TCTA(11)	-1 stutter Al 16b
	14	23	4	27	TCTA(1)	TCTG(3)	TCTA(10)	-2 stutter Al 16c
	15	612	520	1132	TCTA(1)	TCTG(1)	TCTA(13)	Allele 15a
	15	536	494	1030	TCTA(1)	TCTG(2)	TCTA(12)	Allele 15b
	15	88	63	151	TCTA(1)	TCTG(3)	TCTA(11)	-1 stutter Al 16c
	16	9	11	20	TCTA(1)	TCTG(1)	TCTA(14)	+1 stutter Al 15a
	16	666	439	1105	TCTA(1)	TCTG(2)	TCTA(13)	Allele 16b
	16	650	491	1141	TCTA(1)	TCTG(3)	TCTA(12)	Allele 16c
	17	14	1	15	TCTA(1)	TCTG(2)	TCTA(14)	+1 stutter Al 16b
90:10 Mixture	13	9	2	11	TCTA(1)	TCTG(1)	TCTA(11)	<1% of highest allele
	14	117	84	201	TCTA(1)	TCTG(1)	TCTA(12)	-1 stutter Al 15a
	14	76	45	121	TCTA(1)	TCTG(2)	TCTA(11)	-1 stutter Al 15b
	15	924	808	1732	TCTA(1)	TCTG(1)	TCTA(13)	Allele 15a
	15	978	566	1544	TCTA(1)	TCTG(2)	TCTA(12)	Allele 15b
	15	70	16	86	TCTA(1)	TCTG(3)	TCTA(11)	-1 stutter Al 16c
	16	23	3	26	TCTA(1)	TCTG(1)	TCTA(14)	+1 stutter Al 15a
	16	110	65	175	TCTA(1)	TCTG(2)	TCTA(13)	Allele 16b
	16	177	102	279	TCTA(1)	TCTG(3)	TCTA(12)	Allele 16c

Table S5 - Sequence numbers and allele structure of D13S317 Short Tandem Repeat.

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3	Mixture interpretatio
Sample 1	11	5	6	11	ATCT(9)	ATCA(2)		
	12	210	192	402	ATCT(10)	ATCA(2)		
	12	7	10	17	ATCT(9)	ATCA(3)		
	13	3061	3039	6100	ATCT(11)	ATCA(2)		
	13	249	222	471	ATCT(10)	ATCA(3)		
	14	59	59	118	ATCT(12)	ATCA(2)		
	14	3218	2864	6082	ATCT(11)	ATCA(3)		
	14	53	49	102	ATCT(10)	ATCA(4)		
	15	14	30	44	ATCT(12)	ATCA(3)		
Sample 2	12	7	13	20	ATCT(10)	ATCA(2)		
	13	182	160	342	ATCT(11)	ATCA(2)		
	13	8	10	18	ATCT(10)	ATCA(3)		
	14	2495	1982	4477	ATCT(12)	ATCA(2)		
	14	151	181	332	ATCT(11)	ATCA(3)		
	15	47	42	89	ATCT(13)	ATCA(2)		
	15	2409	2014	4423	ATCT(12)	ATCA(3)		
	15	22	29	51	ATCT(11)	ATCA(4)		
	16	24	21	45	ATCT(13)	ATCA(3)		
50:50 Mixture	12	35	38	73	ATCT(10)	ATCA(2)		-1 stutter Al 13b
	13	525	467	992	ATCT(11)	ATCA(2)		Allele 13b
	13	30	32	62	ATCT(10)	ATCA(3)		-1 stutter Al 14c
	14	191	235	426	ATCT(12)	ATCA(2)		Allele 14b
	14	327	324	651	ATCT(11)	ATCA(3)		Allele 14c
	14	6	1	7	ATCT(10)	ATCA(4)		< 1% of highest allele
	15	4	4	8	ATCT(13)	ATCA(2)		< 1% of highest allele
	15	160	229	389	ATCT(12)	ATCA(3)		Allele 15c
	15	2	4	6	ATCT(11)	ATCA(4)		+1 stutter Al14c
90:10 Mixture	12	104	58	162	ATCT(10)	ATCA(2)		-1 stutter Al 13b
	13	751	589	1340	ATCT(11)	ATCA(2)		Allele 13b
	13	48	43	91	ATCT(10)	ATCA(3)		-1 stutter Al 14c
	14	66	41	107	ATCT(12)	ATCA(2)		Allele 14b
	14	691	546	1237	ATCT(11)	ATCA(3)		Allele 14c
	14	4	5	9	ATCT(10)	ATCA(4)		<1% of highest allele
	15	28	71	99	ATCT(12)	ATCA(3)		Allele 15c

Table S6 - Sequence numbers and allele structure of TH01 Short Tandem Repeat.

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3	Mixture Interpretation
Sample 1	8	233	112	345	CATT(8)			
	8.3	3	22	25	CATT(4)	CAT(1)	CATT(4)	
	8.3	91	34	125	CATT(3)	CAT(1)	CATT(5)	
	9	5280	2526	7806	CATT(9)			
	9.3	4626	2417	7043	CATT(3)	CAT(1)	CATT(6)	
	10	24	12	36	CATT(10)			
Sample 2	6	114	34	148	CATT(6)	•		
	6.3	8	4	12	CATT(5)	CAT(1)	CATT(1)	
	6.3	18	2	20	CATT(4)	CAT(1)	CATT(2)	
	7	5435	1412	6847	CATT(7)			
	8.3	70	24	94	CATT(3)	CAT(1)	CATT(5)	
	9.3	5789	1507	7296	CATT(3)	CAT(1)	CATT(6)	
50:50 Mixture	6	20	23	43	CATT(6)	•		-1 stutter Al 7
	7	514	494	1008	CATT(7)			Allele 7
	8	5	14	19	CATT(8)			-1 stutter Al 9
	8.3	9	4	13	CATT(3)	CAT(1)	CATT(5)	-1 Stutter Al 9.3
	9	239	235	474	CATT(9)			Allele 9
	9.3	671	591	1262	CATT(3)	CAT(1)	CATT(6)	Allele 9.3
90:10 Mixture	7	95	60	155	CATT(7)	-		Allele 7
	8	29	23	52	CATT(8)			-1 stutter Al 8
	8.3	0	11	11	CATT(4)	CAT(1)	CATT(4)	<1% of highest allele
	8.3	16	7	23	CATT(3)	CAT(1)	CATT(5)	-1 stutter Al 9.3
	9	873	404	1277	CATT(9)			Allele 9
	9.3	1044	584	1628	CATT(3)	CAT(1)	CATT(6)	Allele 9.3

Table S7 - Sequence numbers and allele structure of D21S11 Short Tandem Repeat (6 most frequent alleles for every sample.)

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3	Repeat Motif 4	Repeat Motif 5	Repeat Motif 6	Repeat Motif 7	Repeat Motif 8	Repeat Motif 9
Sample 1	31	1593	2968	4561	TCTA(6)	TCTG(5)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(12)
	30	1527	2443	3970	TCTA(4)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(12)
	30	151	261	412	TCTA(6)	TCTG(5)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(11)
	29	167	169	336	TCTA(4)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(11)
	30	28	102	130	TCTA(5)	TCTG(5)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(12)
	31	36	80	116	TCTA(4)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(13)
Sample 2	29	1182	4645	5827	TCTA(4)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(11)
	31	1275	3041	4316	TCTA(5)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(12)
	30	200	286	486	TCTA(5)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(11)
	28	65	252	317	TCTA(4)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(10)
	30	130	158	288	TCTA(4)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(12)
50:50 Mixture	29	256	298	554	TCTA(4)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(11)
	30	209	264	473	TCTA(4)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(12)
	31	156	186	342	TCTA(6)	TCTG(5)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(12)
	31	115	177	292	TCTA(5)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(12)
	30	17	15	32	TCTA(5)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(11)
	28	14	17	31	TCTA(4)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(10)
90:10 Mixture	31	421	634	1055	TCTA(6)	TCTG(5)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(12)
	30	377	550	927	TCTA(4)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(12)
	29	40	58	98	TCTA(4)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(11)
	30	34	47	81	TCTA(6)	TCTG(5)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(11)
	31	24	46	70	TCTA(5)	TCTG(6)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(12)
	32	37	10	47	TCTA(6)	TCTG(5)	TCTA(3)	TATCTA(1)	TCTA(2)	TCA(1)	TCTA(2)	TCCATA(1)	TCTA(13)

Table S8 - Sequence numbers and allele structure of D16S539 Short Tandem Repeat (6 most frequent alleles for every sample.)

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3
Sample 1	12	12	57	69	GATA(12)		
	11	1	1	2	GATA(11)		
Sample 2	13	3	34	37	GATA(13)		
	12	0	27	27	GATA(12)		
50:50 Mixture	12	5	8	13	GATA(12)		
	13	0	8	8	GATA(13)		
	11	1	0	1	GATA(11)		
90:10 Mixture	12	7	8	15	GATA(12)		
	11	0	1	1	GATA(11)		

Table S9 - Sequence numbers and allele structure of TPOX Short Tandem Repeat (6 most frequent alleles for every sample.)

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3
Sample 1	11	3	8	11	AATG(11)		
	9	6	2	8	AATG(9)		
	10	0	1	1	AATG(10)		
Sample 2	11	19	7	26	AATG(11)		
	8	24	2	26	AATG(8)		
	10	1	2	3	AATG(10)		
50:50 Mixture	11	1	2	3	AATG(11)		
	10	1	1	2	AATG(10)		
	9	2	0	2	AATG(9)		
	8	1	0	1	AATG(8)		
90:10 Mixture	11	3	4	7	AATG(11)		
	9	4	0	4	AATG(9)		
	8	0	1	1	AATG(8)		

Table S10 - Sequence numbers and allele structure of vWA Short Tandem Repeat (6 most frequent alleles for every sample.)

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3	Repeat Motif 4	Repeat Motif 5
Sample 1	16	67	86	153	TCTA(1)	TCTG(4)	TCTA(11)		
	15	3	5	8	TCTA(1)	TCTG(4)	TCTA(10)		
	16	0	5	5	TCTA(1)	TCTG(5)	TCTA(9)	TCTG(1)	
Sample 2	17	34	50	84	TCTA(1)	TCTG(4)	TCTA(12)		
	16	39	42	81	TCTA(1)	TCTG(4)	TCTA(11)		
	15	5	1	6	TCTA(1)	TCTG(4)	TCTA(10)		
	15	0	1	1	TCTA(1)	TCTG(3)	TCTA(11)		
	14	0	1	1	TCTA(1)	TCTG(4)	TCTA(9)		
	17	1	0	1	TCTA(1)	TCTG(4)	TCTA(10)	TCCA(1)	TCTA(1)
50:50 Mixture	16	14	5	19	TCTA(1)	TCTG(4)	TCTA(11)		
	17	3	6	9	TCTA(1)	TCTG(4)	TCTA(12)		
	17	0	1	1	TCTA(2)	TCTG(3)	TCTA(12)		
90:10 Mixture	16	12	15	27	TCTA(1)	TCTG(4)	TCTA(11)		
	16	1	1	2	TCTA(1)	TCTG(4)	TCTA(10)	TCCA(1)	
	16	1	0	1	TCTA(1)	TCTG(5)	TCTA(10)		

Table SII - Sequence numbers and allele structure of D7S820 Short Tandem Repeat (6 most frequent alleles for every sample.)

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3
Sample 1	7	3268	3037	6305	GATA(7)		
	12	1535	1847	3382	GATA(12)		
	11	130	102	232	GATA(11)		
	6	75	69	144	GATA(6)		
Sample 2	9	2927	3561	6488	GATA(9)		
	8	140	170	310	GATA(8)		
50:50 Mixture	7	584	454	1038	GATA(7)		
	9	326	322	648	GATA(9)		
	12	311	289	600	GATA(12)		
	8	28	22	50	GATA(8)		
	11	24	19	43	GATA(11)		
	6	11	17	28	GATA(6)		
90:10 Mixture	7	548	474	1022	GATA(7)		
	12	284	333	617	GATA(12)		
	9	28	42	70	GATA(9)		
	11	15	29	44	GATA(11)		

Table S12 - Sequence numbers and allele structure of FGA Short Tandem Repeat (6 most frequent alleles for every sample.)

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3	Repeat Motif 4	Repeat Motif 5	Repeat Motif 6	Repeat Motif 7
Sample 1	21	7	0	7	TTTC(3)	TTTTTCT(1)	CTTT(13)	CTCC(1)	TTCC(2)		
	17	1	0	1	TTTC(3)	TTTTTCT(1)	CTTT(9)	CTCC(1)	TTCC(2)		
Sample 2	20	5	0	5	TTTC(3)	TTTTTCT(1)	CTTT(12)	CTCC(1)	TTCC(2)		
	24	5	0	5	TTTC(3)	TTTTTCT(1)	CTTT(16)	CTCC(1)	TTCC(2)		
	19	1	0	1	TTTC(3)	TTTTTCT(1)	CTTT(11)	CTCC(1)	TTCC(2)		
	21	1	0	1	TTTC(3)	TTTTTCT(1)	CTTT(13)	CTCC(1)	TTCC(2)		
50:50 Mixture	17	2	0	2	TTTC(3)	TTTTTCT(1)	CTTT(9)	CTCC(1)	TTCC(2)		
	20	1	0	1	TTTC(3)	TTTTTCT(1)	CTTT(8)	CCTT(1)	CTTT(3)	CTCC(1)	TTCC(2)
	20	1	0	1	TTTC(3)	TTTTTCT(1)	CTTT(12)	CTCC(1)	TTCC(2)		
90:10 Mixture	21	2	0	2	TTTC(3)	TTTTTCT(1)	CTTT(13)	CTCC(1)	TTCC(2)		

Table S13 - Sequence numbers and allele structure of D8S1179 Short Tandem Repeat (6 most frequent alleles for every sample.)

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3	Repeat Motif 4
Sample 1	10	12	13	25	TCTA(10)			
	13	18	3	21	TCTA(1)	TCTG(1)	TCTA(11)	
	12	2	0	2	TCTA(1)	TCTG(1)	TCTA(10)	
Sample 2	13	13	17	30	TCTA(13)			
	14	10	11	21	TCTA(14)			
	12	0	4	4	TCTA(12)			
	14	1	0	1	TCTA(3)	TCTG(1)	TCTA(10)	
	14	1	0	1	TCTG(1)	TCTA(13)		
50:50 Mixture	13	4	2	6	TCTA(1)	TCTG(1)	TCTA(11)	
	10	1	2	3	TCTA(10)			
	14	0	2	2	TCTA(14)			
	13	1	1	2	TCTA(13)			
90:10 Mixture	10	1	5	6	TCTA(10)			
	13	1	3	4	TCTA(1)	TCTG(1)	TCTA(11)	
	14	1	0	1	TCTA(14)			
	13.1	0	1	1	TCTA(1)	TCTG(1)	TCTA(10)	TCTAT(1)

Table S14 - Sequence numbers and allele structure of CSF1P0 Short Tandem Repeat (6 most frequent alleles for every sample.)

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3
Sample 1	10	7	26	33	AGAT(10)		
	13	6	10	16	AGAT(13)		
	12	1	1	2	AGAT(12)		
	9	0	1	1	AGAT(9)		
	8	1	0	1	AGAT(8)		
Sample 2	12	19	106	125	AGAT(12)		
	13	28	67	95	AGAT(13)		
	11	1	4	5	AGAT(11)		
	10	0	1	1	AGAT(10)		
50:50 Mixture	13	1	7	8	AGAT(13)		
	12	2	1	3	AGAT(12)		
	10	0	2	2	AGAT(10)		
	11	0	1	1	AGAT(11)		
90:10 Mixture	10	1	8	9	AGAT(10)		
	13	0	3	3	AGAT(13)		
	8	0	1	1	AGAT(8)		

Table S15 - Sequence numbers and allele structure of D18S51 Short Tandem Repeat (6 most frequent alleles for every sample.)

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3
Sample 1	14	0	35	35	AGAA(14)		
	15	0	19	19	AGAA(15)		
	13	0	1	1	AGAA(13)		
Sample 2	13	0	57	57	AGAA(13)		
	14	0	35	35	AGAA(14)		
	12	0	8	8	AGAA(12)		
	15	0	2	2	AGAA(15)		
50:50 Mixture	14	0	13	13	AGAA(14)		
	13	0	2	2	AGAA(13)		
	15	0	1	1	AGAA(15)		
90:10 Mixture	14	0	10	10	AGAA(14)		
	13	0	4	4	AGAA(13)		
	15	0	3	3	AGAA(15)		

Table S16 - Sequence numbers and allele structure of D5S818 Short Tandem Repeat (6 most frequent alleles for every sample.)

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3
Sample 1	14	2049	2825	4874	AGAT(11)	AGAG(1)	
	17	1626	2140	3766	AGAT(14)	AGAG(1)	
	16	118	241	359	AGAT(13)	AGAG(1)	
	13	106	152	258	AGAT(10)	AGAG(1)	
Sample 2	14	2118	4512	6630	AGAT(11)	AGAG(1)	
	15	1708	3351	5059	AGAT(12)	AGAG(1)	
	13	114	261	375	AGAT(10)	AGAG(1)	
50:50 Mixture	14	517	765	1282	AGAT(11)	AGAG(1)	
	15	258	313	571	AGAT(12)	AGAG(1)	
	17	156	169	325	AGAT(14)	AGAG(1)	
	13	33	38	71	AGAT(10)	AGAG(1)	
	16	8	36	44	AGAT(13)	AGAG(1)	
90:10 Mixture	14	340	529	869	AGAT(11)	AGAG(1)	
	17	282	354	636	AGAT(14)	AGAG(1)	
	16	24	56	80	AGAT(13)	AGAG(1)	
	15	27	32	59	AGAT(12)	AGAG(1)	
	13	19	39	58	AGAT(10)	AGAG(1)	

Table S17 - Sequence numbers and allele structure of Penta E Short Tandem Repeat (6 most frequent alleles for every sample.)

	Total # of repeats	Forward sequences	Reverse sequences	Total # of sequences	Repeat Motif 1	Repeat Motif 2	Repeat Motif 3
Sample 1	17	0	6	6	AAAGA(17)		
	12	0	2	2	AAAGA(12)		
Sample 2	7	0	139	139	AAAGA(7)		
90:10 Mixture	13	0	2	2	AAAGA(13)		

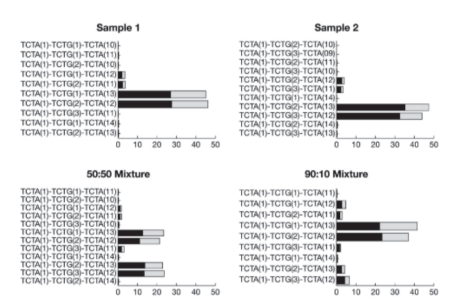


Figure S2 – The percentage of reads supporting detected allele-structure of D3S1358 Short Tandem Repeat. Black bars depict reads supporting the forward strand and grey bars correspond to the proportion of reads supporting the reverse complement.

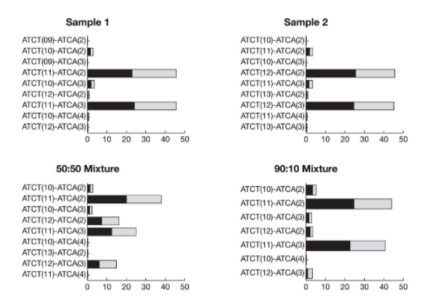


Figure S3 – The percentage of reads supporting detected allele-structure of D13S317 Short Tandem Repeat. Black bars depict reads supporting the forward strand and grey bars correspond to the proportion of reads supporting the reverse complement.

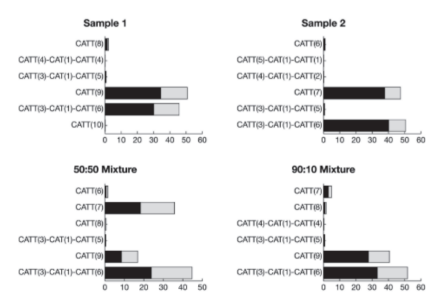


Figure S4 – The percentage of reads supporting detected allele-structures of THO I Short Tandem Repeat. Black bars depict reads supporting the forward strand and grey bars correspond to the proportion of reads supporting the reverse complement.

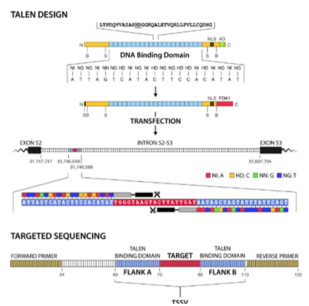


Figure S5 - Schematic representation of TALEN design and targeted resequencing TALEN-pairs were designed to specifically target intron 52 of the hDMD in mouse ES Cells. The binding sites of TALEN-pairs are 21bp long (blue). After successful transfection, TALENs initiate a double strand break within the target locus of 19bp (red). The 135bp fragment encompassing the entire targeted region was PCR-amplified, sequenced, and analysed using TSSV.

Table S18 – Target sequence of the TALENS targeting intron 52 of the hDMD gene.

	TAL1	TAL2
Nucleotide Sequence	ATTAGTCATACTTCCACATAT	AATAGCTAGTATTTATTCAGT
RVD sequence	NI NG NG NI NN NG HD NI NG NI HD NG NG HD HD NI HD NI NG NI NG	NI HD NG NN NI NI NG NI NI NI NG NI HD NG NI NN HD NG NI NG NG

Table S19 – Basic statistics on sequencing reads from TALEN-treated and control ES Cells.

 Table S19 - Basic statistics on sequencing reads from TALEN-treated and control ES Cells.

	# Reads	# Unique New Alleles	# Reads for New Alleles	# Reads with no Start	# Reads with no End	# Unrecognized Reads
TALEN-treated ES Cells	466,633	3,868	46,337	4,225	2,861	7,229
Control ES Cells	423,674	1,458	12,317	3,614	5,756	7,414

Table S20 – Basic statistics on sequencing reads from 5 male patients and a female carrier.

	# Reads	# Unique New Alleles	# Reads with no Start	# Reads with no End
PG090-01				
Sample 1	16,346	1,214	2,304	673
Sample 2	15,589	1,161	2,215	2,208
Sample 3	22,397	1,593	3,533	3,387
Sample 4	12,773	1,009	1,867	2,094
Sample 5	14,834	1,093	2,206	2,581
Sample 6	20,599	1,307	2,626	3,241
PG090-02				
Sample 1	14,337	498	1,536	620
Sample 2	13,420	511	1,251	2,095
Sample 3	18,881	594	2,004	3,098
Sample 4	10,995	421	1,040	2,004
Sample 5	12,769	429	1,266	2,483
Sample 6	18,488	571	1,599	3,237
PG109-02				
Sample 1	11,612	447	1,411	381
Sample 2	11,857	390	1,253	2,484
Sample 3	17,386	510	1,964	4,018
Sample 4	9,849	351	978	2,407
Sample 5	11,545	365	1,200	2,949
Sample 6	16,816	507	1,637	4,164

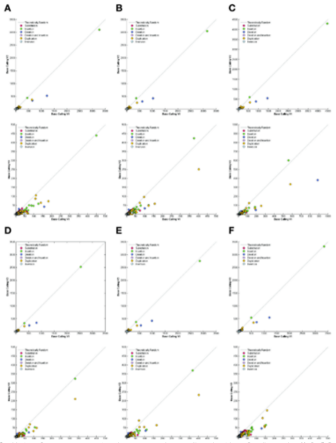


Figure S8 – Variant frequency comparison between two base-calling algorithms for all samples (A, B, C, D, E, F). For each panel, the first scatter plot shows all variants and the second zooms in to variants with frequency less than 500.

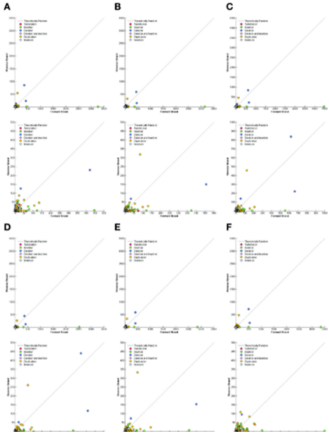


Figure 59 – Strand specificity of observed variants in dataset PG090-01 for all samples (A, B, C, D, E, F). For each panel, the first scatter plot shows all variants and the second zooms in to variants with frequency less than 500 on each strand.

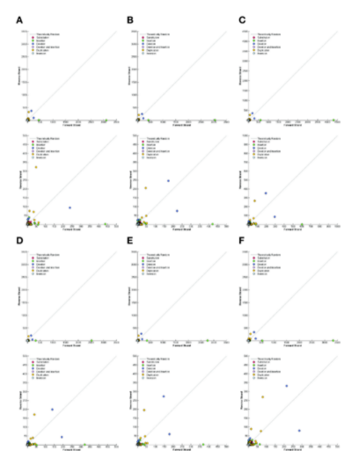


Figure \$10 — Strand specificity of observed variants in dataset PG090-02 for all samples (A, B, C, D, E, F). For each panel, the first scatter plot shows all variants and the second zooms in to variants with frequency less than 500 on each strand.

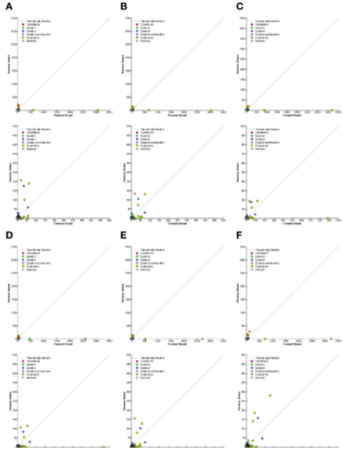


Figure S11 – Strand specificity of observed variants in dataset PG109-02 for all samples (A, B, C, D, E, F). For each panel, the first scatter plot shows all variants and the second zooms in to variants with frequency less than 500 on each strand.

Table S21 – General statistics on identification of informative reads for STR profiling of samples using lobSTR.

	Sample 1	Sample 2	50:50 Mixture	90:10 Mixture
Total number of reads	146,968	178,282	33,127	32,201
Number of aligned reads	114,122	127,327	25,339	24,379
Number of stitched reads	0	0	0	0
Number of single-end reads	114,122	127,327	25,339	24,379
Number of supporting end reads	0	0	0	0
Percent of partially aligned reads	29.03%	37.90%	30.42%	31.26%
Percent of reverse strand reads	53.07%	50.95%	53.34%	51.73%
Percent of non-unit allele reads	14.00%	18.71%	16.02%	51.73%

Table S22 – lobSTR performance in identification and characterization of allele specific STR structures.

							Supporting Reads		ads	Validation	
	Official Name	Chr. Position	STR Repeat	Ref. Copy Number	lobSTR Calls *	Coverage	Total	Allele 1	Allele2	Copy Number	Structure
Sample 1	D3S1358	3p21.31	TCTA	16	-4,-4	23,661	20,810	NA	NA	Yes	No
	D13S317	13q31.1	ATCT	11	0,0	14,536	13,228	NA	NA	Yes	No
	TH01	11p15.5	CATT	7	0,11	19,072	16,828	9,038	7,790	No	No
Sample 2	D3S1358	3p21.31	TCTA	16	0,0	21,535	18,668	NA	NA	Yes	No
	D13S317	13q31.1	ATCT	11	0,5	10,588	9,994	5,738	4,256	No	No
	TH01	11p15.5	CATT	7	11,0	16,802	15,353	7,973	7,380	Yes	No
50:50 Mixture	D3S1358	3p21.31	TCTA	16	-4,0	4,769	4,340	2,754	1,586	No	No
	D13S317	13q31.1	ATCT	11	0,0	2,913	2,227	NA	NA	No	No
	TH01	11p15.5	CATT	7	0,11	3,935	3,642	1,887	1,755	No	No
90:10 Mixture	D3S1358	3p21.31	TCTA	16	-4,-4	5,262	4,327	NA	NA	No	No
	D13S317	13q31.1	ATCT	11	0,0	3,264	2,822	NA	NA	No	No
	TH01	11p15.5	CATT	7	11,0	3,720	3,391	1,755	1,636	No	No

lobSTR calls column represents the number of nucleotide differences between the called alleles and the length of the reference STR (STR leng-