

# A multiplex ligation-dependent probe amplification-based next-generation sequencing approach for the detection of copy number variations in the human genome

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**Abstract.** The aim of the present study was to describe a multiplex ligation-dependent probe amplification (MLPA)-based next-generation sequencing (NGS) assay that exhibited a significantly higher efficiency in detecting copy number variations (CNVs) and known single-nucleotide variants, compared with traditional MLPA. MLPA polymerase chain reaction products were used to construct a library with indexed adapters, which was subsequently tested on an NGS platform, and the resulting data were analyzed by a series of analytical software. The reads from each probe reflected genetic variations in the target regions, and fragment differentiation was based on the specific base composition of the sequences, rather than fragment length, which was determined by capillary electrophoresis. The results of this approach were not only consistent with the MLPA results following capillary electrophoresis, but also coincided with the CNV results from the single-nucleotide polymorphism array chip. This method allowed high-throughput screening for the number of fragments and samples by integrating additional indices for detection. Furthermore, this technology precisely and accurately performed large-scale detection and quantification of DNA variations, thereby serving as an effective and sensitive

method for diagnosing genetic disorders caused by CNVs and known single-nucleotide variations. Notably, MLPA-NGS circumvents the problems associated with the inaccuracies of NGS in CNV detection due to the use of target sequence capture.

## Introduction

Deletions, duplications, or other genomic rearrangements, may result in dosage imbalance of gene(s), which leads to the loss or gain of genetic material (1). The mechanisms underlying copy number variant (CNV) formation are based on recombination and replication. Compared with single-nucleotide polymorphisms (SNPs), the *de novo* locus-specific mutation appearance rate for CNVs is significantly higher (2). CNVs may cause Mendelian or sporadic traits, or be associated with complex diseases, and the molecular mechanisms include gene dosage, disruption, fusion and position effects, among others (3). CNVs are highly significant in human disease and population diversity (4). Other common complicated neuropsychiatric disorders, such as autism and schizophrenia, are also affected by CNVs (5). Various genome analysis platforms may perform CNV analysis. The golden standards for CNV detection are array comparative genomic hybridization (aCGH) and SNP genotyping platforms (6). However, SNP array or aCGH are not sufficient for detecting smaller CNVs. Multiplex ligation-dependent probe amplification (MLPA) may make up for the shortcomings of these technologies (7).

MLPA is an accurate and reliable technique for identifying CNVs, including large and small deletions, as well as single-nucleotide aberrations, with several advantages over other detection methods (8-12). Compared with conventional assays, including Southern blotting, fluorescence *in situ* hybridization and Sanger sequencing, MLPA is a good alternative to array-based techniques and has high accuracy, specificity and efficiency. In addition, MLPA is cost-effective and has less technical complexity relative to array comparative genomic hybridization, which often requires further validation

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using other methods, such as quantitative polymerase chain reaction (qPCR) (13-22). More importantly, MLPA may overcome the limitations of CMA and SNP array to some extent. For example, to our knowledge, the CNVs of the CYP21A2 gene cannot be analyzed correctly by CMA, SNP array or NGS due to the presence of its pseudogene (23,24). However, MLPA easily solves this problem, using the P050-C1 CAH kit (MRC-Holland, Amsterdam, The Netherlands). At least 300 commercial probe sets are currently available for detecting relatively common genetic disorders, such as Duchenne muscular dystrophy and spinal muscular atrophy, as well as rare genetic conditions, such as antithrombin deficiency and Birt-Hogg-Dubé syndrome (7).

MLPA mainly involves the separation of amplification products by size, limiting the maximum number of target sequences that can be screened in parallel to ~50 (12). However, it does not meet the requirements for detecting genetic disorders that are caused by diverse DNA variations. Although MLPA has a higher throughput compared with qPCR, it is currently not suitable for the large-scale screening of target regions, although efforts are currently focused on improving throughput (11,12). CNV-plex is the most representative of the existing methods, but is limited by the number of fluorescent groups and fragment length, with 384 base pairs (bp) as the maximum fragment size that can be detected in one reaction (21-22,25-29). The use of additional fluorescent groups also introduces technical complications into the detection of CNVs, increasing the complexity of data analysis.

The rapid development of next-generation sequencing (NGS), which performs sequencing via synthetic processes, provides a sensitive and accurate tool for detecting known or unknown genomic variations, including CNVs (25-40). However, the statistical approaches to CNV identification are limited, although there are several auto-calculation software types for CNV detection that utilize data generated from whole-exome or whole-genome sequencing (32-42).

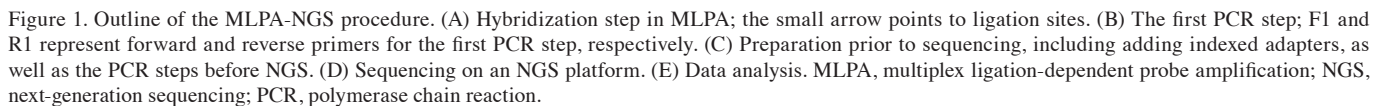
In the present study, a novel and robust method of MLPA-based NGS (MLPA-NGS) was introduced, which utilized MLPA products to construct a library that may be transferred into an NGS procedure to detect CNVs with improved accuracy and high throughput. MLPA PCR products with indexed adapters were tested on an NGS platform, and the resulting data were analyzed by using a series of analytical software, including FastQC, Burrows-Wheeler Alignment (BWA) tool and Genome Analysis Toolkit (GATK). The reads from each probe reflected genetic variations in the target regions, and fragment differentiation was based on the specific base composition of the sequences, rather than fragment length, which was determined by capillary electrophoresis. As such, the probe set may be designed to be within the same range of lengths, thereby allowing consistent detection efficiencies among reads. Furthermore, this approach ensures efficiency in amplification and purification of PCR products. This method also detects a significantly higher number of fragments compared with earlier methods, circumventing the 50 fragment detection limit per run. The novel approach of the current study also circumvented configuration of the stuffer sequence for different lengths in the probe. The synthesized probes did not involve the complexities associated with preparing long probes. Furthermore, the addition of indices to the adapters

for distinguishing between different samples allowed the assay to achieve high throughput detection of both sites and samples, while also ensuring quantitative detection of copy number with high accuracy. In summary, MLPA-NGS technology not only possessed all the advantages of MLPA, such as detecting the CNVs of CYP21A2 gene, but also overcame its limitations.

## Materials and methods

**Samples.** A total of 12 peripheral blood samples were collected from 12 unrelated subjects (age, 6-12 years) in the Children's Hospital of Shanghai affiliated to Shanghai Jiao Tong University (Shanghai, China) between June 2015 and May 2017, and included four 22q11.2 deletion syndrome (22q11DS; OMIM no. 611867) samples, five Duchenne muscular dystrophy (DMD; OMIM no. 310200) samples and three healthy controls. The four 22q11DS samples were collected from two female and two male patients. All DMD samples were collected from male patients. The three healthy controls were collected from two female and one male patient. In the present experiments, a male 22q11DS sample (termed PC sample) and a female negative control (termed NC sample) sample are described in detail. All samples were obtained with written informed consent and the study was approved by the Research Ethics Committee of the Children's Hospital of Shanghai. At each collection, a peripheral blood sample containing 3 ml was collected in BD Vacutainer PPT K2EDTA tubes (BD Diagnostics, Milan, Italy), genomic DNA was isolated from fresh blood samples using a DNeasy Blood and Tissue Extraction kit (Qiagen GmbH, Hilden, Germany), according to the manufacturer's protocol, and quantified using NanoDrop 1000 spectrophotometer (Thermo Fisher Scientific, Inc., Wilmington, DE, USA) and stored at -80°C until used.

**MLPA-based NGS protocol.** An MLPA-based NGS protocol was developed. MLPA was performed using the MLPA One-Tube MDP-v005 mix (MRC-Holland), according to the manufacturer's protocol. Briefly, 100 ng isolated DNA (aforementioned) was denatured at 98°C for 5 min, 3  $\mu$ l of the probe mix was added and heated at 95°C for 1 min, and subsequently hybridized overnight at 60°C. The samples were then treated with ligase-65 at 54°C for 15 min. The reactions were stopped by incubating at 98°C for 5 min. PCR amplification was performed with the specific SALSA FAM PCR primers (a 10  $\mu$ l mix of 7.5  $\mu$ l dH<sub>2</sub>O, 2  $\mu$ l SALSA PCR primer mix and 0.5  $\mu$ l SALSA polymerase) in the SALSA MLPA PCR kit (MRC-Holland). Amplification conditions were: Initial denaturation at 98°C for 5 min; followed by 35 cycles of 30 sec at 95°C; 30 sec at 60°C; 60 sec at 72°C; and final extension of 20 min at 72°C; hold at 15°C. As the labeled PCR product may interfere through ligation with NGS adapters (Fig. 1), the PCR products were then re-amplified by using universal primers without any label (forward, 5'-GGGTTCCTAAGGGT TGGA-3' and reverse, 5'-GCGCCAGCAAGATCCAATCTA GA-3'; amplification conditions were the same as above). PCR fragments were extracted and purified from a 2% agarose gel (stained with ethidium bromide and visualized under UV light) using a QIAquick Gel Extraction kit (Qiagen, Inc.), according to the manufacturer's protocol, and ligated to adapters.



**MLPA analysis.** DNA was isolated using a QIAamp DNA blood mini kit (Qiagen GmbH) according to the manufacturer's instructions. MLPA analysis of 22q11DS was performed using 52 pairs of probes in the SALSA MLPA probemix P064-C1 Mental Retardation-1 (MRC Holland). The regions targeted by P064-C1 probemix included 1p36, 15q11, 4p16, 16p13, 5p15, 17p13, 5q35, 17p11, 7p21, 20p12, 7q11, 22q11, 8q24, 22q13 and 11p13. The 52 MLPA probes resulted in amplification products between 130 and 483 nucleotides (nt) in length. MLPA was then performed according to the manufacturer's instructions. Briefly, 100 ng DNA was denatured at 98°C for 5 min, then 3  $\mu$ l of the probe mix was added, heated at 95°C for 1 min, and then hybridized overnight at 60°C. The samples were then treated with ligase-65 at 54°C for 15 min. The reactions were stopped by incubating at 98°C for 5 min. Finally,

PCR amplification was performed, following the protocol described in the aforementioned *MLPA-based NGS protocol*. The amplification products were run on an ABI PRISM 3500 Dx Genetic Analyzer (Applied Biosystems; Thermo Fisher Scientific, Inc.). The raw data from the Genetic Analyzer were analyzed with Coffalyser.Net (version 14; MRC-Holland). Briefly, the channel content of the probes was filled with P064-MR-1-C1-0912 (C1); following fragment analysis and comparative analysis with default settings, normalization for MLPA fragment data files was performed. Furthermore, the area encompassing the CNVs in the PC and NC samples relative to that of the reference was calculated using Coffalyser.Net. All quality measures and parameters were within a satisfactory range. All samples were normalized against multiple runs of the reference sample (inter-sample normalization), and all probes were adjusted to the reference probes within each sample (intra-sample normalization).

MLPA analysis of DMD was performed using P034-B2 DMD-1 & P035-B1 DMD-2 kits, which were also purchased from MRC-Holland. The analysis was performed using the same procedures as described above.

**Library construction and sequencing.** As MLPA PCR products are labeled with 5-carboxyfluorescein (FAM), which blocks ligation with adapters, the PCR products were once again amplified using the following primers: MLPA, forward 5'-GGGTTCCCTAAGGGTTGGA-3', reverse 5'-GCGCCA GCAAGATCCAATCTAGA-3'. The reaction was performed in system containing 2  $\mu$ l MLPA PCR product, 5  $\mu$ l 10X HS *Taq* buffer (Takara Biotechnology Co., Ltd. (Dalian, China), 4  $\mu$ l dNTPs (2.5 mM), 1  $\mu$ l MLPA forward primer (20 pM), 1  $\mu$ l MLPA reverse primer (20 pM), 2 U *HSTaq* (Takara Biotechnology Co., Ltd.), which was made up to a 50  $\mu$ l with the appropriate volume of ddH<sub>2</sub>O. PCR was performed using the same temperature profile as that of the MLPA-amplified reaction described above.

The PCR products were purified using Agencourt Ampure XP-PCR purification beads (cat. no. A63880; Beckman Coulter, Inc., Brea, CA, USA) with a Dynal magnetic bead stand (cat. no. 123-21D; Thermo Fisher Scientific, Inc.), according to the manufacturer's protocol. Briefly, 90  $\mu$ l Agencourt beads were mixed with 50  $\mu$ l PCR product, incubated at room temperature for 15 min, placed on a magnetic stand, washed twice with 80% (v/v) ethanol, separated from the ethanol and air-dried for 5 min. The beads were resuspended and incubated in 22  $\mu$ l ddH<sub>2</sub>O. Next, 20  $\mu$ l eluate (plus 4  $\mu$ l of the loading dye) was electrophoresed on a 2% agarose gel with a 100 bp DNA ladder for 2 h at 120V. DNA fragments that were within the size range of 40-550 bp were eluted with 25  $\mu$ l ddH<sub>2</sub>O by gel extraction using a QIAquick Gel Extraction kit (Qiagen, Inc., Valencia, CA, USA), according to the manufacturer's protocol. The products in the eluates were then subjected to end repair and A-tailing by a Kapa Library Preparation kit (Kapa Biosystems), according to the manufacturer's protocol. The Ligation Master mix and the indexed adapters were mixed and incubated at 2°C for 15 min, 35°C for 15 min and 72°C for 20 min, and then held at 4°C to produce paired-end libraries.

The post-ligation products were purified using Agencourt Ampure XP-PCR purification beads (cat. no. A63880; Beckman Coulter, Inc.) with a Dynal magnetic bead stand (cat.

no. 123-21D; Thermo Fisher Scientific, Inc.), according to the manufacturer's instructions. During library amplification, the reaction system was performed in a 0.2 ml tube. Similarly, library amplification purification was performed using Ampure XP beads, according to the manufacturer's instructions, in which the product was quantified using a Qubit DNA HS kit (Thermo Fisher Scientific, Inc.), and the test for fragment quality was conducted using an Agilent 2100 (Agilent, Inc., Santa Clara, CA, USA). The test results generated the expected fragments, which were then sequenced. The library preparations were sequenced on an Illumina HiSeq 2500 platform, and 150 bp paired-end reads were generated.

**Analysis of CNVs by SNP array chip.** HumanCytoSNP-12 BeadChip (Illumina, Inc.) was used to detect CNVs in DNA isolated from PC patient peripheral blood. The sample DNA was amplified, labeled and hybridized as previously described (44), and the data were acquired using Illumina's iScan scanning system. The frequency of the B allele and the log R ratio were analyzed with Illumina Karyo Studio (version 1.4.3). The log R ratio is the logged ratio of observed probe intensity to expected intensity; any deviations from zero in this metric are evidence of copy number alteration. The frequency of the B allele is the proportion of the hybridized sample that carries the B allele, as designated by the Infinium assay.

**Data analysis.** Quality control metrics for the NGS raw sequencing data (FASTQ files) were obtained using FastQC, version 0.10.1 ([www.bioinformatics.babraham.ac.uk/projects/fastqc](http://www.bioinformatics.babraham.ac.uk/projects/fastqc)). The sequences were then aligned to the human reference genome sequence (GRCh37/hg19) using BWA (version 0.6.2). GATK was used to compute read depths within the target region. For GATK analyses, default settings were used, except the mapping quality threshold (Q=5). The same normalization method treatment with the PC relative peak area was also performed for the NGS reads of the PC sample to the NC sample. The adjusted PC reads and the NC reads were compared using Microsoft Excel (Microsoft Corporation, Redmond, WA, USA).

MLPA and NGS data were compared to examine the consistency of the two methods. Following a run on the Genetic Analyzer, relative peak areas of each sample were calculated and compared to five sex-matched controls using the Coffalyser.Net software (MRC-Holland). This program classifies a peak as normal when the ratio to NC is 0.7-1.3, deletes a peak when the ratio is <0.7, and designates a peak as duplicated when the ratio is >1.3. Relative peak area data were extracted from the software and further analyzed using Microsoft Excel.

The PC relative peak areas were calculated using a method similar to the normalization method. Briefly, the sum of all 52 peak areas from the NC were compared to the sum of all 52 peak areas of the PC, thereby resulting in a ratio, and each PC peak area was then multiplied by that ratio, which was then designated as the normalized PC area. The adjusted PC and the NC peak areas were subsequently compared.

Other 22q11DS samples and DMD samples also followed the same analysis procedures. Data were presented as the mean  $\pm$  standard deviation of three repeated experiments.



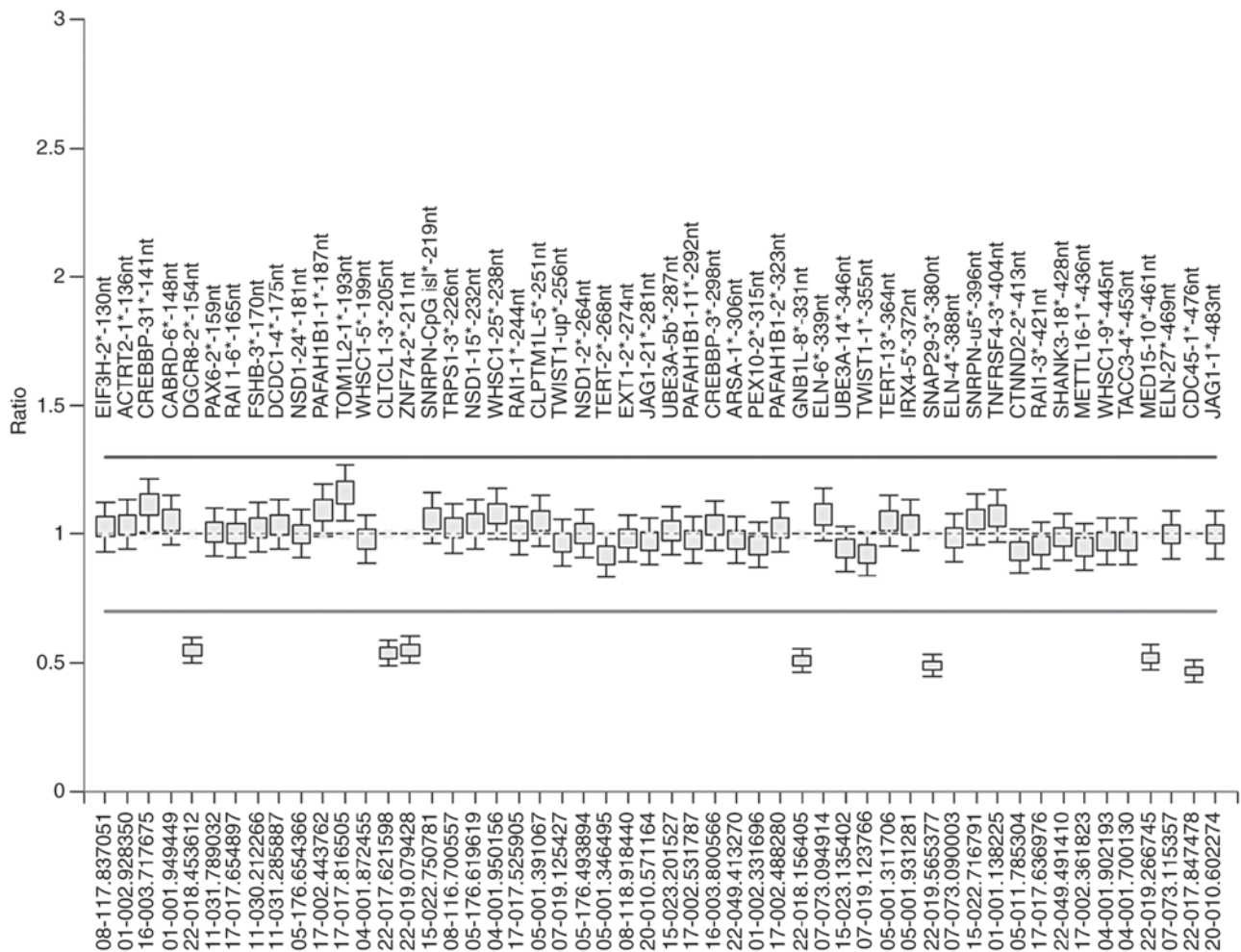


Figure 2. MLPA analysis. The MLPA data of the PC sample were calculated using the Coffalyser.Net software with the NC sample as reference. MLPA represents deletion of the probe in the 22q11 region, including peaks that correspond to 154, 205, 211, 331, 380, 461 and 476 nt. The error bars represent the calculated standard deviations for each probe. MLPA, multiplex ligation-dependent probe amplification; PC, positive control; NC, negative control; nt, nucleotides.

## Results

**MLPA analysis.** MLPA was performed to validate the NGS-MLPA findings of the DNA samples from patients with the 22q11 deletion syndrome or with DMD. For the PC sample, deletions in chromosomal region 22q11, which encompasses the *CLTCL1*, *CDC45*, *GNBIL*, *DGCR8*, *ZNF74*, *MED15* and *SNAP29* genes, are the most frequent cause of DiGeorge syndrome (45).

The 52 pairs of MLPA probes in the P064-C1 set were used to distinguish the seven gene-dosage alterations aforementioned (Fig. 2). In addition, nine control fragments were used, which generated amplicons of <120 nt in size. MLPA was performed using DNA from the NC and PC samples. Analysis using the Coffalyser.Net software identified seven peaks with gene dosage alterations that were clearly distinguishable, using the DNA of the NC sample as a calibrator (Fig. 2).

For the three other 22q11DS samples, it was identified that one female patient carried deletions of *CLTCL1*, *CDC45*, *GNBIL* and *DGCR8* genes, and the other samples carried the same deletions as the PC sample. Using identical protocols, it was determined that four DMD samples carried hemizygous deletions of exons 51, 3-44, 45-48 and 3-11 (PC-Del sample),

and one carried hemizygous duplications of exons 16-44 (PC-Dup sample).

**MLPA-NGS analysis.** All the genes within chromosome 22q11 were detected as single copies in the PC samples, compared with the same fragments from the NC samples, in which the read number at each site was reduced by half. This finding closely matched the MLPA results (data not shown).

The number of reads of the target sequence that was calculated using GATK utilized an alignment score, which affected the final results. For example, the number of NC and PC reads for the seventh fragment (165 bp; number 16526-L20951, RAI1) were 2,745.19 and 3,780, respectively, at a default score of 20; when the score was adjusted to 5, those values were converted into 183,901.82 and 190,662.17, respectively. The latter data demonstrated that on-target rate improved when the alignment score of '5' was used in this experimental condition, for the sequence of these added reads is manually verified to be consistent with the seventh fragment, so we use '5' as the alignment score.

For the PC sample, the PCR products derived from MLPA were re-amplified, purified, gel extracted, ligated with indexed adapters and processed with other protocols, and

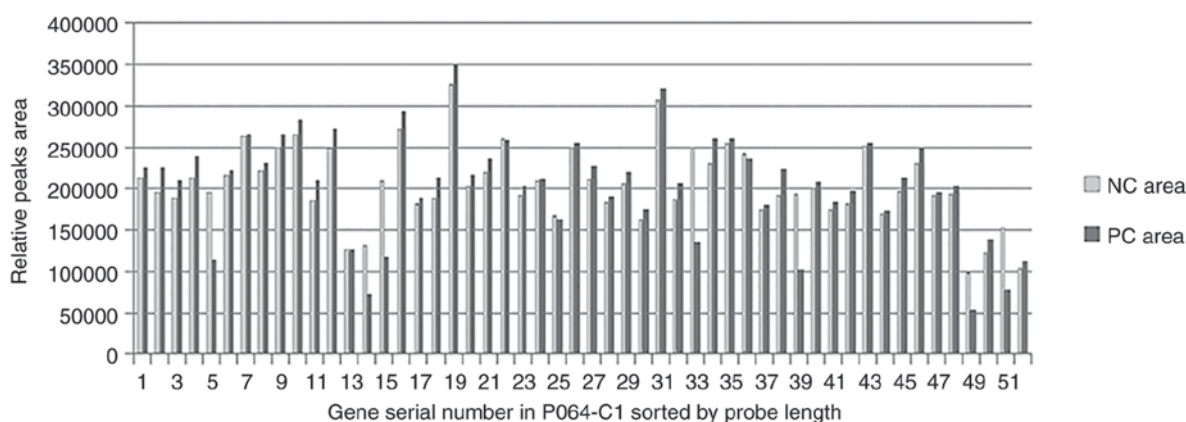


Figure 3. Comparison of MLPA peak areas for NC and PC. The x-axis represents the number of discontinuous peaks. The y-axis represents the peak areas. The PC peak areas were compared to the corresponding peak area from the NC sample. MLPA, multiplex ligation-dependent probe amplification; PC, positive control; NC, negative control.

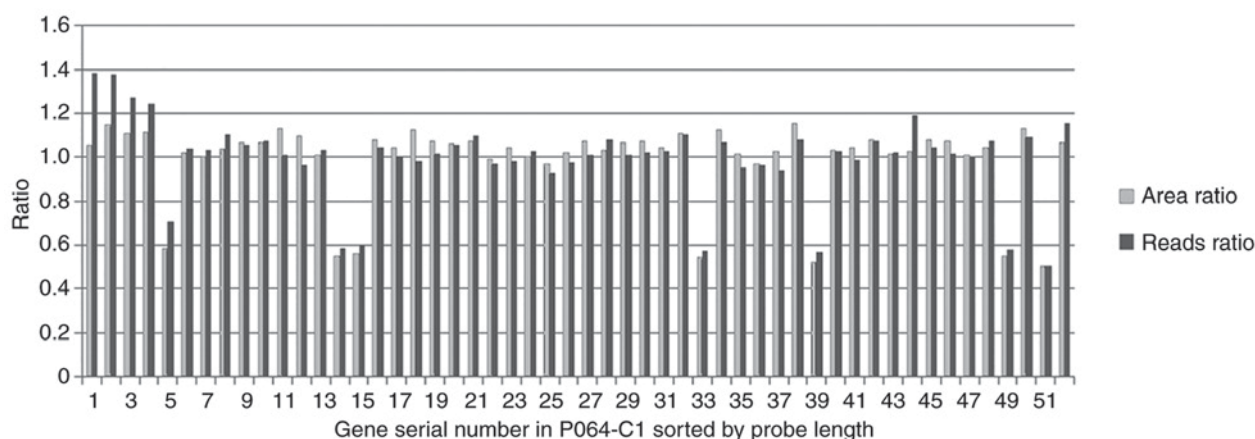


Figure 4. Comparison of area ratios between MLPA and NGS. The x-axis represents the number of discontinuous peaks. The y-axis represents the area ratio and the reads ratio. The area ratio and the reads ratio were compared peak by peak. MLPA, multiplex ligation-dependent probe amplification; NGS, next-generation sequencing.

finally sequenced on an Illumina HiSeq 2500, which generated reads that were analyzed. The NGS reads contained 52 segments, excluding fragments <120 bp in size (which refers to the 92 nt benchmark probe; 64, 70, 76 and 82 bp-long Q-fragments; 88 and 96 bp-long D-fragments; 100 and 105 bp X & Y fragments, and other quality control fragments). The PC read fragments were normalized using similar methods to the relative PC peak area normalization, thereby resulting in normalized PC reads, which were subsequently compared with the corresponding original NC reads, for which the ratio was designated as the reads ratio.

Data from the relative peak areas were extracted from the Coffalyser.Net software and analyzed in Excel, from which the normalized PC and NC peak areas were compared (Fig. 3). The mean peak area of the normalized PC  $\pm$  standard deviation (SD) was  $205,609.19 \pm 46,075.30$  (range, 54,129.87-351,745.90), whereas  $SD/mean = 0.22$ ; the mean peak area of NC  $\pm$  SD was  $205,609.19 \pm 61,965.87$  (range, 98,580-326,506), whereas  $SD/mean = 0.30$ . The area of the peaks, such as numbers 5, 14, 15, 33, 39, 49 and 51 of the PC sample, were notably smaller compared with the corresponding peak area of the NC sample, approximately half of which indicated a gene-dosage

mutation, which was consistent with the results generated by the Coffalyser.Net software.

For the other 22q11DS samples and DMD sample, following the same protocols, the read number at each site was analyzed and the results were consistent with the MLPA results.

**Comparison of MLPA area ratio and NGS reads ratio.** For the PC and NC samples, the area ratio and reads ratio of each peak was compared (Fig. 4). The results of the two methods were in agreement, as peaks 5, 14, 15, 33, 39, 49 and 51 had similar area and reads ratios, both of which were  $\sim 0.5$ -fold lower compared with the other ratios. However, the ratio of the normalized PC reads and the original NC reads was slightly higher compared with the ratio of the corresponding MLPA peak area for peaks 1, 2, 3, and 4, thereby suggesting an error in the analysis.

The normalized PC reads and the NC reads were compared (Fig. 5). The mean peak reads of the normalized NC  $\pm$  SD was  $22,0881.73 \pm 127,415.63$  (range, 36,223.14-559,197.7), whereas  $SD/mean = 0.58$ ; the mean peak reads of PC  $\pm$  SD was  $220,881.73 \pm 136,905.55$  (range, 19,374.58-604,290.46), whereas  $SD/mean = 0.62$ . The

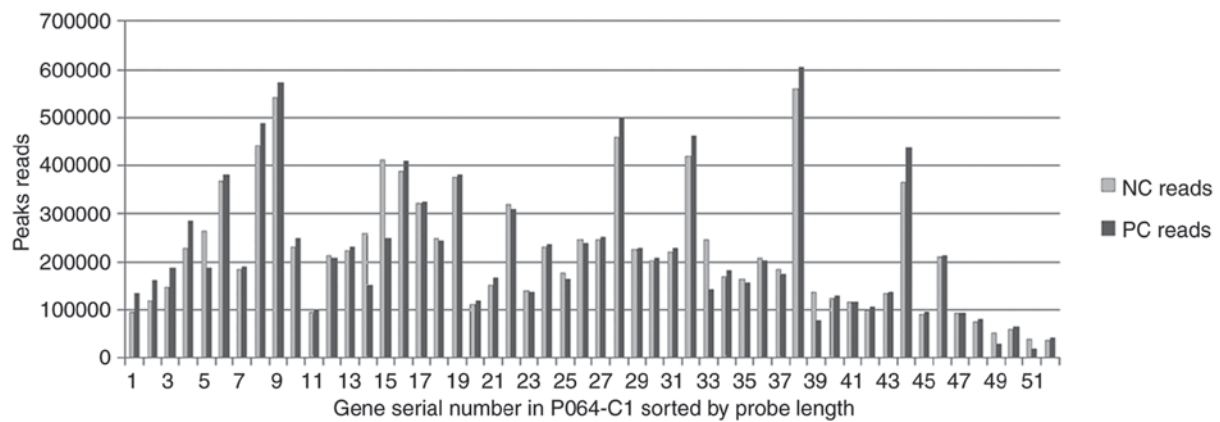


Figure 5. NGS reads for NC and PC. The x-axis represents the number of discontinuous peaks. The y-axis represents the NGS peak reads. The PC peak reads were compared to the corresponding NC peak reads. NGS, next-generation sequencing; PC, positive control; NC, negative control.

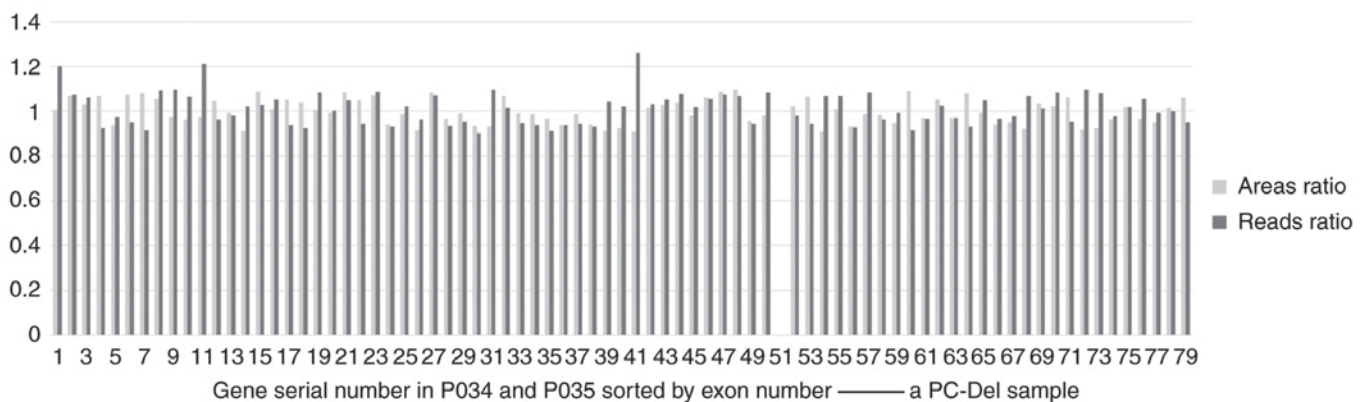


Figure 6. Comparison of area ratios between MLPA and NGS. This was a case of Duchenne muscular dystrophy, which carried hemizygous deletions of exon 51. The x-axis represents the number of exons. The y-axis represents the area ratio and the reads ratio. The area ratio and the reads ratio were compared peak by peak. MLPA, multiplex ligation-dependent probe amplification; NGS, next-generation sequencing.

SD/mean ratio of the NGS reads was higher compared with the SD/mean ratio of the MLPA peak area, thereby illustrating that variations in the NGS data were wider than those of the MLPA data from peak to peak. However, the reads of certain peaks in the PC sample were markedly lower than the corresponding NC peak areas, including peaks 5, 14, 15, 33, 39, 49 and 51, approximately half of which also indicated a gene-dosage mutation, which coincided with the MLPA peak area.

As for the other three 22q11DS patients and five DMD patients, the ratio of the normalized PC reads was consistent with the ratio of the corresponding MLPA peak area for each peak (data not shown). The hemizygous deletions of exons were not detected by both the MLPA-NGS and MLPA (Fig. 6; this case carries hemizygous deletions of exons 51), while the hemizygous duplications of exons were detected by both methods (Fig. 7). As such, it was concluded that the results generated by the two technologies were in good agreement.

**SNP array results.** For the PC Sample, the CNVs in the DNA extracted from peripheral blood were detected by using HumanCytoSNP-12 BeadChip. A 3 Mb deletion within the 22q11 region was observed (Fig. 8), which coincided with the observed absence in the MLPA results.

## Discussion

In the present study, an MLPA product was sequenced using NGS. Following read analysis, large differences were observed among fragments relative to the MLPA results, as indicated by higher standard deviations and mean values from read ratios following NGS, compared with those indicated by the peak area ratios using MLPA capillary electrophoresis. The relatively large standard deviation value reduced confidence in the analysis of the initial copy number of the template. Provided that the number of reads for each fragment is proportional to the amount of the initial template, a fragment containing CNVs may be deduced based on the relative read values of each amplified fragment when no reference sample is present (28). Unfortunately, the read standard deviation for different segments was too excessive to allow this rigorous form of analysis.

Several aspects that contributed to these differences were considered. Initially, amplification bias may have occurred during the first PCR amplification in MLPA and the second amplification with labeled MLPA products as the templates and non-labeled primers to get non-labeled MLPA products. Subsequent PCR steps were performed to ligate adapters to the fragments, further increasing heterogeneity in the

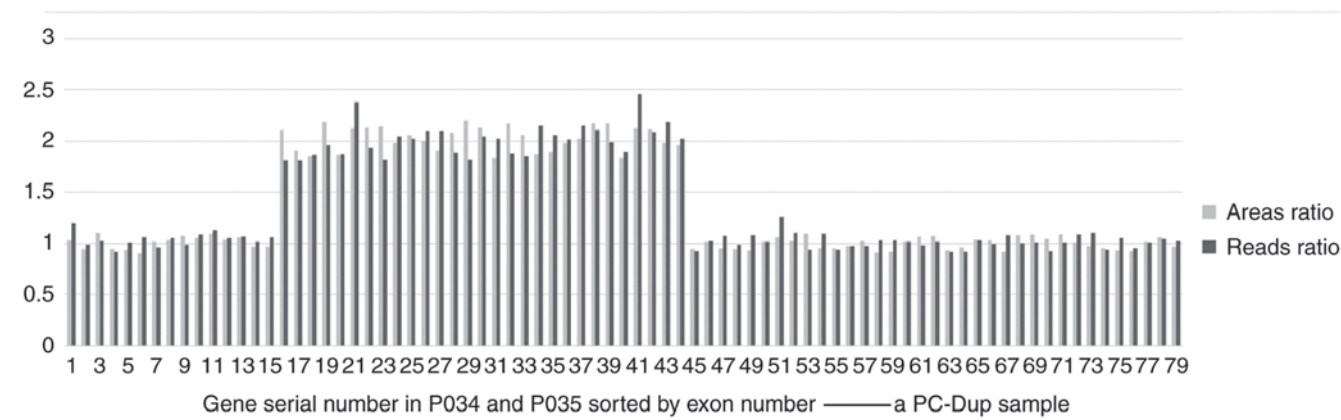


Figure 7. Comparison of area ratios between MLPA and NGS. This was a case of Duchenne muscular dystrophy, which carried hemizygous duplications of exons 16-44. The x-axis represents the number of exons. The y-axis represents the area ratio and the reads ratio. The area ratio and the reads ratio were compared peak by peak. MLPA, multiplex ligation-dependent probe amplification; NGS, next-generation sequencing.

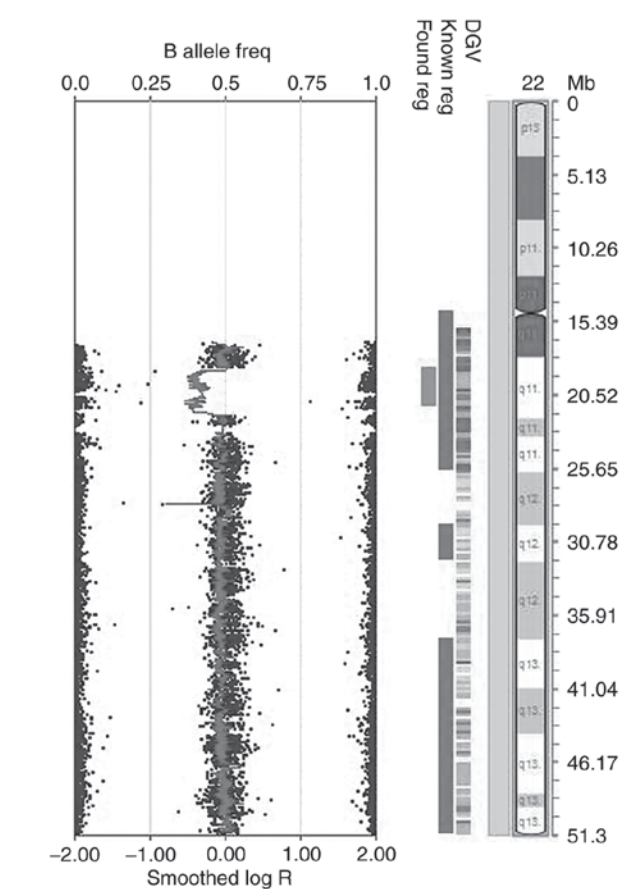


Figure 8. SNP array for the positive control sample. The red line represents the smoothed average log R (bottom plots), which is the ratio between the observed and expected probe intensities, thus indicating CNVs. The B allele frequency (top plots) parameters, which represent the frequency of B alleles at a given SNP, also exhibited signature profiles that specifically identified structural variants. SNP, single-nucleotide polymorphism; CNV, copy number variations.

fragment number. In addition, the gel extraction step after PCR was performed to remove fragments of the wrong size, in order to improve target segment detection. *Taq* polymerase was used for the aforementioned steps, which may have caused greater bias. There are many *Taq* enzymes used in

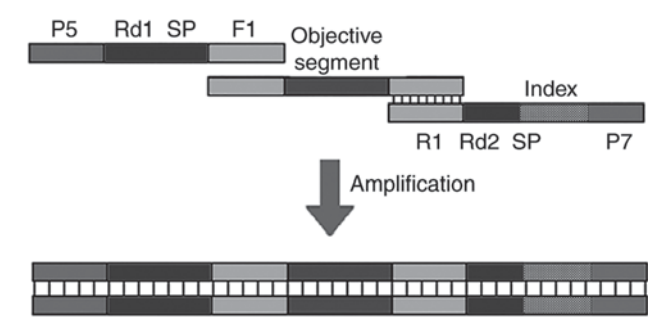


Figure 9. Library construction for the developed MLPA-NGS assay. The left adapter comprised of P5 and Rd1 SP. The right adapter consisted of R1, Rd2 SP, index and P7. MLPA, multiplex ligation-dependent probe amplification; NGS, next-generation sequencing.

the amplification of high-throughput sequencing, which have high requirements for fidelity and bias, but this enzyme is not included (44). Gel electrophoresis revealed that fragments over a certain size range were distributed upstream and downstream of the peak rather than within the peak, and were the brightest sites following ethidium bromide staining. Furthermore, fragments that were closer to the peak were also more abundant. The majority of the fragments were evidently not within the size range (88-480 bp) and were thus discarded during gel extraction, based on the boundaries of 40-550 bp. Thus, the reads for these fragments were relatively low. Furthermore, it was almost impossible to maintain fragment sizes during alignment, particularly for different samples used in gel extraction. The reads of a few peaks at one edge exhibited greater deviation when the position of the gel piece was slightly offset. Overall, the ratio of the normalized PC reads to the original NC reads was >1.2 for several smaller segments, such as 130, 136, 141 and 148 bp, which may have been caused by inconsistent cutting sites. Thus, it was deduced that gel extraction should be excluded from this approach. Fragment purification, which was performed using magnetic beads, caused fragment retention during library construction, which resulted in low yields for very small fragments and subsequently relatively few reads for small fragments. Additionally, there was a large number of



base-pair alterations in the probes during the alignment, as not all reads could be aligned with the real target area during sequence alignment against the human reference genome sequence (GRCh37/hg19), using BWA. Thus, the number of reads of the target sequence that was calculated using GATK utilized a revised alignment score, thereby affecting the final results. Lastly, errors in sequencing may have resulted in further errors, based on the length of the 150 bp pair-end reads generated in both directions. In regards to the small fragments, such as the 130 bp fragment, the homologous sequences could be sequenced twice, thereby doubling the resulting number of reads. For larger-sized fragments, homologous sequences may only be measured from one end, whereas medium-sized fragments are likely to be partially sequenced repeatedly, thereby increasing deviations in length differences between the reads following statistical analysis and finding the actual fragment number.

Overall, there was a high level of diversity of reads among the different fragments following NGS, which was mainly due to the broad range in fragment length, whereas there were only slight differences among the same fragments from different samples. The number of reads of the same fragment from the two samples was highly similar when no CNVs were detected within the area. Taken together, these findings supported the conclusions reached by the MLPA-NGS method, which demonstrated that, relative to the results of MLPA and SNP array chip, the PC sample harbored a single copy of the 22q11 region. These consistent results prove the reliability of the MLPA-NGS results.

The results of other three 22q11DS samples and five DMD samples turned out that a secondary amplification with labeled PCR products as the templates and non-labeled primers would not interfere with accurate CNV detection, as long as the gels were cut accurately. It was assumed that the PCR bias was caused by the standard *Taq* polymerase, instead of the secondary amplification. Using a specific *Taq* polymerase for NGS library construction, such as High-Fidelity 2X PCR MasterMix specified by Illumina, Inc., would likely avoid PCR bias, thus improving the accuracy in CNV detection.

The MLPA-NGS method described herein was a reliable method that would be suitable for detecting the CNVs of target genes at a large scale when performing sample detection together with normal controls. This reliability was based on a certain depth of sequencing. In general, the depth in this experiment was >1,000-fold higher compared with that of ordinary whole-exome sequencing (average depth of 100x), although the absolute extent by which sequencing depth reduced dependability on the final results is unclear.

In the MLPA-NGS method, several further studies are required to improve the assay: First, in the probe designing stage, the MLPA products obtained should all be roughly of the same length, considering that there is no need to distinguish different fragments based on length during capillary electrophoresis. This ensures amplification and purification efficiency of the different fragments, thus rendering consistent detection efficiency for all reads, as well as eliminating the requirement for the number of fragments to be <50 per run. The addition of indices to this high-throughput method may also increase the number of samples in an assay, although this has yet to be verified. To further improve accuracy, limiting

amplification to a one-step PCR method by redesigning the primers may be useful. By using ligase-65 at 54°C during the ligation step of MLPA and stopping the reaction at 98°C, it was possible to remove the stuffer sequences from the ligated product as the adapters were being simultaneously added. Following PCR of the ligated product, the 5'-terminus was subsequently used as an adapter for the NGS, which was made possible by using new adapters for amplifying ligated products, thereby simplifying the MLPA-NGS process. This allowed sample fragments to be detected on the NGS platform following amplification and purification, which was beneficial in reducing both PCR bias and workload. The forward primer (comprising P5, Rd1 SP and F1) and the reverse primer (consisting of R1, Rd2 SP, index or barcode, and P7) contained PCR sequences that were later used in library construction, sample differentiation and NGS sequencing (Fig. 9). Second, due to the simplicity of MLPA-NGS, this method may serve as a powerful tool in classifying tumors or genetic disorders caused by CNVs. MLPA-NGS may be used for the analysis of both CNVs and certain types of variations in genomic DNA derived from peripheral blood, along with various genetic disorders, such as 22q11 deletion syndrome. In addition, point mutation-specific MLPA probes may be designed to detect currently known single nucleotide variants (28). Finally, CNV detection by MLPA-NGS, as well as other types of DNA variations, can be simultaneously analyzed on the same NGS platform. Samples marked with different indices will not interfere with their respective sequencing. Appropriate correction or algorithm optimization will render it more adaptable to data analysis, and the supporting software for this method can be exploited.

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#### Availability of data and materials

The datasets generated and analyzed in the present study are available from the corresponding author upon reasonable request.

#### Authors' contributions

YY and HZ designed the experiments. DW, KX, JJ, ZZ, CX and WX performed the experiments. ZZ and YY analyzed the data. YY and CX wrote the paper. HZ reviewed and edited the manuscript. All the authors have read and approved the final version of this manuscript.

#### Ethics approval and consent to participate

Not applicable.

## Patient consent for publication

Not applicable.

## Competing interests

The authors declare that they have no competing interests.

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