ORIGINAL INVESTIGATION



Low-pass genome sequencing: a validated method in clinical cytogenetics

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Abstract

Clinically significant copy-number variants (CNVs) known to cause human diseases are routinely detected by chromosomal microarray analysis (CMA). Recently, genome sequencing (GS) has been introduced for CNV analysis; however, sequencing depth (determined by sequencing read-length and read-amount) is a variable parameter across different laboratories. Variating sequencing depths affect the CNV detection resolution and also make it difficult for cross-laboratory referencing or comparison. In this study, by using data from 50 samples with high read-depth GS (30×) and the reported clinically significant CNVs, we first demonstrated the optimal read-amount and the most cost-effective read-length for CNV analysis to be 15 million reads and single-end 50 bp (equivalent to a read-depth of 0.25-fold), respectively. In addition, we showed that CNVs at mosaic levels as low as 30% are readily detected, furthermore, CNVs larger than 2.5 Mb are also detectable at mosaic levels as low as 20%. Herein, by conducting a retrospective back-to-back comparison study of low-pass GS versus routine CMA for 532 prenatal, miscarriage, and postnatal cases, the overall diagnostic yield was 22.4% (119/532) for CMA and 23.1% (123/532) for low-pass GS. Thus, the overall relative improvement of the diagnostic yield by low-pass GS versus CMA was ~3.4% (4/119). Identification of cryptic and clinically significant CNVs among prenatal, miscarriage, and postnatal cases demonstrated that CNV detection at higher resolutions is warranted for clinical diagnosis regardless of referral indications. Overall, our study supports low-pass GS as the first-tier genetic test for molecular cytogenetic testing.

Matthew Hoi Kin Chau, Huilin Wang and Yunli Lai contributed equally.

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Introduction

Copy-number variants (CNVs) are defined as gains or losses of genomic segments, the majority of which contribute to human genomic diversity (1000 Genomes Project

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Consortium 2015; Iafrate et al. 2004). However, some are known to cause human diseases (Trost et al. 2018) such as DiGeorge syndrome (also known as 22q11.2 recurrent deletion syndrome; OMIM# 188400) and Charcot-Marie-Tooth disease (17p12 duplication; OMIM# 118220) due to the involvement of dosage sensitive genes (e.g., haploinsufficiency for TBX1 and triplosensitivity for PMP22). Chromosomal microarray analysis (CMA) has been recommended by various professional societies as the first-tier genetic test for detecting clinically significant CNVs in prenatal (Hay et al. 2018), miscarriage (Foyouzi et al. 2012), and postnatal (Miller et al. 2010) applications. However, CMA platforms have different detection resolutions attributed by different probe densities and detection methods (Lin et al. 2015; Wang et al. 2019). In addition, in some applications, the experimental repeat rate is as high as 4.5% due to inherent factors of particular sample types such as amniotic fluid samples (Wang et al. 2020a).

The advantages of next-generation sequencing-based CNV analysis include lower costs and higher throughputs, thus leading to the development of different clinical applications (such as non-invasive prenatal screening and invasive genetic diagnostic testing). As such, low-coverage and highthroughput (low-pass) genome sequencing (GS) has been introduced for CNV analysis in various applications (Chen et al. 2017; Dong et al. 2016, 2019a; Gross et al. 2019; Redin et al. 2017; Wang et al. 2018, 2020b). However, there are various sequencing platforms utilizing different sequencing parameters including sequencing read-amount (ranging from 2 million reads to > 15) and read-length (from 26 to > 150 bp). In addition, detection resolutions for CNVs across different platforms and laboratories are also variable, resulting in difficulties for clinical implementation, genetic counseling and cross-laboratory referencing. Up to date, the majority of published studies focus on the diagnostic yield of NGS-based CNV analysis alone (Chen et al. 2017; Wang et al. 2018, 2020b). In a few studies with limited sample sizes, verification of CNV detection accuracy of low-pass GS-based CNV analysis was compared with CMA in retrospective analyses (Dong et al. 2016; Gross et al. 2019; Li et al. 2014; Liang et al. 2014). Furthermore, no study has evaluated the accuracy of detecting mosaic CNVs. This warrants a study to benchmark and standardize sequencing parameters to ensure accurate and precise detection of CNVs across different laboratories.

In our previous study, simulation of "various CNVs" from YH cell line (Dong et al. 2016) concluded that the minimal read-amount for detecting CNVs at a resolution of 50 kb was 15 million reads, which was equivalent to 0.25-folds (50 bp sequencing read-length) (Dong et al. 2016). Although previously, the accuracy of our detection method was validated on Illumina (Dong et al. 2016) and BGISEQ-500 sequencing platforms (Wang et al. 2020a), in this study, we aimed to

determine the optimal read-amount, sequencing read-length and the minimal mosaic threshold for CNV detection. In addition, our previous prospective back-to-back study demonstrated low-pass GS provided a 1.7% additional diagnostic yield of clinically significant CNVs (smallest 19 kb) in 1023 pregnancies referred for invasive prenatal diagnostic testing. Herein, we evaluate the performance of low-pass GS in prenatal, products of conception and postnatal cohorts in a back-to-back comparison with different designs of CMA assays.

Materials and methods

Samples analyzed

The study was approved by the institutional review boards of each collaborative site. Written consent for sample storage and genetic analyses of peripheral blood and invasive diagnostic samples was obtained from each participant. DNA was previously extracted from prenatal [chorionic villus sampling (CVS), amniotic fluid (AF), cord blood (CB)], products of conceptions (POC) and postnatal samples [peripheral blood (PB)] for CMA testing. The extracted DNA samples were retrieved retrospectively for low-pass GS. Routine quantitative fluorescence polymerase chain reaction (QF-PCR) with short tandem repeat (STR) markers was performed prior to the CMA experiment for exclusion of maternal cell admixture and polyploidies as described (Cheng et al. 2013).

Chromosomal microarray analysis

Two CMA platforms were used in this study. The 8X60K Fetal DNA Chip v2.0 (Agilent Technologies) including array comparative genomic hybridization (aCGH) and singlenucleotide polymorphism (SNP) probes was used for prenatal diagnosis and POC testing (Chau et al. 2019; Huang et al. 2014). The CMA used for postnatal genetic analysis was the 4X180K postnatal array with higher probe densities (Dharmadhikari et al. 2014; Gu et al. 2016). Both were performed according to the manufacturers' protocols and the CNVs were analyzed via the CytoGenomics 5.0 software (Chau et al. 2019; Huang et al. 2014). A minimum of three consecutive probes were required for a positive call on the Fetal DNA Chip, while at least five consecutive probes were required for the postnatal array (Brown et al. 2012). Based on the guidelines of manufacturers, the derivative log ratio spread (DLRS) value of larger than 0.2 was considered a technical failure that required repeat testing (Wang et al. 2020a). Cases with absence of heterozygosity (AOH) detected by CMA platforms were excluded from this study as AOH detection is out of the scope of the current protocol



of low-pass GS (0.25-fold), which is unable to obtain genotype information.

Low-pass genome sequencing

Library construction for low-pass GS was performed with a modified protocol as described in our previous study (Wang et al. 2020a). In short, 50 ng of genomic DNA was fragmented to 200-300 bp by fragmentation end-repair restriction enzyme (MGI Tech Co., Ltd., Shenzhen, China), followed by adaptor ligation and PCR amplification. The concentration of each library was quantified using the Qubit dsDNA HS Assay Kit (Invitrogen, Carlsbad, California, USA) and pooled with equal molality (20–24 samples per lane) and sequenced with a minimum of 15 million reads per sample (single-end 50 bp) on the BGISEQ-500 platform (MGI Tech Co., Ltd., Shenzhen, China) (Wang et al. 2020a). The read depth is estimated to be 0.25-fold, which is determined by multiplying the reads (15 million) and the read length (50 bp), divided by the size of human reference genome (3 Gb).

Data simulation

In this study, high read-depth GS data (30-fold, paired-end 100 bp) from 50 fetuses with increased nuchal translucency published in our previous study (Choy et al. 2019) were downloaded and used for the evaluation of low-pass GS sequencing parameters. Among them, nine pathogenic or likely pathogenic (P/LP) CNVs and nine variants of uncertain significance (VUS) were reported previously. The detection of these CNVs under different simulated sequencing parameters were used to benchmark the optimal read-length and read-amount for low-pass GS. Read 1 (or fastq file 1) from each sample was used as single-end sequencing data (single-end 100 bp, SE100). For shorter read-lengths such as 50 bp and 35 bp, each read was trimmed to the target size. Evaluation of CNV detection using different read amounts was performed by in silico filtering through a process of random sampling to generate datasets with the target readamounts (60, 45, 30, 15 and 10 millions). In addition, for evaluating the mosaic resolution of CNV detection, the nine P/LP CNVs (from eight cases) were analyzed under different simulated mosaic levels. Reads from each sample were pooled with a sex-matched normal control totaling to 15 million reads with single-end 50 bp according to a series of percentages (10%, 20%, 30%, 40% and 50%) to simulate different mosaic levels for subsequent CNV analyses.

Data quality control and CNV analysis

The data QC was performed as previously described (Wang et al. 2020a). After QC filtering, sequencing reads were

aligned to the human reference genome (GRCh37/hg19) using the Burrows–Wheeler Aligner (BWA) (Li and Durbin 2009) with "Aln" and "Samse" alignment modules. Uniquely aligned reads were selected and deposited into adjustable sliding windows (50 kb in length with 5 kb increments) and adjustable non-overlapping windows (5 kb), respectively. The coverage of each window was calculated by the sum of read-amounts undergoing a two-step bias correction (GC correction and population-scale normalization). Genomewide standard deviation (SD) of the copy ratios, calculated from the mean of SD of all windows with the exception of those located on chromosomes with numerical abnormalities, was used as a quality control statistic. In this study, QC cut-off value (genome-wide SD) was also set as 0.1 based on the validation result in our previous study (Wang et al. 2020a). Detection of constitutional and mosaic aneuploidies, homozygous/hemizygous/heterozygous deletions and duplications/triplications was performed with parameters consistent as our reported study (Wang et al. 2020a). The resolution for CNV detection for homozygous/hemizygous deletions was set as 10 kb; and for the other CNVs, it was set at 50 kb. Detection of homozygous/hemizygous deletions was performed by identifying nonoverlapping windows lacking aligned reads. A homozygous/hemizygous deletion would be reported if there are two or more consecutive nonoverlapping windows without aligned reads (copy ratio < 0.1). Therefore, the resolution of homozygous/hemizygous deletion detection is equal to the size of two windows (approximately 10 kb).

Detection of mosaic aneuploidies and copy number variants

Mosaic levels of aneuploidies and CNVs were estimated by the differences of copy-ratio compared with a normal copy ratio (expected as 1). In brief, the mosaic level (percentage) of a duplication with a copy-ratio N was calculated by (N-1)/M, where M is the copy-ratio of one copy in autosomal or sex chromosomes (M equals to 0.5 when the CNV located in autosomal chromosomes or X chromosome in a female, while M equals to 1 when the CNV is located on the X or Y chromosomes in a male). For a mosaic deletion, the mosaic level (percentage) with a copy-ratio N was calculated by (1-N)/M. For mosaic CNV detection, the minimal mosaic level would be set as 20% when the genome-wide SD of windows' copy-ratios was 0.1 (Wang et al. 2020a).

Variant interpretation

Variant interpretation was performed in accordance to the recently published guideline of the American College of Medical Genetics and Genomics (ACMG) (Riggs et al. 2019) using the criteria, methods, and references to online



and in-house datasets such as Database of Genomic Variants (DGV). CNVs were classified into five categories including pathogenic, likely pathogenic, benign, likely benign and variants of uncertain significance (VUS) based on the scoring system recommended by the joint consensus recommendation of the ACMG and the Clinical Genome Resource (ClinGen) (Riggs et al. 2019). The additional findings (P/LP/VUS) from low-pass GS were selected for further verification.

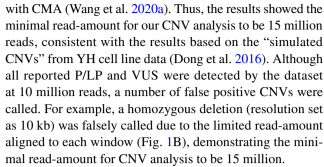
Verification of CNVs and parental confirmation

Quantitative PCR (qPCR) was performed with primer-pairs specific to the candidate regions designed with Primer 3 Web, Primer-Blast (NCBI) and In-Silico PCR (UCSC) based on the GRCh37/hg19 human reference genome (Suppl. Table 1). Melt curve analysis was carried out for each primer-pair to ensure specificity of the PCR amplification and the standard curve method was used to determine PCR efficiency (ranging from 95 to 105%). Each reaction was performed in triplicate in 10-µl reaction mixtures in cases and control (in-house normal male and female controls) on a 7900HT Real-Time PCR System (Applied Biosystems, Foster City, CA, United States) with SYBR Select Master Mix (Applied Biosystems) with the default thermal cycling conditions. The copy number was determined by using the $\Delta\Delta$ Ct method, which compares the difference in Ct (cycle threshold) of the candidate region with a reference primerpair targeting the universally conserved element in the cases and the difference in the controls. Two independent primer pairs (Suppl. Table 1) were used in triplicate for verification of each candidate CNV. Confirmation of parental inheritance was performed whenever parental DNA samples were available.

Results

Sequencing parameters evaluation

To evaluate the optimal read-amount required for the CNV analysis at a resolution of 50 kb (10 kb for homozygous/hemizygous deletion), we generated different datasets with different read-amounts (Fig. 1A) from 50 GS data with nine P/LP and nine VUS ranging from 240 kb to 47 Mb reported by CMA and GS from our previous study (Choy et al. 2019). As SD of the copy-ratios among all windows from each sample is one of the major factors affecting the CNV detection accuracy as demonstrated before (Dong et al. 2016), we calculated the SDs from each read-amount among these 50 cases (mean values and SDs shown in Fig. 1A). Our previous study demonstrated that the cut off of SD values would be 0.1 based on the back-to-back validation data



The sequencing read-lengths utilized by other published studies varied from 26 to 150 bp. We simulated different read-lengths (100 bp, 50 bp and 35 bp) with the same readamount (15 million) for the CNV analysis. Although the dataset with shortest read-length provided the highest alignment rate, the read-utility (considered as the percentage of uniquely aligned reads over the raw reads) in the dataset with 35 bp was the lowest (Mann–Whitney test, P < 0.0001), at approximately 10% less than the dataset with 100 bp (Fig. 1C). However, in theory, longer sequencing readlengths require longer turn-around-times and higher costs. Compared with dataset utilizing 35 bp reads, the dataset utilizing 50 bp reads would be able to provide a significantly lower SD (Mann Whitney test, P = 0.0042, Fig. 1D) due to the significantly higher read-utility obtained (Mann Whitney test, P < 0.0001, Fig. 1C). Although all P/LP CNVs and VUS were detected by each dataset, meaning different readlengths did not affect the sensitivity of P/LP CNV detection, false positives shown in data with 10 million reads as described above (Fig. 1B) demonstrated the false positive detection would be contributed by higher SDs. Thus, it indicated that dataset generated using 50 bp reads would be the most cost-effective read-length for CNV analysis when comparing the performance among datasets with various read-lengths (100 bp, 50 bp and 35 bp).

Although in our previous study, mosaic CNVs were detected and reported with mosaic level ranging from 20 to 40% (Wang et al. 2020a), objective evaluation of the minimal cutoff for mosaic CNV detection is still required. We simulated mosaic CNVs at different levels by pooling each of the eight cases with nine P/LP CNVs ranging from 751 kb to 47 Mb in a series of different artificially reconstituted mosaic levels (from 10 to 50% at 10% intervals) with a sex-matched control sample and performed CNV detection. The results showed that none of the CNVs were detected in datasets at a 10% mosaic levels (Fig. 2A), while only four CNVs out of nine (larger than 2.5 Mb) were detected in datasets at 20% mosaic levels (Fig. 2A). All nine simulated mosaic CNVs were detected when the mosaic levels were at 30% or higher (Fig. 2A). We further replicated the data simulation twice for the datasets with a mosaic level of 30% and all "mosaic CNVs" were detected. The mosaic levels estimated by our methods for all nine P/LP CNVs among



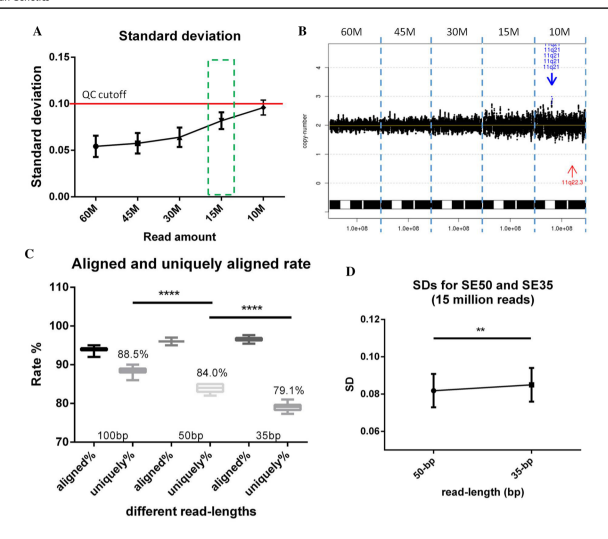


Fig. 1 Evaluation of sequencing parameters for CNV Detection. **A** Distribution of SDs among 50 samples with different read-amounts. Each error bar represents the standard deviation among these 50 samples when filtered to the indicated read-amounts. The *X* axis represents different read-amounts, while the *Y* axis indicates the genome-wide SD values. The red line indicates the cutoff SD threshold validated by our previous prospective study (Wang et al. 2020a), while the dotted green frame shows the optimal read-amount (15 millions) concluded. **B** Distribution of copy-ratios on chromosome 11q14.3q22.3 among different read-amounts simulated in the same sample. The *X* axis shows the genomic location, while the *Y* axis shows the copy-number. In each figure, read amount is shown at the top of the figure, while each black dot indicates an adjustable sliding window (50 kb in size with 5 kb increment). Blue arrows indicate

duplication or copy number gains detected, while red arrow indicates a homozygous deletion identified. **C** Comparison of the percentages of total aligned and uniquely aligned reads over total reads (15 million) among the datasets with 100 bp, 50 bp and 35 bp read-lengths, respectively. The average percentages of uniquely aligned reads for different read-lengths are shown in each bar. Significant difference of the probabilities by Mann–Whitney tests between datasets with 100 bp and 50 bp, and between 50 bp and 35 bp are shown in the top panel (**** indicates P < 0.0001). **D** Distributions of SDs among 50 samples with 15 million reads, sequencing length of 50 bp (SE50) and 35 bp (SE35), respectively. Each error bar represents the SDs among these 50 samples at the indicated read-amounts. The probability of Mann–Whitney test between datasets with 50 bp and 35 bp is shown in the top panel (** indicates P < 0.01)

the three datasets (one from original and two from replications) were on average 29.7% with a SD of 2.3% (Fig. $2A_b$). In addition, the deviation of estimated mosaic levels for each sample across three datasets had a median value of 0.9% (Fig. $2A_c$). Furthermore, of P/LP CNVs selected for data simulation, the smallest was 751 kb. By setting the resolution as 500 kb [based on the minimum size (751 kb) of the P/LP CNVs used as reference] for mosaic CNVs analysis, the specificity of mosaic CNVs detection was 100%.

Prenatal samples

Among the 208 prenatal cases, low-pass GS not only identified all 13 P/LP CNVs reported by CMA (Suppl. Tables 2 and 3), but also provided two additional P/LP CNV findings. First, low-pass GS identified a homozygous deletion of *HBA1* and *HBA2* in a fetus (15BA0039) who presented with multiple ultrasound anomalies, consistent with the Southeast Asian (SEA) type α-thalassemia deletion,



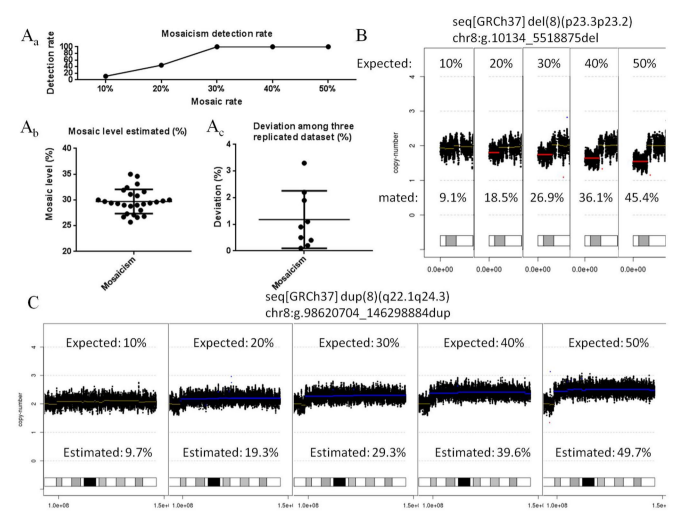


Fig. 2 Evaluation of mosaic CNVs detection. A_a Detection rate for nine "mosaic" pathogenic/likely pathogenic CNVs with different expected "mosaic" levels. A_b Distribution of the estimated mosaic levels in each dataset when the mosaic level is simulated as 30%. The dataset includes the original simulation and two replicates. A_c SD of the mosaic levels detected among three simulated datasets from each case when the mosaic level was set as 30%. Detection of a 5.5 Mb deletion (**B**) and a 47 Mb duplication (**C**) in different mosaic levels with exact genomic coordinates shown in the top panel. **B** and **C**

show the distributions of copy-ratios with each black dot indicating the copy number of an adjustable sliding window (50 kb in size with 5 kb increment). The X axis shows the genomic location, while the Y axis shows the copy number. In each figure, expected mosaic level is shown at the top of each figure, while estimated mosaic level from the dataset is indicated at the bottom. Blue lines indicate mosaic duplication or copy number gains detected, while red lines indicate mosaic deletion or copy number loss identified

which was missed by CMA due to insufficient probe coverage (Fig. 3A). In addition, low-pass GS revealed a likely pathogenic CNV in another fetus (15C0667) with left-hand pre-axial polydactyly. A de novo 122.0 kb intragenic duplication seq[GRCh37] dup(18)(q21.2)dn chr18:g.52903795_53025806dup (Fig. 3B) involving exons 2–10 of gene *TCF4* was detected. Loss of function variants in *TCF4* are known to cause Pitt-Hopkins syndrome in an autosomal dominant manner (OMIM# 610954) (Pontual et al. 2009). In addition, 53% of the reported patients with Pitt-Hopkins syndrome had finger and toe anomalies including over-riding toes, syndactyly,

and polydactyly (Goodspeed et al. 2018), consistent with the malformation in this case. This duplication was classified as a likely pathogenic variant. Among reported postnatal cases, patients often suffer from intellectual disability and epilepsy (Goodspeed et al. 2018) warranting a follow-up study. After investigation, even though there were five probes located in the duplicated region reported by low-pass GS, targeted reanalysis with the CMA software still failed to report this duplication. Overall, low-pass GS provided a 1.0% additional finding of P/LP CNVs (Tables 1 and 2) and one more VUS (Suppl. Table 4) compared with CMA in the prenatal group.



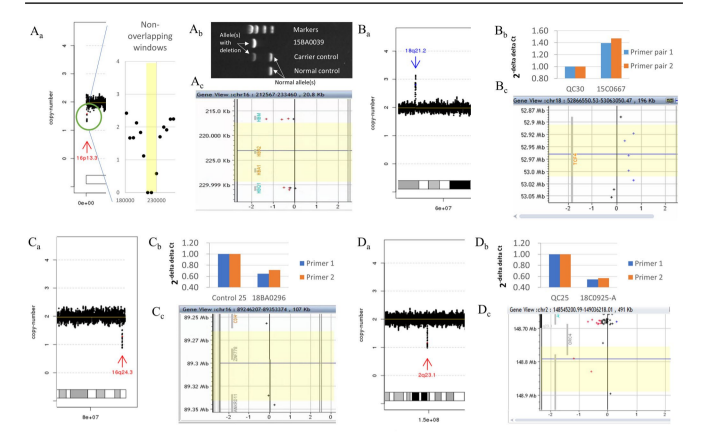


Fig. 3 Additional findings reported by low-pass GS. Low-pass GS identified **A** a homozygous deletion seq[GRCh37] del(16)(p13.3) chr16:g.216050_229978del involving *HBA1* and *HBA2*, known as the Southeast Asian (SEA) type α-thalassemia deletion (15BA0039), **B** a de novo intragenic duplication seq[GRCh37] dup(18)(q21.2) dn chr18:g.52903795_53025806dup affecting gene *TCF4* in a fetus (case: 15C0667), **C** a heterozygous deletion seq[GRCh37] del(16)(q24.3) chr16:g.89264235_89335546del involving partial of *ANKRD11* in a POC 18BA0296, and **D** a de novo heterozygous deletion seq[GRCh37] del(2)(q23.1)dn chr2:g.148755869_148864775del in a postnatal case (18C0925). **A_a**, **B_a**, **C_a** and **D_a** Distributions of copy-ratios with each black dot indicating the copy number of an adjustable sliding window (50 kb in size with 5 kb increment). The

X axis shows the genomic location, while the Y axis shows the copynumber. In the figure, blue arrows indicate duplication or copy number gain, while red arrows indicate heterozygous deletion or one copy number loss. In ${\bf A_a}$ the distribution of copy-ratio with adjustable nonoverlapping windows (5 kb) is also shown with the homozygous deletion region highlighted by yellow box. ${\bf A_b}$ The gap PCR results indicating the SEA deletion. ${\bf B_b}$, ${\bf C_b}$ and ${\bf D_b}$ Bar charts show the results of qPCR validation. Two independent pairs of primers residing in the candidate region were used for CNV verification and all showed consistent findings (duplication or deletion) with low-pass GS compared with a normal control sample. ${\bf A_c}$, ${\bf B_c}$, ${\bf C_c}$ and ${\bf D_c}$ Probe distribution on the CMA platform with the candidate region reported by low-pass GS highlighted in yellow

Table 1 Diagnostic yields in prenatal, products of conception (POC) and postnatal cases by chromosomal microarray (CMA) and low-pass genome sequencing (GS)

Cohort	Sample size	Aneuploidies		P/LP CNVs		Overall		Additional
		CMA, n (%) ^a	Low-pass GS, n (%) ^a	CMA, n (%) ^a	Low-pass GS, n (%) ^a	CMA, n (%) ^a	Low-pass GS, n (%) ^a	diagnostic yield by low-pass GS, n (%)
Prenatal	208	5 (2.4)	5 (2.4)	8 (3.8)	10 (4.8)	13 (6.3)	15 (7.2)	2 (1.0)
Products of conception	221	91 (41.2) ^b	91 (41.2) ^b	10 (4.5) ^b	11 (5.0) ^b	98 (44.3)	99 (44.8)	1 (0.5)
Postnatal	103	0 (0)	0 (0)	8 (7.8)	9 (8.7)	8 (7.8)	9 (8.7)	1 (1.0)
Total	532	96 (18.0) ^b	96 (18.0) ^b	26 (4.9) ^b	30 (5.6) ^b	119 (22.4)	123 (23.1)	4 (0.8)

^aPositive findings are calculated based the number of cases detected with aneuploidy or P/LP CNVs



^bMultiple findings: both aneuploidy and P/LP CNV(s) detected in three cases

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Case ID	Cohort	Indications	CMA platform	CMA result	platform CMA result Low-pass GS result	Interpretation	Reason for missed detection by CMA
15C0667	Prenatal	Ultrasound: left hand pre-axial polydactyly	8X60K	Normal	seq[GRCh37] dup(18)(q21.2)dn chr18;g.52903795_53025806dup	Likely pathogenic: 122 kb de novo intragenic duplication of <i>TCF4</i> . Pitt-Hopkins syndrome in an autosomal dominant manner (OMIM# 610954)	Software/algorithm failed to auto call (5 probes)
15BA0039 Prenatal	Prenatal	Ultrasound: multiple structural abnormalities	8X60K	Normal	seq[GRCh37] del(16)(p13.3) chr16:g.216050_229978del	Pathogenic: <i>HBA1/HBA2</i> homozygous deletion, also known as Southeast Asian (SEA) type α-thalassemia deletion	Lack of probes
18BA0296 POC	POC	Missed abortion	8X60K	Normal	seq[GRCh37] del(16)(q24.3) chr16:g.89264235-89335546del	Likely pathogenic: 71.3 kb deletion involving <i>ANKRD11</i> . KBG syndrome (OMIM#148050)	Lack of probes
18C0925	Postnatal	Postnatal Developmental delay, failure to 4X180 thrive, speech delay	4X180K	Normal	seq[GRCh37] del(2)(q23.1)dn chr2:g.148755869_148864775del	Pathogenic: 103 kb de novo heterozygous deletion involving <i>MBD5</i> . Haploinsufficiency of <i>MBD5</i> causes mental retardation, autosomal dominant 1 (OMIM#156200)	Insufficient probes for detection (2 probes available) ^a

CMA chromosomal microarray, POC products of conception

^aA gain or loss encompassing a minimal of 5 consecutive probes is the threshold for CNV detection on the 4X180K array



Products of conception

Among 221 POCs, CMA identified aneuploidies and P/ LP CNVs in 98 cases and VUS in five cases. In contrast, low-pass GS consistently reported all abnormalities identified by CMA and also detected three additional findings including one LP CNV (Table 2) and two cases with VUS (Suppl. Table 4). In an early miscarriage case (18BA0296) with normal CMA results, low-pass GS reported a 71.3 kb heterozygous deletion (Fig. 3C) seq[GRCh37] del(16) (q24.3) chr16:g.89264235 89335546del involving the gene ANKRD11. Disruptions in ANKRD11 is known to cause KBG syndrome in an autosomal dominant manner (OMIM # 148050). Studies have demonstrated heterozygous premature terminations would result in reduction of RNA expression. This CNV was classified as likely pathogenic based on the current joint ACMG and ClinGen guideline (Riggs et al. 2019). However, the parental samples were not available for further confirmation of inheritance. In another fetus with fetal death at 19 weeks of gestation, low-pass GS detected a heterozygous deletion seq[GRCh37] del(8) (q21.13) chr8:g.82352080_82389112del involving the entire gene *PMP2* (Suppl. Table 4), whereas the CMA was normal. PMP2 mutations reported in the literature cause Charcot-Marie-Tooth disease, demyelinating, type 1G (OMIM # 618279) through a toxic gain of function mechanism (Motley et al. 2016). This heterozygous deletion was classified as a VUS as there was no such deletion reported in the DGV database and our in-house datasets and no parental samples were available for confirmation of inheritance. It is uncertain whether whole gene deletion of PMP2 would result in any phenotypic consequences. All of the three additional findings were due to insufficient probe densities at the targeted regions, hence were not reported by CMA (Suppl. Figure 1 and Suppl. Table 4).

Postnatal samples

Among the group of 103 postnatal cases, low-pass GS identified all eight P/LP CNVs reported by CMA, and one additional pathogenic CNV missed by CMA. A 2.5-year-old girl with developmental delay, failure to thrive and speech delay had a normal CMA report. Low-pass GS identified a de novo 103 kb heterozygous deletion (Fig. 3D) seq[GRCh37] del(2) (q23.1)dn chr2:g.148755869_148864775del involving gene MBD5 (Table 2). Disruptions of MBD5 causes mental retardation, autosomal dominant 1 (OMIM # 156200) in an autosomal dominant manner. The reported patients commonly suffer from delay of motor development, severe psychomotor retardation and language impairment, which were consistent with the presentation of our patient. In addition, seizures were reported in some of the patients. Although our patient did not suffer from epilepsy, further follow-up and

precautionary management for seizures would be beneficial for the patient care. The reason behind the missed detection by CMA was again, insufficient probe density at the targeted region (Fig. 3D_c).

Together, among all 532 samples, low-pass GS detected all P/LP CNVs reported by previous CMA testing and provided an overall 0.8% increased diagnostic yield of P/LP CNVs compared with routine CMA.

Discussion

In the present study, we not only provided the critical sequencing parameters for constitutional and mosaic CNV detection, but also demonstrated that low-pass GS is able to provide an overall ~ 3.4% relative improvement of additional diagnostic yield regardless of referral indications compared with routine CMA.

Our evaluation first demonstrated the optimal readamount for CNV analysis to be 15 million reads at a resolution of 50 kb (10 kb for homozygous/hemizygous deletion). We also show decrease in read-amounts led to an increase in the rate of false-positive calling (Fig. 1A, B). In addition, although read-length would not affect the sensitivity of P/ LP CNV detection in this study, longer read-length would provide significantly higher read utility (Fig. 1C) and significantly lower genome-wide SD (Fig. 1B, D), which would minimize the probability of false calling, particularly for the small CNVs. However, due to longer sequencing turnaround-times and higher costs for longer read-lengths, 50 bp would be the most cost-effective read-length for CNV analysis when comparing with read-lengths of 100 bp and 35 bp. Furthermore, this study also demonstrated that mosaic CNVs can be readily detected using our validated sequencing parameters (15 million reads and single-end 50 bp) for mosaic CNVs at levels as low as 30%, while there were four out of nine CNVs (larger than 2.5 Mb) detected in datasets expected to have 20% mosaic level (Fig. 2). In our previous study, there were two prenatal cases with normal CMA results but mosaic CNVs cryptic to CMA, which indicated that low-pass GS had a higher sensitivity in detecting clinically significant mosaic variants compared to the CMA platforms used.

We further conducted a retrospective back-to-back comparison detecting clinically significant CNVs by low-pass GS (15 million reads, single-end sequencing 50 bp, 0.25-fold) among 532 samples including 208 fetuses, 221 POCs and 103 postnatal cases, compared with CMA. For prenatal and POC applications, the routine CMA platform used was demonstrated to be able to detect > 98% of the known P/LP CNVs reported in the literature for prenatal diagnosis (Chau et al. 2019), while for postnatal analysis, the routine CMA platform employed had higher



probe density (Dharmadhikari et al. 2014; Gu et al. 2016). Nonetheless, low-pass GS yielded an overall diagnostic rate of 23.1% and was able to provide additional findings among different referral cohorts. Since the baseline diagnostic yield by CMA was 22.4%, this resulted in an overall relative improvement of diagnostic yield by low-pass GS to be ~ 3.4% (4/119). However, our previous study reported a 1.7% increase in diagnostic yield in a prenatal cohort of 1,023 cases, which is different from that of our current study (0.75%) among 523 cases. This may be explained by the various referral cohorts (prenatal, postnatal and products of conception) and the sample size of prenatal cohort is limited (n = 208). Interestingly, among the four additional P/LP CNVs identified by low-pass GS, insufficient probe coverage on the CMA was the reason for missed detection in three of them. An intragenic duplication with 122.0 kb in size was missed by CMA even though there were five probes located in the targeted region reported by low-pass GS. After targeted reanalysis of the raw data, the CMA software still reported normal results, suggesting potential flaws exist in the current CMA platforms or software. Furthermore, variant interpretation might be reclassified or changed based on the continuously published updates and guidelines (Riggs et al. 2019). As the variant should receive the same variant classification (P, LP, VUS, etc.), regardless of the clinical phenotype or presentation in the patient, the 71.3 kb heterozygous deletion detected in a POC case, involving gene ANKRD11, was classified as a LP CNV. As such, since small clinically significant CNV exists in POC, our study demonstrates POC analysis may require investigation at higher resolution (Chen et al. 2017; Wang et al. 2020b).

Currently, the resolution for CNV detection suitable for paediatric application is still under debate in previously reported studies with higher read-depth methods (Gross et al. 2019; Trost et al. 2018; Zhang et al. 2019). In this study, although the overall resolution of CNV detection was set at 50 kb as in the application in POC and prenatal samples, low-pass GS was still able to provide 1.0% increased diagnostic yield by comparing with high probe density (4X180K) CMA in the application of postnatal cases. As studies with genotyping SNP arrays show a higher sensitivity in detecting mosaicisms ($< \sim 10\%$) (Markello et al. 2012), it is higher than low-pass GS. However, as the CMA platforms used in this study were not SNP-based arrays, no such comparison could be evaluated. Nonetheless, such study is warranted in future for further evaluation of the performance of low-pass GS.

Finally, by comparing with various genome-wide CNV analysis methods including CMA (aCGH and SNP-array) (Choy et al. 2019)] and GS with different genome readdepths and library construction methods [small-insert (Trost et al. 2018) or mate-pair/large-insert (Dong et al. 2019a, b)], low-pass GS (with single-end 50 bp and small-insert library) described in this study would be the most cost-effective approach (Table 3) as the first-tier genetic testing for CNV analysis from different referral indications. This table does not include data review and variant interpretation time and efforts. In fact, the average number of raw CNVs detected by low-pass GS are more than that

Table 3 Comparison various CMA and sequencing methods for genome-wide CNV detection

	CMA ^a	Low-pass genome sequencing ^b	Mate-pair sequencing ^b (70 mil- lion read-pairs)	Genome sequencing (30-fold) ^b
Turn-around-time				
Experiment (days)	2.5	3 (1 for library construction; 2 for sequencing)	7 (2 for library construction; 5 for sequencing)	6 (1 for library construction; 5 for sequencing)
Data processing and CNV analysis (hours)	1	2	4	24
Reagent cost (USD) ^d	150	100	250	1,000
Resolution of CNV detection	100 kb	50 kb	50 kb ^c	1 kb

^aCMA includes two platforms: aCGH and SNP-array aCGH refers to the 8X60K platform used in this study, and SNP-array refers to CytoScan 750K (Applied Biosystems, Affymetrix, Inc., Santa Clara, CA) in our previous study (Choy et al. 2019). The turn-around-time represents the average of the two platforms

^dThe reagent cost was estimated from routine procurement at our diagnostic laboratory (does not include staff cost for data review and interpretation efforts)



^bThe estimation was based on the BGISEQ-500 platform used in this study and the factors may vary upon different sequencing platforms used. The sequencing depth for low-pass genome sequencing was 0.25-fold as described in this study, while mate-pair sequencing and GS (small-insert) was at ~4.6-fold and 30-fold, respectively

^cThe resolution for CNV analysis by mate-pair genome sequencing was based on our previous study, while the resolution by genome sequencing (30-fold, small-insert) was reported by Trost and colleagues' study (Trost et al. 2018)

of CMA (48 versus 16 in our cohort). However, after filtering the variants against our in-house database, the number of remaining variants that require clinical interpretation are not significantly higher (on average 4.3). Furthermore, there were only three additional VUS identified by lowpass GS in this study, providing the additional yield of VUS as 0.6%. As shown in our previous study, we highlight the importance of referencing public databases such as DGV and constructing a local reference dataset to filter out variants that are not of clinical significance but otherwise due to polymorphisms.

Conclusion

In summary, our study demonstrated the optimal read-amount and the most cost-effective read-length for CNV analysis to be 15 million reads and 50 bp, respectively. In addition, we also demonstrated CNVs with a mosaic level of 30% or higher can be readily detected, while CNVs larger than 2.5 Mb were also detectable at mosaic levels as low as 20%. Our study demonstrated that low-pass GS provided an overall 0.8% of additional diagnostic yield (3.4% relative improvement) of clinically significant CNVs in different applications including prenatal, miscarriage and postnatal cases compared with routine CMA. With its robust performance, our study supports that low-pass GS is suitable and should be recommended as the first-tier genetic test for CNV detection in the various applications.

Author contributions ZD, MHKC, HW, YL, TYL, YK, LX, YZ, WW, and KWC designed the study. HW, YL, TYL, JPWC, SCC, YZ, LX and WW collected the samples and followed up. MHKC, HW, YL, FX, YT, YW and ZC performed CMA and low-pass GS. ZD, MHKC and YK performed the analysis and data interpretation. MHKC, HW and YL conducted the validation. ZD, MHKC, HW, YL, and KWC wrote the manuscript.

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Availability of data and material The genome sequencing data in this study have been made available on the CNGB Nucleotide Sequence Archive (CNSA: https://db.cngb.org/cnsa) under the accession number CNP0000901. Code availability: All relative programs are available at https://sourceforge.net/projects/increment_ratio-of-coverage-v2/files/Increment_Ratio_of_Coverage_V2.0.tar.gz/download.

Compliance with ethical standards

Conflict of interest The authors declare no conflicts of interest or competing interest.

Ethics approval The study was approved by the institutional review boards of each collaborative site. Written consent for sample storage and genetic analyses of peripheral blood and invasive diagnostic samples was obtained from each participant.

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