MapCaller – An integrated and efficient tool for short-read mapping and variant calling using high-throughput sequenced data

Hsin-Nan Lin and Wen-Lian Hsu*

Institute of Information Science, Academia Sinica, Taipei, Taiwan

*To whom correspondence should be addressed (hsu@iis.sinica.edu.tw)

Abstract

With the advance of next-generation sequencing (NGS) technologies, more and more medical and biological researches adopt NGS technologies to characterize the genetic variations between individuals. The identification of personal genome variants using NGS technology is a critical factor for the success of clinical genomics studies. It requires an accurate and consistent analysis procedure to distinguish functional or disease-associated variants from false discoveries due to sequencing errors or misalignments.

In this study, we integrate the algorithms for read mapping and variant calling to develop an efficient and versatile NGS analysis tool, called MapCaller. It not only maps every short read onto a reference genome, but it also detects single nucleotide variants, indels, inversions and translocations at the same time. We evaluate the performance of MapCaller with existing variant calling pipelines using three simulated datasets and four real datasets. The result shows that MapCaller can identify variants accurately. Moreover, MapCaller runs much faster than existing methods. We demonstrate MapCaller is both time and space efficient for NGS data analysis. MapCaller is available at https://github.com/hsinnan75/MapCaller.

1 Introduction

With the advance of next-generation sequencing (NGS) technologies, it is becoming affordable to support various applications of precision medicine in the near future (Ku and Ronkos, 2013). More and more medical and biological researches adopt NGS technologies to characterize the genetic variations between individuals (Dong, et al., 2015; Kulkarni and Frommolt, 2017). Such genetic variations can be classified into three types: (1) single nucleotide variant (SNV, also referred to as SNP); (2) insertion and deletion (indel); and (3) structural variant (SV, including translocation, inversion, copy number variation and indels of size at least 50 bp).

The identification of genome variants is a critical factor for the success of clinical genomics studies (Hwang, et al., 2015). It requires an accurate and consistent analysis procedure to distinguish true variants from false discoveries. This procedure often involves the steps of short read alignment, alignment rearrangement, and variant calling. In each step, one or more tools are applied to generate desired output. For example, BWA (Li and Durbin, 2009), Bowtie (Langmead and Salzberg, 2012; Langmead, et al., 2009), GEM (Marco-Sola, et al., 2012), Subread (Liao, et al., 2013), HISAT/HISAT2 (Kim, et al., 2015), and KART (Lin and Hsu, 2017) are read aligners that can map NGS short reads onto a reference genome and generate their alignments. Samtools (Li, et al., 2009) and Picard (Carver and Stubb, 1997) provide various utilities for manipulating read alignments. For variant calling, the Genome Analysis Tool Kit (GATK) (DePristo, et al., 2011), Freebayes (Erik Garrison, 2012), Platypus (Rimmer, et al., 2014), VarScan (Koboldt, et al., 2009) and SAMtools are widely used.

Different combinations of those tools produce various analysis pipelines. Thus different variant calling pipelines may generate substantial disagreements of variant calls. Several studies (Hwang, et al., 2015; Yen, et al., 2017) have been conducted to confirm the disagreements. Besides, all existing variant calling pipelines are time and space consuming. They require read alignments are sorted and stored in desired format.

In this study, we present MapCaller, an efficient and versatile NGS analysis tool, by integrating the algorithms for read mapping and variant calling. For read mapping, we adopt a divide-and-conquer strategy to separate a read into regions with and without gapped alignment. With this strategy of read partitioning, SNVs, indels and breakpoints can be identified efficiently. For variant calling, MapCaller maintains a position frequency matrix to keep track of every nucleobase’s frequency at each position of the reference genome while mapping the input read sequences.

Since MapCaller collects all information required for variant identification while reads are mapped onto the reference genome, variants can be called directly in the same process. Therefore, the conventional analysis pipeline can be simplified greatly. Most existing variant callers can only detect a few specific types of variants, however MapCaller can detect multiple types of variations, including SNVs, indels, inversions and translocations. We demonstrate that MapCaller not only produces comparable performance on variant calling, but it also spends much less time compared to some selected variant calling pipelines. MapCaller was developed under Linux 64-bit environment and implemented with standard C/C++. It takes read files (FASTA/FASTQ) as input and outputs all predicted variants in VCF format. The source codes of MapCaller and benchmark datasets are available at https://github.com/hsinnan75/MapCaller.

2 Methods

MapCaller aligns every short read onto a reference genome and collects the alignment information during read mapping to deduce sequence variants. MapCaller uses a modified algorithm of KART to perform read mapping. It also maintains a position frequency matrix to keep track of every
nucleobase’s frequency at each position in the reference genome and collects all insertion and deletion events that are identified during read mapping. Furthermore, MapCaller learns all possible breakpoints from discordant or partial read alignments. Finally, MapCaller finds sequence variants based on all of the above-mentioned information. The novelty of our algorithm derives from the integration of read mapping and variation information gathering into a coherent system for variant calling.

2.1 Read mapping and alignment profiles
The details of read mapping method, KART can be found in our previous study. Here we focus on the high-level methodology description. KART adopts a divide-and-conquer strategy to handle matches and mismatches separately between read sequence and reference genome. KART identifies all locally maximal exact matches (LMEMs). We then cluster them according to their coordinates and fill gaps between LMEMs to create final alignments. In doing so, KART divides a read alignment into two groups: simple region pairs (abbreviated as simple pairs) and normal region pairs (normal pairs), where all simple pairs are LMEMs and normal pairs are gaps between simple pairs and might require gapped alignment (due to mismatches or indels). Thus, all SNVs and indels should be found in normal pairs, and breakpoints can be deduced from normal pairs at either end of read alignment. We will explain the identification metrics of each variant type below.

For the convenience of describing the methodology of MapCaller, we define the following notations. Given a read sequence \( R \), the reference genome \( G \), let \( R_i \) be the \( i \)-th residue of \( R \) and \( G[j, i] \) be the substring between \( R_j \) and \( R_i \). Likewise, let \( G_j \) be the \( j \)-th nucleotide of \( G \) and \( G[j, j] \) be the substring between \( G_j \) and \( G_j \). A simple pair (or a normal pair) consists of a read’s substring and its counterpart of reference’s substring. They can be represented as \( (R[i, i], G[j, j]) \). One or more simple/normal pair forms a candidate alignment. We perform pairwise alignment for each normal pair in a candidate alignment.

We also check the alignment quality of a normal pair at either end of the read sequence to determine whether they should be removed. The quality evaluation is as follows. Given an alignment of a normal pair at either end, MapCaller counts the number of mismatches and state transitions of the alignment. A state transition is an alignment state change at the end reads Read3 and Read4 are mapped with same orientation due to an inversion event in the sample genome. each short read, we only keep the alignment with the highest alignment score.

MapCaller creates a position frequency matrix (PFM) to count the nucleobase frequencies at each position. PFM is a matrix of \( 4 \times L \), where \( L \) is the reference genome size. Therefore, each column of PFM represents the frequencies of nucleobases A, C, G and T at that position of the reference genome. MapCaller updates the nucleobase frequency in each column according to read alignments. Fig. 2 shows an example to illustrate how PFM works in this study. Five reads are mapped onto the reference genome. MapCaller counts the frequencies of each involved column. For example, PFM[3] = (1, 0, 0, 4) indicates there are one ‘A’ and four ‘T’s aligned at the third position of the reference genome. Any insertion and deletion events are kept opposite. We use a 3-tuple, \( \text{Ins}(Gpos, R[i, j], k) \) and \( \text{Del}(Gpos, G[m, n], k) \) to represent an insertion and deletion event, where \( Gpos \) indicates the occurrence location, \( R[i, j] \) and \( G[m, n] \) are the indel strings, and \( k \) is the number of occurrences. MapCaller would create the following three 3-tuples: \( \text{Ins}(3, T, 1), \text{Del}(5, C, 1) \) and \( \text{Del}(8, CG, 1) \) based on the example cases in Fig 2. We describe how the SNVs, indels, insertions and translocations are identified based on the PFM and by MapCaller below. It is noteworthy that the quality measurement of each type of variant is described in the Supplementary.

2.2 SNV detection
MapCaller uses PFM to keep track of the frequencies of the four nucleobases at each position in the reference genome. The depth of position \( p \), denoted as \( \text{Depth}(p) \), where \( \text{Depth}(p) = \text{PFM}[p, A] + \text{PFM}[p, C] + \text{PFM}[p, G] + \text{PFM}[p, T] \). We partition the reference genome sequence into blocks of 100 nucleotides. For each block \( i \), MapCaller determines a threshold, denoted as \( \text{depth}(i) \), which is the half of the average depth for all the nucleotides within the block.

A nucleobase at position \( p \) is considered an alternative allele if its frequency is above \( \text{FreqThr}(p) = \text{Depth}(p) \times \text{MinAlleleFrequency} \), where \( \text{MinAlleleFrequency} \) is a user defined threshold (the default value is 0.2 for germline mutation and 0.02 for somatic mutation). Thus, a nucleobase \( b \) at position \( p \) is considered an SNV if the following two conditions are satisfied: (1) \( \text{Depth}(p) \geq \text{depth}(i) \); (2) \( \text{PFM}[p, b] \geq \text{FreqThr}(p) \). Since the sample genome may carry multiple sets of chromosomes, the genotype at the same position can be homozygous or heterozygous. MapCaller considers both haploid and diploid scenarios. If only one nucleobase is called and its frequency is less than \( 1\times \text{MinAlleleFrequency} \), then the locus is considered heterozygous, otherwise it is homozygous.

2.3 Indel detection
3.1 Experiment design

We develop a simulator to generate genome variations using the E.Coli K-12 strain, human chromosome 1 and human reference genome (GRCh38).

The simulator (described in the Supplementary material S8) randomly generates sequence variations with occurrences of 2700 substitutions, 180 small indels (1-10 bp), 45 large indels (11-50 bp), 1 translocation (TNL, size ranges 1000-2000bp), 1 inversion (INV, size ranges 1000-2000bp) and 1 copy number variation (CNV, size ranges 300-1300bp) for every 1,000,000 base pairs. We use WGSIM (https://github.com/lh3/wgsim) to generate simulated short read sequences for each mutant genome. The simulated read coverage is 30X, sequencing error rate is 0.02, and read length is 100bp. The synthetic datasets are referred to as Sim_Ecoli, Sim_Ch1 and Sim_HG respectively. We also download four real NGS datasets from SRA web site, two from sample of HG001-NA12878 (RUN: SRR6062143 and SRR7781445) and two from sample of HG002-NA24385 (RUN: SRR3440404 and SRR6691661), where the dataset of SRR3440404 includes short reads from RUNs of SRR3440404 to SRR3440422 in order to have enough read depth. Note SRR6062143 and SRR7781445 are two separate runs of Illumina sequencing data of NA12878. SRR3440404 and SRR6691661 are also two separate runs of NA24385. GIAB (The Genome in a Bottle Consortium) provides high-confidence SNP, small indel calls for the two sample genomes. They can be found at ftp://ftp-trace.ncbi.nlm.nih.gov/giab/ftp/release/. Those variant calls were made by GATK, Freebayes and SentiOn. Table 1 shows the number of short reads of each dataset as well as the number of each type of variants. It is noteworthy that GIAB does not provide structural variant annotation for samples NA12878 and NA24385.
A conventional analysis pipeline includes a read mapper, SAM/BAM file processing, and a variant caller. Thus, in study, we compared the performance of Map Caller with different combinations of those read mappers and variant callers. For read mapping, we selected KART, BWA-MEM (BWA for short), Bowtie2, and GEM. For the variant calling, we selected Genome Analysis Tool Kit Haplo typeCaller (GATK for short), Freebayes and SAMtools mpileup (Mpileup for short). For SAM/BAM file processing, we used SAMtools view/sort to perform file format converting and alignment sorting. We also compared the performance of structural variant calling with existing methods, including DELLY (Rausch, et al., 2012), LUMPY (Layer, et al., 2014), and SVDetect (Zeitouni, et al., 2010). The commands as well as the argument setting used for each pipeline are shown in Table S1 (Supplementary material).

It is noteworthy that since Map Caller integrates the algorithms of read mapping and variant calling, it handles the whole procedure of analysis pipeline alone. The run time was estimated from the read mapping to variant calling. We estimated the precision and recall for each dataset. We define a true positive case (TP) as a true variant call; a false positive case (FP) as a false variant call; and a false negative case (FN) as a true variant that is not called. A predicted SNP event is considered an TP if its genomic coordinate is correct without any tolerance, otherwise it is a FP. A predicted indel event is considered an TP if the genomic coordinate is within 10 bp, otherwise it is a FP.

3.2 Performance comparison on synthetic datasets

Table 1 summarizes the comparison result on the three synthetic datasets. We tested every combination of read mapper and variant caller. We found that BWA-MEM combined with any variant caller generally performed better than any other selected read mapper did; therefore, we only showed the performance of pipelines involved with BWA-MEM here to compare to Map Caller. The complete experiment result can be found in Supplementary (Table S2 and the excel file).

It can be observed that Map Caller and Freebayes performed comparably on the three synthetic datasets; however, Map Caller produced better performance on Sim_HG with respect to recalls. The SNP and indel recalls of Map Caller on Sim_HG are 96.9% and 96.3% respectively. When further analyzed some of the false negative cases, we found that most of false negatives occur at repetitive or highly similar regions. Freebayes and mpileup produced higher precisions on indel detection on Sim_HG; however, they compromised the recalls. The precisions and recalls of Freebayes are 99.9% and 91.6%, and those of mpileup are 99.9% and 91.0%. Map Caller produced balanced result on indel detection. Its precision and recall are 99.1% and 96.3% respectively.

Surprisingly GATK produced poor recalls on SNP and indel detection on all of the three datasets. The recalls on SNP detections were all below 80%, and those on indel detection were all below 20%. In this experiment, we fed all the three variant callers with the same read alignments; however, the fact that Freebayes and mpileup produced much higher recalls on the three datasets suggested BWA could process those indel events correctly in the alignments. We tried to diagnose the reason by configuring the pipeline but it did not help. We then increased the read depth from 30X to 100X (Sim_Ecoli) and we found both of the recalls improved from 76.4% to 95.7% (SNV) and from 15.9% to 65.7% (indel). If we further increased the read depth to 200X (Sim_Ecoli), the two recalls achieve 96.7% and 87.2% respectively. It suggested GATK performs much better on NGS data with deeper coverage. The performance regarding different read depths is shown in Supplementary (Table S3). We also found that if the NGS data were error-free, then the performance of GATK would be very good. We simulated an NGS data without any sequencing errors (read depth: 30X, Sim_Ecoli) and tested with GATK combined with BWA, the precision and recall on SNP detection were 100% and 99.4% respectively.
and those on indel detection were 99.5% and 99.0% respectively. It implies that GATK is more sensitive to sequencing errors.

Another noteworthy observation is that Mpileup performed much worse on Sim_HG. The precision on SNV detection was only 44.7%. When we further analyzed the output, we found that Mpileup generated many false positive SNVs at loci with very shallow depth. Fortunately, we can filter out some of the false calls according to their confidence scores. We will investigate the relationship between variant calling performance and confidence score later.

In terms of run time, it can be observed that MapCaller is much more efficient than the selected pipelines. For example, MapCaller only spent 148 minutes to handle Sim_HG. However, the other three pipelines spent 3414, 6954, and 36812 minutes respectively on the dataset. Since the variant analysis pipeline consists of multiple steps, we can further decompose the runtime into multiple parts. Take the pipeline of BWA+GATK for example, it spent 1444 minutes for read mapping, 249 minutes for SAM/BAM file processing, and then another 35119 minutes for variant calling. If we only considered the run time for variant calling, MapCaller only spent 2.6 minutes (155 seconds). MapCaller is much more efficient because it not only adopts a very efficient read mapping algorithm, but it also saves run time by skipping processing SAM/BAM files and collects variant information directly during read mapping.

3.3 Performance comparison on real datasets

We downloaded four real NGS datasets sequenced for genomes of NA12878 (HG001) and NA24385 (HG002). However, the two genomes do not have ground truth variant annotation. GIAB provides reference calls that were made by GATK, Freebayes and Senteon for the sample genomes. In this study, we used the reference calls as the gold standard to estimate the performance of each method.

Table 3 summarizes the comparison result. It can be observed that MapCaller, Freebayes and GATK performed comparably. It is not obvious that which method performed the best. Each method has its strength and weakness. For example, MapCaller and GATK generally produced higher precisions on SNV detections than the other two methods. Freebayes and GATK generally produced higher recalls on indel detections. However, it seemed like Mpileup produced relatively lower precisions on SNV detection and lower recalls on indel detection. Mpileup still generated many false positive cases on SNV detection. Its precisions were all below 20%. We also found that all of false calls made by MapCaller were due to ambiguous alignments of indels. If we relaxed the tolerance of locus difference for indel events to 50bp, the precisions and recalls of MapCaller on indel detection would be increased by around 10% and 4% respectively.

Moreover, some of false negative cases were due to poor alignment quality and their read alignments were discarded by MapCaller. For example, the reference calls reported three adjacent variants, which were chr1:20761541 (an insertion: AGAG), chr1:20761544 (SNV: C), and chr1:20761545 (a deletion: AT). Since there were more than three state transitions in the corresponding alignment, MapCaller chose not to report such variants. If two adjacent variants appear in two separated alignments, they could be still called by MapCaller. It is estimated NA12878 contains 11,018 indel events (2.1%) that are adjacent to one another within five nucleotides, and NA24385 contains 10,016 variants (2.0%). By contrast, MapCaller produced around 0.7% of indels that are adjacent within five nucleotides in average.

Since the reference calls were made by integrating three different callers, we analyzed the performance of overlapping calls by integrating MapCaller, Freebayes, Mpileup and GATK on the dataset of SRR6062143. If an SNV was called by all the four methods, the precision was 83.6%. If an indel was called by all the four methods, the precision was 79.2%. It suggested that the overlaps of variant calls could increase calling accuracy significantly. Moreover, we also found the union of the four callers could cover 99.9% of reference SNVs and 98.6% of reference indels. The performance with respective to overlaps among the four callers for each dataset was shown in Table S4 (Supplementary material).

In terms of run time, MapCaller was still much faster than any other pipeline. It is around 100 times faster than BWA+GATK. It spent around one or two hours to handle a human genome data with around 30X of read depth. If we only consider the run time for variant calling, MapCaller is much faster than GATK. For example, MapCaller spent three minutes on variant calling for SRR6062143, while GATK spent 203.8 hours for the same dataset. Though MapCaller runs very fast, it produces comparable result as Freebayes and GATK do in this analysis. Thus, we demonstrated MapCaller is a highly efficient variant calling method.

3.4 Variant filtering

We estimate the performance of MapCaller and each selected pipeline using the raw result. However, most variant callers provide specific filters to remove unlikely variants based on their algorithm design and statistic models; it may complicate the comparison if we try to optimize the filtering strategy for each caller. According to the performance comparison result using synthetic datasets, we found that MapCaller and the other selected methods produced high precisions on SNV and indel detection, except Mpileup lost its SNV accuracy on the dataset of Sim HG. It suggested that MapCaller, Freebayes and GATK could produce reliable variant calls without any specific filters. However, Mpileup tends to produce more SNV calls. It is necessary to filter out false calls to improve the accuracy. Since all variant callers give an estimate on how likely a variant
call is true with a quality score (the QUAL column), we analyzed the SNV accuracy regarding Mpileup on Sim_HG, SRR6062143, SRR7781445, SRR3440404 and SRR6691661 to investigate the relationship between precision/recall and QUAL values.

We filter out SNV calls with different QUAL thresholds. Fig 4 illustrates the analysis result. The QUAL threshold is from 0 to 180. Note that almost all of the quality scores of SNV calls for Sim_HG are below 150. It can be observed that the precision improves more and more when the QUAL threshold becomes higher and higher. On the other hand, the recall deteriorates at the same time. It is interesting that the quality scores of Mpileup are not consistent with the accuracy. For example, if the QUAL threshold is 90, we could observe the SNV accuracy on the five datasets varies a lot. It is much higher on Sim_HG, but it is lower on the real datasets. The highest precision is 82.3 while the lowest is only 21.5%. It implies the quality scores do not actually reflect the calling confidence. That MapCaller produced high precisions and recalls on all the three simulated datasets. In particularly, MapCaller produced the highest precisions among these callers. LUMPY produced high precisions and recalls on inversion detections. However, it produced relatively lower precisions and recalls on translocation detections. SVDetect lost precisions on translocation and inversion detection, but it generated much higher recalls on the two types of structural variants. Similar to LUMPY, DELLY generally produced high precisions and recalls on inversion detections, but it could not detect any translocation events on the three datasets. In summary, MapCaller was the only method that handled both translocation and inversion detection at high precision and recall.

4 Conclusions
In this manuscript, we present MapCaller, an integrated system for read mapping and variant calling. MapCaller collects read alignment information during read mapping and maintains a position frequency matrix to keep track of alignments at each position of the reference genome. We evaluate the performance of MapCaller and the selected variant calling pipelines using three synthetic datasets and four real data sets from human genomes. The comparison results show that MapCaller not only identifies highly accurate variants, but it also spends the least amount of time. MapCaller is also versatile. It is capable of identifying SNVs, INDELs and structural variants simultaneously.

All current variant calling pipelines require SAM/BAM files as inputs and take multiple steps to rearrange the alignments for variant calling. It is not only time consuming, but also requires a huge amount of disk space. MapCaller identifies variants directly from the NGS short reads. Thus, it is not necessary to output SAM/BAM files and it saves a lot of disk space. Pipelines involved with GATK require even more disk space for the additional preprocess of alignments. Thus, MapCaller is both time and space efficient. In particularly, MapCaller is very user-friendly. It takes only one command line to handle read mapping and variant calling. Since more and more medical and biological researches adopt NGS technologies to characterize the genetic variations between individuals, we believe MapCaller is a fast and reliable variant calling method.

Acknowledgements
Funding: This work was supported by Bioinformatics Core Facility for Translational Medicine and Biotechnology Development/Ministry of Science and Technology (Taiwan) 105-2319-B-400-002.
Conflict of Interest: none declared.

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