Towards Better Understanding of Artifacts in Variant Calling from High-Coverage Samples

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ABSTRACT

Motivation: Whole-genome high-coverage sequencing has been widely used for personal and cancer genomics as well as in various research areas. However, in the lack of an unbiased whole-genome truth set, the global error rate of variant calls and the leading causal artifacts still remain unclear even given the great efforts in the evaluation of variant calling methods.

Results: We made ten SNP and INDEL call sets with two read mappers and five variant callers, both on a haploid human genome and a diploid genome at a similar coverage. By investigating false heterozygous calls in the haploid genome, we identified the erroneous realignment in low-complexity regions and the incomplete reference genome with respect to the sample as the two major sources of errors, which press for continued improvements in these two areas. We estimated that the error rate of raw genotype calls is as high as 1 in 10–15kb, but the error rate of post-filtered calls is reduced to 1 in 100–200kb without significant compromise on the sensitivity.

Availability: BWA-MEM alignment and raw variant calls are available at http://bit.ly/1g8XqRt. Scripts and miscellaneous data at https://github.com/h3varcomp.

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1 INTRODUCTION

Since the sequencing of the first personal genome (Levy et al., 2007) and in particular the first genomes sequenced with the Illumina technologies (Bentley et al., 2008; Wang et al., 2008), resequencing has been widely used for personal and cancer genomics (Watson et al., 2013), for the discovery of de novo mutations associated with Mendelian diseases (Bamshad et al., 2011), for the reconstruction of human population history (Li and Durbin, 2011) and for the understanding of mutation processes (Veltman and Brunner, 2012; Campbell and Eichler, 2013). In most of these studies, mapping based SNP/INDEL calling plays a central role. The accuracy of the calls has a fundamental impact on the biological interpretation. In this context, various research groups have attempted to evaluate the performance of variant calling.

The simplest approach to the evaluation of variant calling is to simulate variants and reads from a reference genome (Li et al., 2008). However, we are unable to simulate various artifacts, such as the non-random distribution of variants, dependent errors, incomplete reference genome and copy number variations. An improved version is to incorporate real variants instead of using simulated variants (Talwalkar et al., 2013), but it does not address the artifacts caused by large-scale effects, either. A better simulation is to take the reads sequenced from one sample with a finished genome, map them to another finished genome, call variants and then compare the calls to the differences found by genome-to-genome alignment (Li et al., 2008). However, this approach is limited to small haploid genomes. There are attempts to apply a similar idea to mammalian genomes (Li et al., 2013; Bolosky et al., 2014), but as the mammalian reference genomes are frequently incomplete and the whole-genome alignment is imperfect, such a simulation is still different from realistic scenarios.

The difficulties in simulation have motivated us to focus more on real data. One simple approach is to thoroughly sequence a small target region with mature technologies, such as the Sanger sequencing technology, and take the resultant sequence as the ground truth (Harismendy et al., 2009). It does not capture large-scale artifacts, though. Another more commonly used method is to measure accuracy either by comparing variant calls from different pipelines, or by comparing calls to variants ascertained with array genotyping or in another study (Clark et al., 2011; Li, 2012; Lam et al., 2012a,b; Boland et al., 2013; Liu et al., 2013; Goode et al., 2013; O’Rawe et al., 2013; Zook et al., 2014; Cheng et al., 2014). However, array genotyping is biased to easier portions of the genome and may have a higher error rate per assayed site than the variant calling error rate (Bentley et al., 2008); simply comparing call sets would only give us an estimate of the relative accuracy – if two pipelines are affected by the same artifact that a third pipeline does not have, the third pipeline will appear worse even though it is in fact better. In addition, comparative studies usually measure the accuracy with summary statistics such as the fraction of calls present in dbSNP or the transition-to-transversion ratio. They do not tell us the wrong sites.

Many studies also experimentally validated typically up to a few hundred variants with MiSeq or Sanger sequencing or Sequenom genotyping. Nonetheless, such experiments are biased towards easier regions and may also be subjected to other artifacts such as on-primer variants and non-specific amplification (the 1000 Genomes Project analysis subgroup, personal communication). Calling heterozygotes from Sanger sequence data is also challenging by itself.

In the author’s view, it is better to evaluate variant calling by comparing samples from a pedigree (Zook et al., 2014), or from the same individual (Nickles et al., 2012), including cancer samples (Löwer et al., 2012). Because we expect to see only tens to hundreds of somatic mutations or Mendelian errors per...
2 DATA SETS AND DATA ANALYSIS

2.1 Data sets

In this study, we focused on deep Illumina sequencing data from two cell lines, the CHM1hTERT cell line Jacobs et al. (1989) and the NA12878 cell line. A crucial and unusual feature of CHM1hTERT, or briefly CHM1, is that this cell line is haploid, which suggests that any heterozygous variant calls are errors. A calling method producing fewer heterozygotes is in theory better. Meanwhile, to avoid overestimating a variant calling method with low sensitivity on heterozygotes, we also used NA12878 as a positive control.

The entire CHM1 data set (AC:SRP017546) gives over 100-fold coverage. We are using 6 SRA runs with the accessions ranged from SRR642636 to SRR642641. The 6 runs are from the same library, yielding about 65-fold coverage before the removal of potential PCR duplicates.

We acquired the NA12878 data set (AC:ERR194147) from the Illumina Platinum Genomes project. The library was constructed without PCR amplification. We are only using paired-end data, which yields about 55-fold coverage.

2.2 Alignment and post-alignment processing

We mapped the CHM1 reads with Bowtie2 Langmead and Salzberg (2012) and BWA-MEM Li (2013) and mapped the NA12878 reads with BWA Li and Durbin (2009) in addition to Bowtie2 and BWA-MEM. The detailed command lines can be found in Table 1. Except in Section 2.4, we mapped the reads to hs37d5, the reference genome used by the 1000 Genomes Project in the final phase.

After the initial alignment, we run Picard’s MarkDuplicates on both data sets. Picard identified 20% of CHM1 reads as PCR duplicates. We are only using 6 SRA runs with the accessions ranged from SRR642636 to SRR642641. The 6 runs are from the same library, yielding about 65-fold coverage before the removal of potential PCR duplicates.

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2.3 Calling SNPs and short INDELs

We called SNPs and short INDELs with FreeBayes Garrison and Marth (2012), GATK UnifiedGenotyper, Platypus, SAMtools and GATK HaplotypeCaller. The command lines can be found in Table 1. Additional details are as follows.

2.3.1 Resolving overlapping variants Platypus and SAMtools may produce many overlapping variants. To avoid overcounting variants, for two overlapping variants, we always keep the one with the higher variant quality. We repeated this procedure until no overlapping variants remain.

2.3.2 Re-calling genotypes Given the same genotype likelihood, different callers may produce different genotypes. For example, SAMtools estimates genotypes assuming the prior of seeing a heterozygote being $10^{-3}$, but GATK does not apply a prior. GATK is more likely to call a heterozygote than SAMtools. Genotype calling for a single sample is relatively simple. To avoid the subtle difference in this simple step complicating the final results, we re-call the genotypes from genotype likelihoods provided by the callers. We multiplied $10^{-3}$ to the likelihood of heterozygotes and then called the genotype with the maximum likelihood.

Platypus does not give genotype likelihoods for multi-allelic variants. We kept the reported genotypes in the VCF.

2.3.3 Decomposing complex variants Both FreeBayes and Platypus may report a variant composed of multiple SNPs and/or INDELs. We decomposed such variants into individual events such that the results are more comparable. FreeBayes uses a CIGAR string to describe how a complex variant is aligned to the reference. We extracted SNPs and INDELs from the CIGAR. Platypus does not report these numbers. DS has been identified to be an effective filter on cancer data (Roberts et al., 2013; Kim and Speed, 2013). We re-applied this filter to cancer data (Roberts et al., 2013; Kim and Speed, 2013) to cancer data (Roberts et al., 2013; Kim and Speed, 2013).

2.4 Variant filtering

All the callers used in this study come with filtering programs or a recommended set of filters. However, applying caller-specific filters may complicate comparison and obscure artifacts. We decided to choose several universal filters applicable to most callers:

1. Low-complexity filter (LC): filtering variants overlapping with low-complexity regions (LCRs) identified with the mdust program (http://bit.ly/mdust-LC), which is a stand-alone implementation of the DUST algorithm first used by BLAST. In GRCh37, 2.0% of A/C/G/T bases on autosomes are identified to be LCRs.

2. Maximum depth filter (DP): filtering sites covered by excessive number of reads. It should be noted that different callers may define the depth differently. For example, Platypus apparently only counts reads with unambiguous realignment. The read depth reported in the Platypus VCF is noticeably smaller in comparison to other callers.

3. Allele balance filter (AB): filtering sites where the fraction of non-reference reads is too low.

4. Double strand filter (DS): filtering variants if either the number of non-reference reads on the forward strand or on the reverse strand is below a certain threshold. This filter is not applicable to GATK calls as GATK does not report these numbers. DS has been identified to be an effective filter on cancer data (Roberts et al., 2013; Kim and Speed, 2013).

5. Fisher strand filter (FS): filtering sites where the numbers of reference/non-reference reads are highly correlated with the strands of the reads. More precisely, we counted the number of reference reads on the forward strand and on the reverse strand, and the number of non-reference reads on the forward and reverse strand.
four numbers, we constructed a 2-by-2 contingency table and used the
P-value from a Fisher’s exact test to evaluate the correlation.

6. Quality filter (QF): filtering sites with the reported variant quality
below a threshold.

Among these filters, LC is a regional filter and is entirely independent
of alignment and variant calling. Although DP is computed from called
variants, its effect is usually not greatly dependent on the mapper and the
caller, either. The remaining filters may be dependent of the error models
used by the callers. For example, SAMtools effectively gives a higher weight
on variants supported on both strands; FreeBayes seems to require a variant
to be supported 20% of reads covering the site. The optimal thresholds for
the AB, DS and FS filters are caller dependent.

2.5 Measuring accuracy
The CHM1 and the NA12878 data sets share many properties. They are both
sequenced with 100bp Illumina reads to a similar coverage after the removal
of PCR duplicates. The number of called variants per haplotype is also very
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3 RESULTS
When studying the effect of filters on variant calling, we initially
applied the filters independently on each call set. However, when
presenting the results in the following, we applied the filters in an
order, with a filter applied later depending on the filters applied
before it. We did this for clarity and to highlight filters having
major effects. Figure 1 overviews the breakdown of various filters
across multiple call sets. If we consider that there might be true
heterozygotes in CHM1 potentially due to somatic mutations, call
sets generally have an error rate approximately 1 in 100–200kb (i.e.
15000–30000 false heterozygotes per genome) after filtering.

3.1 Checking the ploidy of CHM1
Although the CHM1hTERT cell line is supposed to be haploid,
we may still see heterozygous variant calls potentially because: a)
the cell line is not truly haploid; b) there are somatic mutations
in the cell line; c) there are library construction and sequencing
effects (Robasky et al., 2014), which ought to be considered by the
calling algorithms; d) mapping or variant calling algorithms have
flaws. In this study, we are focusing on c) and d), but first we should
make sure heterozygotes resulted from a) and b) occur at a much
lower rate.

We note that if the sample submitted for sequencing is not haploid
either due to biological artifacts or massive somatic mutations,
a large number of heterozygotes should be evident from the
sequencing data and get called by all callers. In contrast, if
heterozygotes are mostly caused by sequencing errors or algorithm
artifacts, due to the differences in algorithm and error modeling,
callers will call a subset of errors with different characteristics,
which will result in low consistency between call sets. The small
call set intersection in Figure 1 suggests the latter is the case.

We also manually reviewed tens of heterozygotes called by
multiple callers both on the data in this study and on Illumina
data generated from other libraries (AC: SRR642626–SRR642635
and AC:SRR642750), which were mapped with the original BWA
algorithm (Li and Durbin, 2009) by the 1000 Genomes Project
analysis group. Reviewing the read evidence using an alignment
viewer, it appears that more than half of the SNPs are real. Most
of these SNPs have averaged read depth, non-overlap with known
segmental duplications (http://bit.ly/eelabdb) and are not associated
with known error-prone motifs in Illumina sequencing (Nakamura
et al., 2011). On the other hand, many INDELs in low-complexity
regions look like systematic errors called by all callers (see also
Section 3.2). We speculate there may be 5–20k heterozygotes

| Symbol | Algorithm | Version | Command line |
|--------|-----------|---------|--------------|
| bt2    | bowtie2   | 2.1.0   | bowtie2 \-x ref.fa -I read1.fq -2 read2.fq \-X 500 |
| bwa    | bwa-backtrack | 0.7.6 | bwa aln \-t read1.sai ref.fa read1.fq; bwa sampe ref.fa read1.sai read2.fq read1.fq read2.fq |
| mem    | bwa-mem   | 0.7.6   | bwa mem ref.fa read1.fq read2.fq |
| fb     | freebayes | 0.9.9   | freebayes \-I ref.fa aln.bam |
| st     | samtools  | 0.1.19  | samtools mpileup \-E ref.fa aln.bam \-bcftools view \-v - |
| ug     | UnifiedGenotyper | 2.7-4 | java \-jar GenomeAnalysisTK.jar \-T UnifiedGenotyper \-R ref.fa \-I aln.bam \-stand_call_conf 30 \-stand_emit_conf 10 \-glm BOTH |
| hc     | HaplotypeCaller | 2.7-4 | java \-jar GenomeAnalysisTK.jar \-T HaplotypeCaller \--genotyping_mode DISCOVERY \-R ref.fa \-I aln.bam \-stand_call_conf 30 \-stand_emit_conf 10 |
| pt     | Platypus  | 0.5.2   | Platypus.py callVariants \--filterDuplicates=1 \--bamFiles=aln.bam \--refFile=ref.fa |
Fig. 1. Effect of filters. Low-complexity filter (LC): not overlapping low-complexity regions identified by the DUST algorithm. Maximum-depth filter (DP): read depth below $d + 3\sqrt{d}$, where $d$ is the average read depth. Miscellaneous filter (misc) includes three filters: allele balance above 30%, variants supported by non-reference reads on both strands, and Fisher strand P-value is above 0.01. Filters are applied in the order of LC, DP, and misc, with DP applied to variants passing LC, and misc applied to variants passing both LC and DP. For each call set, the total height of the bar gives the number of raw variant calls with the reported quality in VCF no less than 30. Note that the Y-axes are scaled differently.

in CHM1 with strong alignment support from multiple Illumina libraries. It is hard to get a more accurate estimate or to further tell the sources of these heterozygotes with the data we are using. As we were writing up this work, Pacific Biosciences released deep resequencing data for the CHM1 cell line. It could be used to isolate errors caused by the Illumina sample preparation and sequencing. However, mapping and variant calling from PacBio human data is still in the early phase. We decided to leave out the comparison to the PacBio data for now.

Anyway, even if we assume the variant calls in the intersection are all present in the CHM1hTERT cell line, we should still be able to measure an error rate up to one error per 170kbp (=3 Gbp/17.7 k). Given that there are 10 times more raw heterozygous calls in NA12878 than CHM1 (Figure 1), it seems likely that CHM1 heterozygotes are likely errors from major sequencing/calling artifacts.

As a side technical note, we applied milder filters in Figure 2 in comparison to Figure 1. We found the intersection between call sets often becomes smaller with more stringent thresholds because stringent thresholds reduce the sensitivity in different aspects of call sets and amplify the subtle differences between calling algorithms. In addition, in Figure 2B, we were clustering INDELs within 20bp from each other. Increasing the distance threshold to 100bp only changed the numbers slightly.

3.2 The low-complexity filter
On CHM1, low-complexity regions (LCRs), 2% of the human genome, harbor 80–90% of heterozygous INDEL calls and up to 60% of heterozygous SNPs (Figure 1). Recall that if we let $N_{h}^{GL}$ be the number of CHM1 heterozygous INDELs in LCRs and $N_{d}^{GL}$ the number of NA12878 heterozygous INDELs in LCRs, $N_{h}^{GL} / N_{d}^{GL}$ estimates the FPR of heterozygotes. The FPR in LCRs is ranged from 10% to as high as 40% depending on call sets. With a similar estimator, the FPR of heterozygous INDELs outside LCRs is much lower, about 1–8% depending on call sets. We have also tried lobSTR (Gymrek et al., 2012). It called 65k heterozygous INDELs from microsatellites, still yielding a high FPR. To understand why
Artifacts in variant calling

3.2 Potential PCR amplification errors

PCR errors are known to be responsible for many INDEL errors in long homopolymer runs (1000 Genomes Project Consortium 2012). On CHM1, we have observed many apparent 1bp heterozygous INDELs (Figure 4) inserted to or deleted from long poly-A or poly-T runs, which may be due to PCR errors. Although most callers deploy advanced models for homopolymer INDELs, they are calling vastly different number of 1bp heterozygous INDELs. It is still not clear to us that we can model PCR errors well. Maybe the most effective solution is to avoid PCR in sample preparation.

Potential PCR errors are not the only error source. On the PCR-free NA12878 data the call set intersection in LCRs is noticeably smaller than in high-complexity regions (Figure 5), which suggests the presence of other error sources in LCRs. In addition, PCR errors introduced during sample preparation are believed to affect SNPs to a lesser extent. The small intersections between CHM1 heterozygous SNP call sets (Figure 2), and between PCR-free SNP call sets in LCRs (Figure 5) should be caused by other types of errors.

3.2.2 Realignment errors

When mapping a read to the reference genome, a read mapper chooses the optimal pairwise alignment for each read independent of others. For reads mapped to the same region, the combination of optimal pairwise alignments does not always yield the optimal multi-alignment of reads. If a variant caller simply trusts the suboptimal multi-alignment, it may produce false variants or genotypes (Figure 3). Therefore, more recent variant callers, including HaplotypeCaller, Platypus and FreeBayes in this study, heavily rely on realignment for both SNP and INDEL calling.

However, with our manual review, we found that variant callers often failed to produce the optimal realignment in LCRs. About 50–70% of the reviewed >1bp heterozygous INDELs from CHM1 can be corrected away with better realignment. Without the thorough understanding of the very details of the realignment process, we are unable to explain why the callers fail even on some obvious cases. Nonetheless, as we can often manually derive a better multi-alignment, it is possible that a good realignment algorithm may replace our manual work and achieve higher accuracy than all the tools in our evaluation.

In the process of manual review, we found local assembly with fermi is frequently more effective than the INDEL callers, which may be due to the independence of the reference sequence, the requirement of long-range consistency and the more powerful topology based error cleaning (Zerbino and Birney 2008). Some difficult errors such as Figure 6 are trivial to resolve with local assembly.

3.3 The maximum read depth filter

3.3.1 The effectiveness of the max depth filter

Other filters require a threshold on a single value. To study which filter is more effective, we used a ROC-like plot, Figure 5. In this figure, the X-axis, the number of heterozygous SNPs in CHM1, is proportional to the false positive rate; the Y-axis, the difference of the number of heterozygous SNPs between NA12878 and CHM1, serves as a proxy to the sensitivity. Similar to a standard ROC plot, a curve closer to the top left corner implies a better classifier of errors.

Figure 5 implies that the maximum depth filter is the most effective against false heterozygotes, especially those found from the BWA-MEM alignment. On our data with depth d ≈ 50, a maximum depth threshold between d + 3√d and d + 4√d removes many false positives with little effect on the sensitivity. These false positives are mostly caused by copy number variations (CNVs) or paralogous sequences not present in the human reference genome.
Fig. 4. Example of misalignment around chr1:2608841 in CHM1. The truth allele is derived from local assembly. Three erroneous read alignments and their correct alignments are shown below it. Each of the three reads is an exact substring of the truth allele, but their alignments are different. The first read ‘errRead1’ is aligned without gaps as the 3’-end of the read is a substring of the 18bp deletion. Read ‘errRead2’ is aligned with a 6bp insertion as this alignment is better than having two long deletions. Read ‘errRead3’ is also aligned without gaps but with seven mismatches. It is possible for an aligner to find its correct alignment given a small gap extension penalty. On this example, Bowtie2 did not align any reads with gaps. BW A-MEM aligned four reads correctly. Except HaplotypeCaller which locally assembled reads, other callers all called multiple heterozygotes around this region.

Fig. 5. Effect of filters after removing variants in low-complexity regions. Each filter is associated with one value. For each filter, the number of heterozygous SNPs called from CHM1 and NA12878 are counted accumulatively from the most stringent threshold on the filter value to the most relax threshold. Thresholds are chosen such that they approximately evenly divide variants into 100 bins. Each chosen threshold yields a point in the plot. An arrow points to a point on the DP curve when the corresponding read depth is right above \( d + 4\sqrt{d} \), where \( d \) is the mean read depth across called variants.

3.3.2 The difference between Bowtie2 and BWA-MEM alignment

It is clear that Bowtie2 is less affected by the presence of CNVs and an incomplete genome (Figure 4). With manual review, it seems to us that in comparison to BWA-MEM, Bowtie2 tends to give the same alignment a lower mapping quality when the read has other suboptimal hits. At the same time, missing paralogous sequences from the reference genome are often associated with existing segmental duplications in the reference genome. Therefore, Bowtie2 is more likely to correctly give a low mapping quality to a read from these paralogous sequences. As variant callers usually distrust mismatches on alignments with low mapping quality, their calls from the Bowtie2 alignment are less susceptible to CNVs or an incomplete reference genome.

However, being conservative on the mapping quality estimate may lead to more false negatives. For example, we found a read pair having one mismatch around 13.7Mbp in chr1 but two mismatches around 13.5Mbp. Both Bowtie2 and BWA-MEM mapped the ends of the pair at the same positions. Bowtie2 gives the pair a mapping quality 6, while BWA-MEM gives a mapping quality 27. The similar scenario happens to the other reads mapped to this region. As a result, a SNP is called from the BWA-MEM alignment, but not from the Bowtie2 alignment. Variants callers usually call more variants from the BWA-MEM alignment (Figure 1), many of which are located in segmental duplications.

Another difference, not relevant to the mapping quality, comes from the alignment around long INDELs. HaplotypeCaller always called more ≥15bp INDELs from the BWA-MEM alignment (data not shown). Other callers made three times as many ≥15bp deletion calls from the BWA-MEM alignment, either in LCRs or not, and called 40% more insertions outside LCRs. Interestingly, except HaplotypeCaller, others called more ≥15bp insertions from the Bowtie2 alignment in LCRs instead. We have not found a good explanation to the apparently conflictive observations.

3.3.3 An alternative to the maximum depth filter

While the maximum depth filter is effective against false heterozygotes, it is only applicable to high-coverage data with uniform read depth. It does not work with exome sequencing data, or is not powerful on data with shallow coverage.

To overcome the limitation, we derived an alternative filter. We obtained unfiltered SAMtools SNP calls from the 1000 Genomes Project and computed the inbreeding coefficient and the Hardy-Weinberg P-value using genotype likelihoods (Li, 2011b). We extracted sites satisfying: 1) the reported read depth above 25.000; 2) the inbreeding coefficient less than zero; 3) the P-value below 10⁻¹⁰. We then clustered the sites within 10kb into regions. These regions are susceptible to common CNVs or artifacts in the
reference genome. We call this filter as the Hardy-Weinberg filter, or HW in brief.

On CHM1, the HW filter is almost as effective as the maximum depth filter. It could be a valid alternative when the maximum depth filter cannot be applied. However, the derivation of the HW filter requires multiple thresholds and depends on populations, the mapper (BWA) and the caller (SAMtools). Therefore, we decided to use the much simpler maximum depth filter here.

3.4 Other filters

The remaining filters, including AB, DS, FS and QU (Section 2.3), can filter additional false heterozygous called from CHM1, but their effectiveness varies with call sets. It is also difficult to find the optimal thresholds on these filters as they affect both the false negative rate and the false positive rate. In the end, we arbitrarily chose reasonable thresholds based on the ROC-like curves (Figure 5), which may not be optimal for all call sets.

3.5 Effect of PCR duplicates

20% of CHM1 data are discarded in our analysis due to PCR duplicates. We have also tried variant calling with mem:hc without the MarkDuplicates step. Before filtering, this approach yields 3% more heterozygous SNPs and 12% more heterozygous INDELs, suggesting INDELs are more susceptible to PCR artifacts than SNPs. After filtering, the total numbers of SNPs and INDELs are about the same with or without duplicates.

We have also mapped the CHM1 reads to the GRCh37 and GRCh38 primary assemblies and called variants. The number of homozygous non-LC SNPs called from each reference is close: 2.408, 2.405 and 2.412 million from GRCh37, hs37d5 and GRCh38, respectively. However, the numbers of heterozygous SNPs/INDELs are distinct (Figure 6). We called twice as many heterozygotes from GRCh37 in comparison to hs37d5. This indicates that the 35.4Mb decoy sequences indeed attracted many mismapped reads and consequently improves the variant calls in chromosomal regions. GRCh38 further resolves 39.8k (=36.9k+2909) heterozygotes called from hs37d5. However, it also retains 36.8k heterozygotes called from GRCh37 but not from hs37d5. Intriguingly, GRCh38 further adds 24.6k autosomal heterozygotes not called from GRCh37 or hs37d5. We are unclear of the source of these false heterozygous SNPs. In general, we conclude that hs37d5 and GRCh38 are more complete than GRCh37.

4 DISCUSSIONS AND CONCLUSIONS

A distinct feature of our works is the use of a haploid human sample, CHM1, from which heterozygous calls are supposed to be errors. This allows us to unbiasedly investigate the causal artifacts and to experiment effective filters with the diploid NA12878 data set as a positive control.

When we called SNPs and INDELs from CHM1, we were surprised to find 10% of raw variant calls were heterozygotes. Honestly, our immediate reaction was that CHM1 was not truly haploid. However, after careful analysis, we have convinced ourselves that the heterozygosity of CHM1 should be of an order of magnitude lower than the raw error rate of variant calling. The vast majority of heterozygotes are calling errors. In the raw call set, we usually see an error per 10–15kb.

It was also to our surprise that the low-complexity filter is the most effective against false heterozygotes, especially short INDELs. Although we knew that INDEL errors may be introduced by PCR during sample preparation, we underestimated its substantial effect. We were also unaware that realignment of INDELs in LCRs remains a great challenge even after the many existing efforts in this direction (Homer and Nelson, 2010; Li, 2011; Albers et al., 2011; Narzisi et al., 2014). Without the suggestion from Peter Sudmant (personal communication), we would not have tried this filter.

Before we understand and resolve the issues in variant calling in LCRs, it might be better to filter out all variants overlapping these regions. Although over 50% of single-sample INDEL calls fall in LCRs (Figure 7), only 1.25% of autosomal INDELs in the ClinVar database overlap with LCRs – most INDELs in LCRs have unknown clinical functionality. For certain applications, it might be safe to drop or downweigh these difficult calls.

Outside LCRs, different call sets usually agree well with each other if the same set of filters are applied (Figure 9). Based on Figure 1, we estimate that a caller usually makes a wrong call per 100–200kb without significant compromise on the sensitivity, similar to the previous estimates (Bentley et al., 2008; Nickels et al., 2012). Many of these errors are likely to be systematic. In the context of somatic or de novo mutation discovery by sample contrast, systematic errors will appear in all samples. They will not lead to false mutation calls, fortunately.
A simple method to improve the variant accuracy is to use two distinct pipelines, take the intersection of the raw calls and then apply caller-oblivious filters to derive the final call set. As callers agree well on post-filtered sites (Figure 3) but badly on false positives (Figure 2), we should be able to remove most errors without much hit to the sensitivity. Such a consensus approach has been applied to cancer data with limited success (Löwer et al., 2012; Goode et al., 2013). Without subclonal mutations, it should be much more effective on the variant discovery from normal samples.

Finally, the advances in sequencing technologies lead the development of algorithms. We are heavily relying on mapping based variant calling because with very short reads or at low coverage, the traditional assembly-and-mapping approach would not work. With increased read lengths and decreased sequencing cost, we might go back to de novo assembly. An assembly does not only encode small variants, but also retains large-scale structural variations and is free of the artifacts in the reference genome. Another possible direction which we mentioned four years ago (Li et al., 2010) is to map sequence reads to the ensemble of multiple genomes. Recently, there have been significant progress towards this goal (Sireń et al., 2010; Paten et al., 2014), but a practical solution is yet to be concluded.

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