A three-caller pipeline for variant analysis of cancer whole-exome sequencing data

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Abstract. Rapid advancements in next generation sequencing (NGS) technologies, coupled with the dramatic decrease in cost, have made NGS one of the leading approaches applied in cancer research. In addition, it is increasingly used in clinical practice for cancer diagnosis and treatment. Somatic (cancer-only) single nucleotide variants and small insertions and deletions (indels) are the simplest classes of mutation, however, their identification in whole exome sequencing data is complicated by germline polymorphisms, tumor heterogeneity and errors in sequencing and analysis. An increasing number of software and methodological guidelines are being published for the analysis of sequencing data. Usually, the algorithms of MuTect, VarScan and Genome Analysis Toolkit are applied to identify the variants. However, one of these algorithms alone results in incomplete genomic information. To address this issue, the present study developed a systematic pipeline for analyzing the whole exome sequencing data of hepatocellular carcinoma (HCC) using a combination of the three algorithms, named the three-caller pipeline. Application of the three-caller pipeline to the whole exome data of HCC, improved the detection of true positive mutations and a total of 75 tumor-specific somatic variants were identified. Functional enrichment analysis revealed the mutations in the genes encoding cell adhesion and regulation of Ras GTPase activity. This pipeline provides an effective approach to identify variants from NGS data for subsequent functional analyses.

Introduction

It is well established that tumorigenesis is attributed to chromosomal instability or accumulated genetic changes, including structure variations, genetic copy number variants, single nucleotide variants (SNVs) and small insertions and deletions (indels) (1-3). Somatic mutations are defined by mutations that are absent in corresponding adjacent tissues; however, they are present in all tumors (4). Somatic mutation calling is a critical step for cancer genome characterization and clinical genotyping. Next-generation sequencing (NGS) has become a popular strategy for genotyping, enabling more precise mutation detection compared with traditional methods due to its high resolution and high throughput. Whole-genome sequencing reveals overall genetic information about the variants, whereas whole-exome sequencing (WES) with effective strategy only points economically at coding regions and is currently offered by more laboratories (5). WES of tumor samples and matched normal controls has the potential to rapidly identify protein-altering mutations across hundreds of patients, potentially enabling the discovery of recurrent events that drive tumor development and growth. Identification of somatic mutations from WES data is an increasingly common technique in the study of cancer genomics, and a large number of somatic alterations have been identified by WES in extensive tumor types (6-9). The most prevalent mutations observed are in the p53 tumor suppressor gene (TP53), Wnt/β-catenin signaling pathway regulatory genes (catenin β1 and AXIN 1), chromatin remodeling complex components [AT-rich interactive domain (ARID) 2 and ARID1A], Janus kinase (JAK)/signal transducer and activator of transcription pathway-regulated JAK1, as well as hepatitis B (HBV) integrations into myeloid/lymphoid or mixed-lineage leukemia 4, telomerase reverse transcriptase and cyclin E1 (10,11).

The calling of accurate somatic mutations using WES data remains one of the major challenges in cancer genomics due to various sources of errors, including artifacts occurring during polymerase chain reaction (PCR) amplification or targeted capture, machine sequencing errors and incorrect local alignments of reads (12). Tumor heterogeneity and normal tissue contamination generate additional difficulties for the identification of tumor-specific somatic mutations (12,13). In recent years, several methods have been developed to improve the accuracy of somatic mutation calling. Despite the variations
in the methodology of somatic mutation algorithms, the aim of each program is to identify tumor-specific variants by comparing the tumor variant data with the dbSNP of paired adjacent tissue and germline variant data in the same patient. Currently the most popular computational algorithms are MuTect (14), VarScan2 (15) and Genome Analysis Toolkit (GATK) (16). GATK calculates the variants in tumors and adjacent tissues separately, and then subtracts the variants identified in the adjacent tissues from those in the tumors. MuTect and VarScan2 directly compare the tumor tissues with the adjacent tissues at each mutation point, which in some cases improves the accuracy of variant calling. MuTect detects somatic mutation sensitively with a Bayesian model at low allele-fractions, whereas VarScan2 applies a powerful heuristic/statistic approach to identify high-quality variants (12). However, it is unclear which is the best strategy for identifying and accurately calling genome variations as well as how well these different tools improve the true positive mutations when they are combined.

The present study integrated the resources of different somatic mutation algorithms and optimized their own parameters in order to identify novel and recurrent mutations more effectively and faster. The present study used one case of hepatocellular carcinoma (HCC) to explain the whole-exome analysis pipeline and identify the key somatic mutations of HCC.

Materials and methods

Patient. A punctured HCC tumor and paired adjacent tissue was obtained from a patient (57 years, male) at the Youan Hospital, Capital Medical University of China (Beijing, China) and complied with the principles of The Declaration of Helsinki. The patient was infected with HBV and received no radiation and chemotherapy prior to radiofrequency ablation.

NGS platforms. The DNA was extracted using an E.Z.N.A.® Tissue DNA Kit (Omega Bio-Tek, Inc., Norcross, GA, USA) and the extracted DNA was captured using Agilent Human All Exon 50 M kit (Agilent Technologies, Inc., Santa Clara, CA, USA) following the protocols recommended by the manufacturer. Sequencing machines generated a large volume of data at a rapid speed by sequencing paired-end DNA fragments in parallel using Illumina His-seq2,000 (Illumina, Inc., San Diego, CA, USA) (17,18). Following a series of library construction and actual sequencing, a large quantity of raw data was produced.

Quality evaluation of the raw reads. Raw reads generated by a sequenator are usually affected by adverse factors, including adaptor contamination, poor base sequence quality and guanine-cytosine (GC) bias (19). Once the raw data was obtained, the quality of raw reads was assessed and the adaptor was clipped using fastq-mcf (version 1.04.636; www.github.com/ExpressionAnalysis/ea-utils/blob/wiki/FastqMcf.md). The sequencing data was then processed using the FastQC tool (http://www.bioinformaticsbabraham.ac.uk/projects/fastqc/) to analyze the distribution of base GC content and sequence quality scores.

Alignment and duplicated PCR removal. Following the quality control analyses, the processed reads were aligned to an established reference genome (version hg19), which was provided by the University of California Santa Cruz (Santa Cruz, CA, USA) (20). Millions of short reads were aligned efficiently to the reference genome using Burrows-Wheeler Aligner (BWA) software with default parameters, which were based on the Burrows-Wheeler transform (21). The aligned reads were then stored in BAM file (.bam) using samtools software (22), which was able to sort and index the BAM file to save space and help subsequent process. For the assembled genome data, the picard tool (http://picard.sourceforge.net/index.shtml) was combined with bamtools to filter out the mismatching and inappropriate reads. In addition, picard removed the read duplicates derived from library PCR. The data distribution and reads coverage were then evaluated using the CalculateHsMetrics package. Recalibration and realignment were performed using GATK (version 2.8; Broad Institute, Cambridge, MA, USA; www.broadinstitute.org/gatk/). Finally, the resulting data were used for further variation identification.

Variant identification. A key step in the analysis of cancer exome sequencing data is the identification of variants. The depth of sequence coverage determines the choice of somatic mutation algorithms used for identification of variants mutation. The different identification abilities in different allele frequencies of GATK (version 2.8.1), MuTect (version 1.1.4; Broad Institute; http://www.broadinstitute.org/cancer/gma/mutect), and VarScan (version 2.3.6; http://varscan.sourceforge.net/), and the joint analysis strategy by combining the three softwares (the three-caller pipeline approach), were taken into consideration when identifying somatic mutations.

Variant annotation. Oncotator (http://portals.broadinstitute.org/oncotator/) was used to annotate the screened variations (23). All of the candidate mutations were validated visually using the Integrated Genomics Viewer (IGV) (24) and were confirmed using Sanger sequencing in paired samples. The tools, Polyphen-2 (www.genetics.bwh.harvard.edu/pph2/index.shtml) and scale-invariant feature transform (SIFT; www.sift.jcvi.org/), were integrated to predict whether mutations affected protein function based on the structure and function of the protein, and the conservation of amino acid residues in different species sequences.

Gene functional enrichment analysis. The gene sets screened were used for functional annotation analysis by the Database for Annotation, Visualization and Integrate Discovery software (25), which consists of the Kyoto Encyclopedia of Genes and Genomes and Gene Ontology database. The significance of gene groups enrichment was defined by a modified Fisher's exact test and P<0.05 was considered to indicate a statistically significant difference.

Results

Establishment of three-caller and HCC data analysis. WES was analyzed in one HCC tumor and paired adjacent tissues with the three-caller approach. The present study acquired
96.30X and 79.18X coverages for the tumor and paired adjacent tissues, respectively, in all of the targeted exonic regions, with 93.4% of the base targeted at 20-fold and ≥99.1% bases by a depth of at least two times. To identify the somatic mutations, a flow chart was created with the following steps: i) Quality evaluation of the raw reads; ii) reads map to a reference genome; iii) somatic mutation identification with the three-caller approach; iv) variant annotation; v) data visualization; and vi) pathway analysis (Fig. 1).

Detecting SNVs in a HCC sample. Variant filtering was performed by GATK with the following filter parameters: Low coverage (DP <5), low quality (QUAL >30.0 and QUAL <5.0), very low quality (QUAL <30), hard to validate [MQ0 ≥4 and MQ0/(1.0*DP)>0.1] and quality-by-depth (QD <1.5). The exome data from the samples were calculated by running these parameters and reserved in a VCF file. GATK was primarily used for identifying somatic mutations in the sequencing data, including SNVs and indels.

In order to identify the low allelic-fraction mutations, MuTect was used to generate more performance in low coverage (12). To illustrate how high the sensitivity was based on allele fraction and sequencing depth, a strategy was established based on the published data to analyze the data (14). As shown in Fig. 2, the sensitivity of mutation was detected by MuTect approaching >90% at allele frequency 10% with >80X sequencing depth and 80% at allele frequency 5% with >80X.

VarScan outperformed the other tools at higher allelic fraction. A threshold of 6X for tumor and 8X for normal was set, with ≥20% variation frequency. Subsequently, the present study preferentially analyzed 20X coverage in the tumor, including alternated variation accounting for 10X coverage, to eliminate false positives.

The present study proposed 75 candidate somatic variants through the three-algorithm strategy (Fig. 3), including 50 nonsynonymous mutations, 2 nonsense mutations,
Table I. Selected somatic mutations predicted by Polyphen to affect protein function.

| Hugo symbol | Amino acid change | SIFT   | SIFT score | Polyphen      | Polyphen score |
|-------------|-------------------|--------|------------|---------------|----------------|
| CSMD1       | Q2192R            | Damaging | 0.04       | Probably damaging | 0.973         |
| FREM1       | H822Q             | Damaging | 0.01       | Probably damaging | 0.972         |
| GP5         | I230N             | Damaging | 0.06       | Probably damaging | 0.997         |
| KCNA1       | E422K             | Tolerated | 0.13       | Benign        | 0.011         |
| CDC7        | P94Q              | Damaging | 0.02       | Probably damaging | 0.998         |
| DMBT1       | R2343W            | Damaging | 0.08       | Benign        | 0.015         |
| FAT2        | V3602I            | Tolerated | 0.13       | Benign        | 0.011         |
| C10orf90    | R188W             | Tolerated | 0.08       | Benign        | 0.015         |

CSMD1, CUB and sushi multiple domains 1; FREM1, FRAS1-related extracellular matrix 1; GP5, glycoprotein V platelet; KCNA1, potassium voltage-gated channel subfamily A member 1; CDC7, cell division cycle 7; DMBT1, deleted in malignant brain tumors 1; FAT2, FAT atypical cadherin 2; C10orf90, chromosome 10 open reading frame 90; SIFT, scale-invariant feature transform.

Table II. Functional categories of the tumor-specific mutation.

| Biological process                        | Count | P-value | Genes                                                                 | Fold enrichment |
|-------------------------------------------|-------|---------|----------------------------------------------------------------------|-----------------|
| Cell adhesion                             | 8     | 0.0089  | GP5, LGALS3BP, FREM1, FAT2, FCGBP, COL5A3, PCDHGB4, MUC16            | 3.29            |
| Regulation of Ras GTPase activity         | 3     | 0.0487  | TBC1D3, AGAP3, TBC1D3B, AGAP4                                        | 8.3             |

GP5, glycoprotein V platelet; LGALS3BP, galectin 3 binding protein; FREM1, FRAS1-related extracellular matrix 1; FAT2, FAT atypical cadherin 2; FCGBP, Fc fragment of IgG binding protein; COL5A3, collagen type V α3 chain; PCDHGB4, protocadherin γ subfamily B, 4; MUC16, mucin 16; TBC1D3, TBC1 domain family member; AGAP, ArfGAP with GTPase domain, ankyrin repeat and PH domain.

Figure 4. Identification of MUC16 variants in a pair of HCC samples. The figure depicts the exome sequencing projects of HCC tumor and paired adjacent tissues. The blue letter C indicates the presence of a non-reference allele, and thus a point mutation (T>C) at position_9056725 in MUC16. MUC16, mucin 16; HCC, hepatocellular carcinoma.
20 synonymous mutations and 3 indels. The nonsynonymous to synonymous somatic SNV ratio was 2.5.

Analysis of somatic mutations. The predictive impact of amino acid substitution on functional evidence was analyzed using PolyPhen-2/SIFT (Table I). The P94Q mutation was predicted to affect the protein function of cell division cycle 7 protein, which may be associated with neoplastic transformation of some tumors and affect protein serine/threonine kinase activity. All of the putative somatic mutations were validated manually using IGV. The T>C transversion at position_9056725 in mucin 16 (MUC16) was identified (Fig. 4), which was then validated by Sanger sequencing.

Pathway analysis. The 75 genes with tumor-specific mutations demonstrated significant functional enrichment of cell adhesion and regulation of Ras GTPase activity (P<0.05; Table II). Notably, the genes encoding cell adhesion demonstrated the most prevalent enrichment (P=0.0089), indicating that the enriched mutations of cell adhesion genes may serve pivotal roles in HCC development.

Discussion

WES technologies have provided extensive profiles of genomic mutations in cancers, however, how to process the generated dataset effectively for downstream analyses, remains a problem. Currently the accuracy of variant calling is still influenced by a number of factors. Firstly, low specificity and sensitivity of the existing high-throughput sequencing may prevent the generation of accurate mutation profiles (26). Secondly, the BWA algorithms may produce incorrect base alignment. Finally, the three algorithm tools, MuTect, VarScan and GATK, used for identifying variants, present their respective limitations. GATK is a semi-automated algorithm that calculates somatic variants. VarScan identifies the most high-quality SNVs preferentially, while MuTect outperforms in low-quality ones. Some true SNVs are hard to differentiate due to a number of factors including clonal heterogeneity, strand bias, low allele frequencies, tumor contamination, high GC content of genomic regions, sequencing errors and non-specificities in short read mapping (12).

Comparisons between SNVs calls analyzed with GATK, MuTect and VarScan, revealed that only a few of the SNVs were called by more than one of the tools (Fig. 3), thus it was difficult to select candidate SNVs for further validation. The disagreement was partially due to prior assumptions underlying each algorithm and different error models. Therefore, further development of more significant and accurate calling algorithms was required (27), however, combining MuTect/GATK with VarScan produced more accurate SNVs. In light of these limitations in genomic studies, the three-caller strategy was designed to obtain accurate mutation information for clinical assessment.

The present study integrated different software programs to form a modular pipeline for processing somatic SNVs and indels. A series of software was used to perform data alignment, data filtering, reducing duplicate and realignment, as well as recalibrating through java. In the study of HCC, WES analysis started with the acquisition of raw data to select several candidate genes, which alluded to the potential effect of cancer-associated somatic mutations on tumor progression. The mutation set-based analysis revealed a number of potential somatic events in HCC, including in CUB and sushi multiple domains 1, FRAS1-related extracellular matrix 1 and MUC16 genes. The mutations at different base positions of the same gene or different genes may lead to disparate functions such as activation and inactivation mutations. This may influence their physicochemical properties and structure in comparison with wild-type proteins. Functional enrichment analysis revealed the biological process enrichment of cancer-specific mutations, including cell adhesion and regulation of Ras GTPase activity. Experiments are required to validate the variants which may affect interactions with other proteins and disorder crucial signaling pathways (28).

In conclusion, the pipeline for HCC exome sequencing data analysis demonstrated in the present study provided a convenient strategy to identify the potentially functional tumor-specific mutations, which may support our understanding of the underlying mechanisms of HCC development.

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References

1. Lengauer C, Kinzler KW and Vogelstein B: Genetic instabilities in human cancers. Nature 396: 643-649, 1998.
2. Bass AJ, Lawrence MS, Brace LE, Ramos AH, Drier Y, Cibulskis K, Sougnez C, Voet D, Saksena G, Sivachenko A et al: Genomic sequencing of colorectal adenocarcinomas identifies a recurrent VTI1A-TCF7L2 fusion. Nat Genet 43: 964-968, 2011.
3. Chapman MA, Lawrence MS, Keats JJ, Cibulskis K, Sougnez C, Schinzel AC, Harvick CL, Brunet JP, Ahmann GJ, Adli M et al: Initial genome sequencing and analysis of multiple myeloma. Nature 471: 467-472, 2011.
4. Jia D, Dong R, Jing Y, Xu D, Wang Q, Chen L, Li Q, Huang Y, Zhang Y, Zhang Z, et al: Exome sequencing of hepatoblastoma reveals novel mutations and cancer genes in the Wnt pathway and ubiquitin ligase complex. Hepatology 60: 1686-1696, 2014.
5. Biesecker LG and Green RC: Diagnostic clinical genome and exome sequencing. N Engl J Med 367: 1170, 2015.
6. Cancer Genome Atlas Research Network, Hammerman PS, Lawrence MS, Voet D, Jing R, Cibulskis K, Sivachenko A, Stojanov P, McKenna A, Lander ES, et al: Comprehensive genomic characterization of squamous cell lung cancers. Nature 489: 519-525, 2012.
7. Cancer Genome Atlas Network, Muzny DM, Bainbridge MN, Chang K, Dinl HH, Drummond IA, Fowler G, Kobar CL, Lewis LR, Morgan MB, et al: Comprehensive molecular characterization of squamous cell lung cancers. Nature 489: 519-525, 2012.
8. Cancer Genome Atlas Network, Koboldt DC, Fulton RS, McLellan MD, Schmidt H, Kaliski-Verzler J, McMichael JP, Fulton LL, Dooling DJ, Ding L, et al: Comprehensive molecular portraits of human breast tumours. Nature 490: 61-70, 2012.
9. Litchfield KD, Summersgill B, Yost S, Sultana R, Labrecque K, Dudakia D, Renwick A, Seal S, Al-Saad R, Broderick P, et al: Whole-exome sequencing reveals the mutational spectrum of testicular germ cell tumours. Nat Commun 6: 5973, 2015.
10. Zhang Z: Genomic landscape of liver cancer. Nat Genet 44: 1075-1077, 2012.
11. Kan Z, Zheng H, Liu X, Li S, Barber TD, Gong Z, Gao H, Hao K, Willard MD, Xu J, et al: Whole-genome sequencing identifies recurrent mutations in hepatocellular carcinoma. Genome Res 23: 1422-1433, 2013.
12. Wang Q, Jia P, Li F, Chen H, Ji H, Hucks D, Dahlman KB, Pao W and Zhao Z: Detecting somatic point mutations in cancer genome sequencing data: A comparison of mutation callers. Genome Med 5: 91, 2013.

13. Gerlinger M, Rowan AJ, Horswell S, Larkin J, Endesfelder D, Gronroos E, Martinez P, Matthews N, Stewart A, Tarpey P, et al: Intratumor heterogeneity and branched evolution revealed by multiregion sequencing. N Engl J Med 366: 883-892, 2012.

14. Cibulskis K, Lawrence MS, Carter SL, Sivachenko A, Jaffe D, Sougnez C, Gabriel S, Meyerson M, Lander ES and Getz G: Sensitive detection of somatic point mutations in impure and heterogeneous cancer samples. Nat Biotechnol 31: 213-219, 2013.

15. Koboldt DC, Zhang Q, Larson DE, Shen D, McLellan MD, Lin L, Miller CA, Mardis ER, Ding L and Wilson RK: Varscan 2: Somatic mutation and copy number alteration discovery in cancer by exome sequencing. Genome Res 22: 568-576, 2012.

16. McKenna A, Hanna M, Banks E, Sivachenko A, Cibulskis K, Kernytsky A, Garimella K, Altshuler D, Gabriel S, Daly M and DePristo MA: The genome analysis toolkit: A mapreduce framework for analyzing next-generation DNA sequencing data. Genome Res 20: 1297-1303, 2010.

17. Mardis ER: Next-generation DNA sequencing methods. Annu Rev Genomics Hum Genet 9: 387-402, 2008.

18. Metzker ML: Sequencing technologies-the next generation. Nat Rev Genet 11: 31-46, 2010.

19. Dohm JC, Lottaz C, Borodina T and Himmelbauer H: Substantial biases in ultra-short read data sets from high-throughput DNA sequencing. Nucleic Acids Res 36: e105, 2008.

20. Nielsen R, Paul JS, Albrechtsen A and Song YS: Genotype and SNP calling from next-generation sequencing data. Nat Rev Genet 12: 443-451, 2011.

21. Li H and Durbin R: Fast and accurate long-read alignment with burrows-wheeler transform. Bioinformatics 26: 589-595, 2010.

22. Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G and Durbin R: 1000 Genome Project Data Processing Subgroup: The sequence alignment/map format and SAMtools. Bioinformatics 25: 2078-2079, 2009.

23. Ramos AH, Lichtenstein L, Gupta M, Lawrence MS, Pugh TJ, Saksena G, Meyerson M and Getz G: Oncotator: Cancer variant annotation tool. Hum Mutat 36: E2423-E2429, 2015.

24. Robinson JT, Thorvaldsdóttir H, Winckler W, Guttman M, Lander ES, Getz G and Mardis ER: Integrative genomics viewer. Nat Biotechnol 29: 24-26, 2011.

25. Huang da W, Sherman BT and Lempicki RA: Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. Nat Protoc 4: 44-57, 2009.

26. Totoki Y, Tatsuno K, Yamamoto S, Arai Y, Hosoda F, Ishikawa S, Tsutsumi S, Sonoda K, Totsuka H, Shirakihara T, et al: High-resolution characterization of a hepatocellular carcinoma genome. Nat Genet 43: 464-469, 2011.

27. Lawrence MS, Stojanov P, Polak P, Kryukov GV, Cibulskis K, Sivachenko A, Carter SL, Stewart C, Mermel CH, Roberts SA, et al: Mutational heterogeneity in cancer and the search for new cancer-associated genes. Nature 499: 214-218, 2013.

28. Kwon SM, Cho H, Choi JH, Jee BA, Jo Y and Woo HG: Perspectives of integrative cancer genomics in next generation sequencing era. Genomics Inform 10: 69-73, 2012.