HomSI: a homozygous stretch identifier from next-generation sequencing data

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ABSTRACT

Summary: In consanguineous families, as a result of inheriting the same genomic segments through both parents, the individuals have stretches of their genomes that are homozygous. This situation leads to the prevalence of recessive diseases among the members of these families. Homozygosity mapping is based on this observation, and in consanguineous families, several recessive disease genes have been discovered with the help of this technique. The researchers typically use single nucleotide polymorphism arrays to determine the homozygous regions and then search for the disease gene by sequencing the genomes within this candidate disease loci. Recently, the advent of next-generation sequencing enables the concurrent identification of homozygous regions and the detection of mutations relevant for diagnosis, using data from a single sequencing experiment. In this respect, we have developed a novel tool that identifies homozygous regions using deep sequence data. Using *.vcf (variant call format) files as an input file, our program identifies the majority of homozygous regions found by microarray single nucleotide polymorphism genotype data.

Availability and implementation: HomSI software is freely available at www.igbam.bilgem.tubitak.gov.tr/softwares/HomSI, with an online manual.

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Supplementary information: Supplementary data are available at Bioinformatics online.

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1 INTRODUCTION

In communities with a high level of consanguineous marriage, the diagnosis of a recessive genetic disorder offers a unique advantage for positional cloning of rare diseases. In such an isolated inbred human population, several individuals may inherit a variation of an ancestor, and the offsprings born of consanguineous union will have a high probability of inheriting two copies of the mutated chromosomal segment and thus expressing the disease. In this respect, homozygosity mapping is an effective approach to identify potential disease loci. Because Lander and Botstein successfully used this technique in 1987 (Lander and Botstein, 1987), both the experimental and the computational methods that are used to generate and analyze relevant datasets have undergone diversification and refinement. Originally, to detect homozygous regions, individuals were genotyped with panels of highly polymorphic short tandem repeat (or microsatellite) markers, typically at genomic intervals of 10–12 cM. With the availability of single nucleotide polymorphism (SNP) microarrays, short tandem repeat microsatellites were later replaced by SNP arrays to survey the genome and to identify large stretches of homozygosity. The overlapping regions among the homozygous stretches of affected individuals are called runs of shared homozygosity (ROSHs), and these regions are expected to contain the disease gene. For this purpose, several computational tools have been developed (Amir et al., 2010; Carr et al., 2006; Kayserili et al., 2009; Papic et al., 2011; Seelow et al., 2009; Uz et al., 2010; Zhang et al., 2011).

Even though the identification of recessive disease loci (homozygous regions) is accomplished using the aforementioned SNP array-based techniques, the next step to detect pathogenic sequence variant is sequencing. Until recently, only the candidate genes in these loci are sequenced. If this disease locus is a large interval or if it includes several genes, the determination of disease causing mutations by Sanger sequencing becomes a breaking task. At this point, next-generation sequencing (NGS) platforms present an alternative solution with their capacity to sequence the entire genome. Owing to the moderate costs and tractable data amounts, exome sequencing is a promising approach to detect novel mutations of human monogenic disorders. Hence, it is now possible to concurrently identify homozygous regions and possibly deleterious sequence variants, using data from a single sequencing experiment. To the best of our knowledge, only two programs have been developed to detect homozygous regions from NGS data, i.e. AgileVariantMapper (Carr et al., 2013) and HomozygosityMapper (Seelow and Schuelke, 2012). But, these programs do not take into account the distribution of the variants within the genomic coordinates. Here, we developed a novel sliding window-based methodology, which provides the advantage of scanning the genome in detail, but at the same time detecting the candidate homozygous regions easily. This type of analysis gives us the opportunity of converting the genotype information into a signal, where we can apply well-known signal processing techniques.

2 DESCRIPTION

HomSI was designed to define homozygous stretches in consanguineous families from NGS data. The overall analysis flow of HomSI is illustrated in Supplementary Figure S1.

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2.1 Processing NGS data

Variant call format (*.vcf) files are standard output files of variant identification programs that process NGS data. These files can be directly used as an input file to HomSI. Because the read depths for some regions might be low, we provide users different filtering options such as quality-based user-defined region or variant-based filtering. Details are available in the user manual.

To identify and visualize the homozygous stretches, HomSI processes each variant and generates several graphs. Here, we illustrate HomSI outputs using autosomal recessive Klippel Feil Syndrome dataset (Bayrakli et al., 2013). In Figure 1(a–b), the genotype information is plotted for different individuals, where the x-axis shows the samples and the y-axis indicates genomic coordinates. The variants of an index case are colored in blue for homozygous and in yellow for heterozygous cases. For other individuals, although a homozygous variant is indicated with blue, the contrasting homozygous is represented with white and the heterozygous with orange. For all individuals, gray indicates no call and yellow indicates non-informative SNPs, which are heterozygous for index case (see Supplementary Table S1 for details). HomSI starts with an unbiased approach and generates homozygosity maps for whole genome (Fig. 1a). Once a continuous blue line, which is shared among affected individuals, but not in unaffected ones, is recognized as in Figure 1a, the user can zoom in this region on the chromosome-based representation (Fig. 1b).

To be able to identify the ROSHs, we normalized the number of homozygous and heterozygous variants within each window separately (named as Hom and Het signals) and plotted these signals for each affected (Fig. 1c) and unaffected (Fig. 1d) individual. Using the differences between the Hom and Het signals, which were plotted for each individual in Figure 1e, a prediction signal was generated in Figure 1f. This final signal indicates the
ROSH for this consanguineous family. The detailed results (Supplementary Figs S2 and S7) and calculations of all signals are presented in Supplementary Material.

2.2 Comparative evaluation
We evaluated HomSI using both a simulated dataset generated by (Seelow and Schuelke, 2012) and a real dataset of three disease genes within the homozygous regions, which have been previously identified using a combination of exome and SNP microarray data (Aldahmesh et al., 2012; Shamseldin et al., 2012). On the simulated dataset, HomSI successfully detected the homozygous region on chromosome 15, as shown in Supplementary Figures S8 and S9. On the real dataset, we investigated whether the exome sequence data could identify a homozygous region, without prior SNP array analysis. For all three patients, HomSI detects the homozygous regions, as shown in Supplementary Figures S10 and S12.

3 CONCLUSIONS
We have developed a state-of-the-art method to identify homozygous stretches using NGS data and created a user-friendly tool. Because the usage of HomSI is so simple and intuitive, the clinicians or researchers can immediately start their data analysis without consulting to any dedicated information technology specialist. We provide a step-by-step tutorial and a detailed documentation on our Web site.

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**REFERENCES**