Insertion-deletions burden in copy number polymorphisms of the Tibetan population

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Introduction

Human genome consists of a diverse array of variations, which are present at various levels and at different frequencies. Some of these variations include single nucleotide polymorphisms (SNPs), insertions-deletions (inDels), segmental duplications, and copy number variations (CNVs). Though the size of the variation matters, however the location of occurrence determines the impact on the phenotype. [1,2] SNPs were thought to be the predominant form of genomic variation and account for much of the normal phenotypic variation. [3,4] However, widespread presence of inDels and CNVs in normal individuals seems to be a significant contributor to the phenotypic variation. [5] InDels are defined as the difference in the genome due to deletion or insertion of the nucleotides <1 kb and are distributed throughout the human genome with an average density of one in Del/7.2 kb of deoxyribonucleic acid (DNA). [6] Despite the fact that inDels are highly abundant in humans and cause a vast amount of variation in human genes, they have received less attention than SNPs and CNVs. Thus, inDels are of great interest since they can alter human traits and can cause human diseases. [7] InDels occur almost exclusively in repeat regions composed of homopolymers and multimeric simple sequence repeats, demonstrating the importance of sequence context for inDel mutations. [8] Several lines of evidence indicate that inDels in the coding region are likely to affect gene function in humans. [9] Further, a limited number of studies have looked into the effect of CNVs along with the burden load of inDel polymorphisms in the population. It will be necessary to look into inDels under CNVs, since they contribute significantly toward the genome structure and...
gene expression. Genome-wide CNV analyses followed by next-generation sequencing can clarify the context and burden of indels, thereby integrating a more detailed consideration of indels with our knowledge of nucleotide substitutions. In view of this, 31 Tibetan individuals were screened for the presence of indels in large copy number polymorphisms (CNPs) which map to functionally significant sites within human genes, and thus, are likely to influence human traits and diseases.

Materials and Methods

For this study, 31 randomly selected normal members residing in Tibet with different age groups were selected for CNP analysis. The raw, unprocessed data from Affymetrix Genome Wide SNP 6.0 (Affymetrix Inc., Santa Clara, CA, USA) array for all the 31 individuals of Tibet population was obtained from the ArrayExpress Archive at the European Bioinformatics Institute (EBI) that was submitted by Simonson et al.[10]

Sampling process and data analysis

Generally genome data are submitted to EBI databases. In a similar way the Tibetan genome data has been obtained from EBI submitted by Simonson et al., 2010.[10] Simonson et al., genotyped Forty-nine individual DNA samples Affymetrix 6.0 SNP Array technology (Affymetrix Inc., Santa Clara, CA, USA) (>900,000 SNPs). Further, they subjected this data for default parameters for the Birdseed algorithm (version 2), the Affymetrix Genotyping Console (GTC) 3.1, (Affymetrix Inc., Santa Clara, CA, USA) principal components analysis, pair-wise genetic distances and the proportion of shared genomic segments to determine relatedness was performed. Their analysis revealed that when pairs of individuals exhibited genetic distances <4.95 × 10^{-2} or had genome-wide identity-by-descent of >400 cM (minimum segment size 2.5 cM). Based on these criteria, a total of 31 unrelated individuals were included in the analyses.[10]

Copy number variation discovery

Genotyping

Genome-wide genotyping was performed using an Affymetrix Genome-wide Human SNP Array 6.0 chip having 1.8 million combined SNP and CNV markers with the median inter-marker distance of 500-600 bases. These chips provide maximum panel power and the highest physical coverage of the genome.[11] Genotyping quality was assessed using Affymetrix GTC Software. Copy number (CN) analysis method offers two types of segmenting methods, univariate and multivariate. These methods are based on the same algorithm, but use different criteria for determining cut-points denoting CNV boundaries.

Birdsuite (v2)

Birdsuite[12] is a suite originally developed to detect known common CNPs based on prior knowledge, as well as to discover rare CNVs, from Affymetrix SNP 6.0 array data (Affymetrix Inc., Santa Clara, CA, USA). To do this, it incorporates two main methods; the “Birdsuite” algorithms and the “Canary.”[13] The Birdsuite algorithm uses a Hidden Markov model (HMM) approach to find regions of variable CN in a sample. For the HMM, the hidden state is the true CN of the individual’s genome and the observed states are the normalized intensity measurements of each array probe. CNV calls from the Canary and Birdsuite algorithms were collated for each sample, and kept as long as they met the following criteria: (i) Birdsuite calls with a log10 of odds (LOD) score (odds ratio) ≥10 (corresponding to an approximate false discovery rate of ~ 5%), (ii) Birdsuite calls with CN states other than 2 were retained; (iii) Canary CNP calls with CN states different from the population mode were retained.

Canary

Copy number polymorphism analysis was performed using the Canary algorithm. Canary[3] was developed by the Broad Institute for making CN state calls in genomic regions with CNPs. Canary algorithm computes a single intensity summary statistic using a subset of manually selected probes within the CNP region. The intensity summaries are compared in aggregate across all samples to intensity summaries previously observed in training data to assign a CN state call.

Copy number variation finder

Copy number variation finder developed at the Welcome Trust Sanger Institute uses a dynamic,
multiple-threshold based approach to allow robust classification of CN changes in data of varying qualities. This algorithm makes two main assumptions (i) That the majority of data points are normally distributed around a log2 ratio of zero, and (ii) that data points falling outside of the centralized log2 ratio distribution are representative of a difference in CN between test and reference genome.

**Genotyping console**

After processing CEL files and the Birdseed to call genotypes, we used the GTC (GTC version 3.0.2) (Affymetrix Inc., Santa Clara, CA, USA) to detect CNVs from the Affymetrix 6.0 array for samples that passed initial quality controls (QCs). The default parameters of >100 Kb size and >15 probes in this algorithm were used.

**Data analysis**

Genome-wide CNV study was carried out using SVS Golden Helix version 7.2 (SVS Golden Helix, 2010, Bozeman, Montana, USA) and Affymetrix GTC software as prescribed in their manuals (Affymetrix technical notes, 2005; 2007; 2008). EIGENSTRAT method was used to avoid possibility of spurious associations resulting from population stratification. Bonferroni correction was employed for multiple testing and the corrected data were then used for CNV testing. Bonferroni methods for population data genotyped on the Affymetrix 6.0 platform was \( \alpha = 0.05 \) thresholds between \( 1 \times 10^{-7} \) and \( 7 \times 10^{-8} \).

Analyzing the collated data from both BirdSuite and Canary algorithms increased the stringency on those meeting the CNP calls with a LOD score \( \geq 10 \) corresponding to a false discovery rate of \( \sim 5\% \). All SNPs that were called using Birdseed version 2 algorithm had a QC call rate of \( >97\% \) across individuals. All the subjects and members with SNPs that passed SNP QC procedures were entered into the CNV analysis. Filters were set for ID call rates for the overall SNPs to identify IDs with poor quality DNA, if any. The CNV calls were generated using the Canary algorithm. In AGCS, contrast QC has to be \( >0.4 \) to be included in the CNV analyses. In this study, contrast QC observed was \( >2.5 \) across all samples showing a robust strength. To control for the possibility of spurious or artifact CNVs, we used the EIGENSTRAT approach of Price et al.\(^{[17]}\) This method derives the principal components of the correlations among gene variants and corrects for those correlations in the testing. CNVs were considered validated when there was a reciprocal overlap of 50% or greater with the reference set. Though, the Jaccard statistic is sensitive to the number of CNVs called by each algorithm (ideally each two algorithms would detect similar number of CNV calls), the relative values between the different comparisons of algorithms/platform/site are very informative. All the overlap analyses performed have handled losses and gains separately except when otherwise stated, and were conducted hierarchically. The calls from the algorithms that were called in both were not considered; instead, they were collated, so that the relative values between the different comparisons of algorithms/platform/site are still very informative.

**Insertions-deletions discovery**

2500 whole genome sequences from populations across the globe were obtained from the 1000 Human Genome project to identify the inDels in the identified CNV regions. Golden Helix GenomeBrowse™ 1.0.6 software (Bozeman, Montana, USA) was used to identify the inDels in the CNV regions found in the Tibetan genomes. The following datasets were used to determine the inDels in the genome.

- Reference Sequence, UCSC (Homo sapiens, GRCh_37)
- Reference Sequence Genes, UCSC (Homo sapiens, GRCh_37)
- 1 kG Phase 1 - Variant Frequencies 2012_04_26 v3, GHI (Homo sapiens, GRCh_37)
- NHLBI ESP5400 Exomes European American - 2011_12, GHI (Homo sapiens, GRCh_37)
- GWAS Catalog
- UCSC (Homo sapiens, GRCh_37)
- 1 kG Phase 1 EUR-Sites 2011_05, GHI (Homo sapiens, GRCh_37)
- 1 kG Phase 1 AFR-Sites 2011_05, GHI (Homo sapiens, GRCh_37)
- 1 kG Phase 1 All-Sites 2011_05 and GHI (Homo sapiens, GRCh_37).
Results

Whole genome CN analysis in 31 individuals from the Tibet population and their subsequent CN breakpoint analysis in 2500 sequenced genomes from the 1000 genomes project revealed the presence of InDel polymorphisms at various frequencies. Analyzing the data from the array and from those meeting the CNP calls with a LOD score ≥10 (corresponding to a false discovery rate of ~5%) criteria were selected for the study. We observed a total of 1051 CN events containing 80% duplications and 20% deletions [Figure 1]. These CNVs were seen scattered across chromosomes and were found impacting genes both entirely and partially. These polymorphisms were found enriched in genes when compared to pseudogenes. These CNVs encompassed a total of 2586 genes, which is ~10.5% of the total genes present in humans [Figure 1], which were identified frequently in multigene families compared to others. There are >10 such multigene families in the human genome where such variations were observed. The genes for alpha and beta chains of the mammalian hemoglobin molecule, actins, immunoglobulins, interferons, tubulins, hemoglobins, histones, ribosomal ribonucleic acid genes, alpha-amylase multigene family (AMY1 and AMY2), glucuronosyltransferase (UGT) gene family divided into two subfamilies, UGT1 and UGT2 are some of the multigene families where CNVs were seen. Of the CNVs identified, 85% of them contained entire coding structure of the genes while ~15% was found to be partially disrupted [Figure 1a].

We identified a total of 95935 InDel polymorphisms in the regulatory, exon and intron regions of the genes.
Table 1: Chromosome wise distribution of insertion and deletions in 31 Tibetan genomes

| Chromosome | Insertion | Deletion |
|------------|-----------|----------|
|            | Number    | Percentage | Number    | Percentage |
| 1          | 1207      | 3.0       | 2083      | 4.0        |
| 2          | 2662      | 7.0       | 4895      | 8.39       |
| 3          | 1328      | 3.5       | 1985      | 3.4        |
| 4          | 2104      | 5.5       | 3199      | 5.4        |
| 5          | 1080      | 2.8       | 1905      | 3.2        |
| 6          | 714       | 1.96      | 1127      | 1.9        |
| 7          | 1019      | 3.0       | 1066      | 1.82       |
| 8          | 2095      | 5.5       | 3743      | 6.4        |
| 9          | 1729      | 4.5       | 3413      | 5.8        |
| 10         | 856       | 2.2       | 1144      | 1.96       |
| 11         | 609       | 1.61      | 971       | 1.6        |
| 12         | 1079      | 2.86      | 1514      | 2.59       |
| 13         | 13        | 0         | 36        | 0.06       |
| 14         | 9300      | 24.7      | 13376     | 22.9       |
| 15         | 3461      | 9.1       | 4866      | 8.34       |
| 16         | 471       | 1.25      | 785       | 1.34       |
| 17         | 826       | 2.1       | 1090      | 1.86       |
| 18         | 231       | 1.61      | 413       | 0.7        |
| 19         | 1767      | 4.6       | 2611      | 4.47       |
| 20         | 155       | 0.41      | 248       | 0.42       |
| 21         | 344       | 0.69      | 598       | 1          |
| 22         | 2261      | 6         | 3539      | 6.03       |
| X          | 2323      | 6.1       | 3694      | 6.33       |
| Total      | 37634     | 87.12     | 58301     | 95.95      |

Subjects studied showed almost equal ratio of inDels in the intron regions and the remaining subjects showed increased number of deletions compared to insertions. InDels in exon regions of five subjects were between 10% and 13% and nine subjects were between 6% and 9% followed by <5% in the remaining subjects [Figure 2b]. Comparing inDel ratios of exon regions, only 16% of the subjects showed equal ratio of inDels and on the contrary, most of the remaining subjects showed increased number of insertions compared to deletions [Figure 2b]. The regulatory region showed the lowest concentration of inDels with six subjects showing inDels from 14% to 24% followed by nine subjects showing 5-9% and the remaining subjects showed variations <4% [Figure 2c].

The genomes of male and females were examined for the presence of inDel bias, if any. Females and males showed almost equal number of insertions 38.7% and 38.8% and deletions being 61.2% and 61.1%. Distribution of inDels was observed across chromosomes in the regions of CNVs. In the regulatory region, the number of deletions was found more in 8th chromosome. The ratio of inDel was observed to be almost equal in chromosomes 1, 9, 15 and 19, however, in chromosomes 3-6, 10, 11, 12 and 22, insertions were more than the deletions, but in chromosomes 7, 8, 14, 16, 17, 21 and X, deletions were more than insertions [Figure 3a]. In the exonic region of CNVs, the number of deletions is highest in the fifteenth chromosome and the ratio of inDel is almost equal in chromosomes 16, 19 and 3. In chromosomes 1, 2, 4, 6, 14, 17, 18 and X, insertions were more than deletions, whereas in chromosomes 5, 7-11, 15, 20-22 deletions were found to be less than insertions [Figure 3b]. In the intronic region of CNVs, the number of deletions is more than the X chromosome, and the ratio of inDel is almost equal in chromosomes 1, 2, 5, 6, 7, 8, 18, 20 and 21. In chromosomes 3, 4, 9, 10, 12, 16, 17 and 19 insertions are more than deletions, whereas in
chromosomes 11, 14, 15, 22 and X, deletions are more than insertions [Figure 3c].

Discussion

Insertions-deletions are an abundant form of structural variations in the genome alongside SNPs and CNVs in terms of numbers and size.\textsuperscript{[18]} It is important to estimate the presence of burden load of inDel polymorphism in the regions of CN to assess its contribution towards the phenotypic variability. These variations in the CN regions will also help in increasing the coverage of the markers for designing high resolution DNA chip. Our study of large CNVRs revealed a range varying from 100 kb to several Mbs scattered across all the chromosomes. Duplications CNVs observed were more frequent than deletion CNVs and were found impacting genes both entirely and partially. The identified inDels in these regions of the CNP of the Tibetan genome revealed a considerable load of single-base pair expansion,
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monomeric base pair expansions and multi-base pair polymorphisms.

About 10.5% of the genes were under the influence of CNP, and a significant number of genes among them were found to contain inDel variations. Deletions of single base pair to multi base pair were more compared to insertion variations. InDels within the regulatory and exon regions of a gene are known to cause syndromes in humans. The presence of inDels in 3 bp manifold maintains the open reading frames of the proteins intact and those, which are not in the manifold of 3 bp cause frame shifts in the encoded proteins, which are expected to disrupt gene function. Deletion variations were observed to be higher than insertion variations in the polymorphic regions across all the chromosomes and interestingly chromosome 14, 15, 2 and 8 were observed bearing a mass of these variations. The presence of either the copy number or the existence of the amount of genes in these chromosomes would hardly be the reason, since the presence of copy number or genes in other chromosomes is almost similar. Hence, this poses a great challenge to identify the reason behind such a complexity, and the genomic architecture in these regions should be further looked into, to reason out for these higher numbers of inDels.

Figure 3: Chromosome wise distribution of insertions-deletions across chromosomes in the (a) Regulatory region, (b) Exonic region and (c) Intronic region of Tibetan genome
The presence of inDels in the copy number regions of the subjects was observed to be in diverse forms and frequency. The array of subjects can be divided into quartiles for better understanding of their inDel structure. The first quartile contains subjects 1-8, the second quartile contains subjects from 9 to 15 and the third and fourth quartile contains 16-23 and 24-31 respectively. The subjects from first quartile show balanced frequency of inDels in intronic and exonic regions, but show very less concentration of inDels in regulatory regions. The subjects in second quartile show balanced frequency of inDels in intronic and regulatory regions, but show a higher frequency in exonic regions. The subjects in third quartile resemble the subjects from first quartile with a slightly higher concentration of inDels and the subjects in the fourth quartile show complex frequency and inconsistent pattern of inDel presence in all the three genomic regions. It is probably this diverse and inconsistent presence of inDels, which enables human to be phenotypically different and unique. This load of inDel polymorphism in the genome, presence of copy number regions, and the load of inDels in the copy number regions of the genome will be complex and intricate in understanding the contribution of these towards the phenotype since each of these polymorphism has to be analyzed individually and in tandem since it occurs on functionally important coding regions of the genome. InDels in the regulatory region alter the methylation patterns of the promoter leading to changes in gene expression. However, inDels that occur within the transcription factor binding sites or enhancers are found to diminish or abolish gene expression. This alteration is thought to be due to the phasing and spacing of DNA sequences within promoters caused by the inDels, which explains the differences in gene expression that has been observed in diverse humans. InDels in the coding region are likely to have a major impact on human biology and diseases due to the insertion or deletions of the amino acid(s), though, in other cases, inDels in the in-frame may not coincide perfectly with codon boundaries enabling additional amino acid changes that may also occur in the region that is altered. The coding regions that were under the burden load of inDels were found to be involved in several medically relevant biological pathways, including drug and lipid metabolism. A study by Zhang et al. showed genetic features contributed by CNVs in Tibetan population revealed distinct CNV allele frequencies and were enriched for genes in the disease classes of human reproduction and biological process categories of “response to DNA damage stimulus” and “DNA repair.”

Chromosomes showed various concentrations of inDels with very few in 13th, 20th and 21st. Chromosomes 14, 2 and 15 show almost equal number of CNVs compared to other chromosomes, but show higher concentration of inDels compared to any other chromosomes, which we believe is due to the presence of some repetitive elements in the CNV regions. Intron region contained the highest number of inDels compared with exon and regulatory region. High prevalence of inDels in the intron region can be owed to the ineffectuality of those variations in the genome. Exon region showed more inDels compared with regulatory region and it is these regions which contribute significantly toward the expression of the phenotype. In general, inDels in the regulatory region reduce the expression of the downstream genes, and similarly, inDels in the exon region truncate the coding structure of the gene depending on the location of the variation. Both male and female genomes were checked for the bias of inDel, but the frequency of inDel was similar in both females and males. Chromosomes 1, 9, 15 and 19 show almost equal number of CNVs and also the ratio of inDel in these chromosomes are almost equal.

A collection of these inDel genetic markers thus helps in, genetic mapping, identification of disease genes and in human identity testing. The identification of these variations adds up to the existing catalogue of variations and this is the first study which looks into the effect of inDels in CNVs on the genome. The identification of the inDel variations in the regions of CNV will further help in increasing the coverage of the genome and in detecting novel CNVs. Further studies are required to understand the genome topology and to analyze the combined effect of inDels and CNVs on the genome structure, gene expression and phenotype.

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