Copy Number Variation of UGT 2B Genes in Indian Families Using Whole Genome Scans

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1. Introduction

CNVs are the presence of segments of DNA longer than 1 kb with >90% sequence identity which differ in the number of copies between the genomes of different individuals [1]. They affect more nucleotides per genome than SNP variation [2] and contribute significantly to variation among normal individuals, both in levels of gene expression and in phenotypes of medical relevance [3, 4]. There are many genes and gene families that show copy number differences in population. Around 19 such loci have been identified in genome-wide surveys; they are found to harbor both deletion and duplication alleles [5]. One of these is the uridine diphospho-glucuronosyltransferase 2B (UGT2B), which is a family of genes, involved in metabolizing steroid hormones and several other xenobiotics. UGT2B family of genes has high sequence identity and similar enzymatic functions with UGT2B4, UGT2B7, UGT2B10, UGT2B11, UGT2B15, UGT2B17, and UGT2B28 [6]. In Chinese population, heterozygous deletion polymorphism of UGT2B17 was higher (86%) than homozygous deletion (73%) [2, 7]. These polymorphisms have also been reported in samples covering American, European, and African populations [8–12]. However, such polymorphisms have not been reported from South Asia, particularly from India. In addition, there are controversial reports regarding the association of UGT2B17 polymorphism with lung cancer, osteoporosis, and prostate cancer [8, 12, 13]. It was therefore of interest to investigate the polymorphic condition of UGT 2B genes in the Indian families as family studies are more robust to population stratification. Here, we report the presence of UGT 2B gene deletion and duplication polymorphisms and also the network analysis of the UGT 2B genes, which predicts the involvement of other possible genes in the uridine diphospho-glucuronosyltransferase activity.
2. Materials and Methods

For this study, 31 normal members from randomly selected eight families residing in Mysore (3), Mandya (1), Davangere (3), and Bangalore (1) certified by All India Institute of Speech and Hearing (AIIISH), Mysore, and National Institute of Mental Health and Neuro Sciences (NIMHANS), Bangalore, were selected. These families were subjected to power analysis using standard parameters [14] by specifying an effect size range of 1550–1750. Performing the 1-sample, 1-sided test using the Type I error rate at 5% resulted in the power being equal to 1. Based on this, we selected 1746 total samples for the study. 5 mL EDTA blood was collected from each member of the family and genomic DNA was extracted using Promega Wizard® Genomic DNA purification kit. The isolated DNA was quantified by biophotometer and gel electrophoresis. The study was conducted for a period of two years from 2009 to 2011. The institutional ethics review board approved this research and informed consent was obtained from all sample donors.

2.1. Genotyping. Genome-wide genotyping was performed using an Affymetrix Genome-Wide Human SNP Array 6.0 chip, which has 1.8 million combined SNP and CNV markers with the median intermarker distance of 680 bases. This chip provides maximum panel power and the highest physical coverage of the genome [15]. Genotyping quality was assessed using Affymetrix Genotyping Console Software (AGCC). Briefly, all SNPs that were called using Birdsseed v2 algorithm [16] had a Quality Control (QC) call rate of >97% across members in families. All the members who passed SNP QC procedures were entered into the CNV analysis. The CNV calls were generated using the Canary algorithm. Contrast QC across all samples was >2.5 as required to be >0.4.

2.2. Data Analysis. Genome-wide CNV study was carried out using SVS Golden Helix Ver. 7.2 [17] and Affymetrix Genotyping Console software as prescribed in their manuals [18–20]. 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.

2.3. Weighted Protein Interaction Network Analysis. We used weighted protein network analysis in a first attempt to identify steroid metabolic process associated modules and their key constituents. Weighted protein network analysis starts from the level of thousands of genes, identifies modules of physically interacting proteins, colocalized proteins, coexpressed proteins, and proteins falling under the common pathway, and relates these modules to clinical variables and gene ontology information. We made use of tools such as GeneMANIA, BIOGRID, and CYTOSCAPE developed for network pathway studies to assess the functional consequences of the network topology. GeneMANIA Protein network association database, a FDR corrected hypergeometric test, was used for enrichment in the UGT 2B network [21].

3. Results

Different age group members ranging within 13–73 years were subjected to whole genome scan from eight families. Nine of them were aged 13–16 years and were studying at high school level; 5 of them were aged 16–25 years and had completed graduation; 5 of them were aged 45–55 years and were employees; 3 of them were aged 45–55 years and were businessmen; 7 of them were aged 35–45 years and were housewives; and 2 of them were aged 70–73 years and are grandparents. None of these members had history of any diseases (Table 1 and Figure 1).

CNV analysis of eight families revealed in the first family 1A-III-3 subject showing a 114 kb duplication polymorphism and 1A-III-4 subject with a 114 kb deletion polymorphism. Both of these subjects are identical twins. 1A-I-1 subject of the same family showed only a deletion polymorphism of 109 kb in UGT2B17. Segregation of 111 kb heterozygous deletion genotype of UGT2B17 was seen in four subjects of the second family (Table 1) (Figure 1(A)). Subject DFIA in the third family had a duplication of 168 kb in UGT2B17 and UGT2B15 and deletion in UGT2B28 of 168 kb size which belongs to the same gene family of UGTs. However, subjects DF1F and DF1U of the same family show a 108 kb and a 105 kb deletion of UGT2B28 only.

Subject 2DF4F in the fourth family had a deletion of 216 kb in UGT2B17 and UGT2B15. However, subjects 2DF4A and 2DF4M of the same family showed 104 kb and 100 kb deletion, respectively, of UGT2B28. In the fifth family, subjects 2DF5A and 2DF5U showed a duplication of 141 kb and deletion of 146 kb in the UGT2B17, respectively. Subject 2DF2F from the sixth family had a duplication of 200 kb in UGT2B17 and UGT2B15, while subject 2DF3A from the seventh family showed a deletion of 114 kb involving UGT2B17. In UGT2B15, duplications in two subjects and deletion in one subject were observed, whereas in UGT2B28 duplication in one individual and deletions in five individuals were seen (Table 1). These CNVs were also validated using SVS Golden Helix Version 7.2 and the duplication breakpoints were found to be novel when checked against the online CNV database.

By taking cognizance of all the 8 families, both duplication and deletion polymorphisms of UGT 2B genes were observed in these families (Table 1) and the copy number state varied from 1 to 3 copies (Figure 2). The CNVs of UGT 2B genes were found to be in heterozygous deletion and duplication (Figure 3). The prevalence of UGT2B17 CNVs was found to be 76.4% and that of UGT2B15 was 17.6%, whereas UGT2B28 showed a prevalence of 35.2% in this study. No zero allelic state was observed in these families and a minimum of one allele to a maximum of three alleles of UGT 2B were observed in the genotypes of the families. The deletions and duplications observed here encompassed the complete gene structure and its flanking regions (Table 1).

A heat map showing the difference in heat emission for normal, deletion polymorphisms, and duplication polymorphisms of the 4q13.2 region with inferred functional copy number for all the members under study can be seen in Figure 2.
Table 1: Functional annotation of proteins, their significant $p$ value, and the number of genes participating in the UGT2B17, UGT2B15, and UGT2B28 mediated steroid metabolism and glucuronidation pathways.

| Symbol          | Entrez Gene Name | Location               | Type(s)                      | Biomarker Application(s)                |
|-----------------|------------------|------------------------|------------------------------|------------------------------------------|
| ABCG1           | ATP-binding cassette, subfamily G (white), member 1 | Plasma membrane           | Transporter                   | Unspecified application                  |
| Androgen        | —                | Other                  | Chemical, other              |                                          |
| Androsterone    | —                | Other                  | Chemical, endogenous mammalian |                                          |
| C/ebp           | —                | Cytoplasm              | Group                        |                                          |
| Diclofenac      | —                | Other                  | Chemical drug                |                                          |
| Dihydrotestosterone | —                | Other                  | Chemical, endogenous mammalian | Diagnosis, efficacy               |
| ELF5            | E74-like factor 5 (ets domain transcription factor) | Nucleus                  | Transcription regulator       |                                          |
| Fluconazole     | —                | Nucleus                 | Transcription regulator       | Disease progression                    |
| FOXA1           | Forkhead box A1  | Nucleus                 | Transcription regulator       |                                          |
| HNF1B           | —                | Other                  | Chemical drug                |                                          |
| Ibuprofen       | —                | Other                  | Chemical drug                |                                          |
| KLK3            | Kallikrein-related peptidase 3 | Extracellular space | Peptidase                   | Diagnosis, disease progression, efficacy, safety, and unspecified application |
| KLK4            | Kallikrein-related peptidase 4 | Extracellular space | Peptidase                   | Diagnosis, efficacy                     |
| Methylseleninic acid | —            | Other                  | Chemical reagent             |                                          |
| miR-619-3p (miRNAs w/ seed ACCUGGA) | —                | Cytoplasm              | Mature microRNA               |                                          |
| NKX3-1          | NK3 homeobox 1   | Nucleus                 | Transcription regulator       |                                          |
| ONECUT1         | One cut homeobox 1 | Nucleus                 | Transcription regulator       |                                          |
| Phenobarbital   | —                | Other                  | Chemical, other              |                                          |
| Progestin       | —                | Nucleus                 | Other                        |                                          |
| RAD23A          | RAD23 homolog A (S. cerevisiae) | Nucleus | Other                        |                                          |
| SLC16A6         | Solute carrier family 16, member 6 (monocarboxylic acid transporter 7) | Plasma membrane | Transporter                   |                                          |
| SLC25A13        | Solute carrier family 25 (aspartate/glutamate carrier), member 13 | Cytoplasm              | Transporter                   |                                          |
| SLC9A3          | Solute carrier family 9, subfamily A (NHE3, cation proton antiporter 3), member 3 | Plasma membrane | Ion channel                   | Unspecified application                  |
| Testosterone    | —                | Other                  | Chemical, endogenous mammalian | Diagnosis, efficacy, prognosis, safety |
| TMPRSS2         | Transmembrane protease, serine 2 | Plasma membrane | Peptidase                   | Diagnosis, efficacy                     |
| trans-3'-hydroxycotinine-glucuronide | —                | Other                  | Chemical reagent             |                                          |
| UGT1A1          | UDP glucuronosyltransferase 1 family, polypeptide A1 | Cytoplasm              | Enzyme                       | Diagnosis, efficacy                     |
| UGT2B15         | UDP glucuronosyltransferase 2 family, polypeptide B15 | Cytoplasm              | Enzyme                       |                                          |
| UGT2B17         | UDP glucuronosyltransferase 2 family, polypeptide B17 | Cytoplasm              | Enzyme                       |                                          |
| UGT2B28         | UDP glucuronosyltransferase 2 family, polypeptide B28 | Cytoplasm              | Other                        |                                          |
3.1. Molecular Protein Interaction Network of UGT 2B Genes.

The network analysis of UGT 2B genes establishes interconnecting pathways of genes involved in steroid hormones processing and xenobiotics metabolizing such as UGT2B7, UGT2B15, UGT2B4, and CYP3A4 (Figure 4). These genes function independently in their own pathways involving coexpression and colocalization of UGT2B4, UGT2B28, and UGT2B15 with UGT2B17 proteins in the protein association network indicating the role of other UGT 2B enzymes in this pathway. The loss or presence of UGT2B17 did not have a significant functional impact in the disease pathways. Since the gene modules in the network correspond to biological pathways, focusing the analysis on modules and their highly connected intramodular hub genes identifies the significant role played by each gene functionally. In this way, the impact of UGT2B17 in its loss or gain status on the biological pathway can be downplayed in the earlier reported diseases. This network also provides predictive genes involved in the steroid metabolizing pathway and xenobiotic metabolizing pathway (Table I).

4. Discussion

Perusal of the literature revealed that UGT2B17 in the 4q13.2 region is highly polymorphic and the frequency of polymorphism in this gene was found to be more in African populations, intermediate in Europe and parts of West Asia, and low in East Asia [8–11]. Polymorphisms have been previously described for 1A UGTs, as well as several members of family 2B UGTs, including UGT2B4, UGT2B7, UGT2B15, and UGT2B17 [7, 22–24]. CNVs in the UGT 2B genes are common in the general population as suggested by Xue et al. [10] and Che et al. [12]. However, the role of polymorphism in UGT2B17 remains controversial since the deletion polymorphism of UGT2B17 was found to be associated with lung cancer, osteoporosis, prostate cancer, and endometrial cancer [9, 11, 25, 26] while there are also studies to relate the duplication polymorphism with lower BMD, thinner CT, higher BR, and osteoporosis [25]. On the contrary, Gallagher et al. [8] reported nonassociation of UGT2B17 polymorphism with the risk of lung cancer. Similarly, Olsson et al. [13] also opined the nonassociation of UGT2B17 polymorphism with prostate cancer risk and, recently, Che et al. [12] disregarded homozygous deletion genotype of UGT2B17 with osteoporosis risk. Either the deletion or duplication polymorphisms in UGT2B15 are occurring along with the polymorphism of UGT2B17, which could be due to its close proximity. However, CNVs in UGT2B28 for the first time have also been identified from this study. This gene also encodes for the uridine diphospho-glucuronosyltransferase protein. The encoded enzyme catalyzes the transfer of glucuronic acid from uridine diphosphoglucuronic acid to a diverse array of substrates including steroid hormones and lipid-soluble drugs (Table I). Two transcript variants encoding different isoforms have been found for this gene [27].

Figure 1: Illustration of pedigrees and heatmap of UGT2B gene regions in subjects under study. (A and B) The pedigrees of families showing the allelic state in family members in 4q13.2/UGT2B17 region. Subject 1 = 1A-I-2; Subject 10 = 1A-III-4; Subject 11 = 1A-III-3; Subject 12 = 1A-II-8; Subject 13 = 1A-I-1; Subject 14 = 1A-II-3 in Pedigree A and Subject 6 = 1B-V-2; Subject 7 = 1B-III-5; Subject 8 = 1B-IV-2; and Subject 9 = 1B-V-1 in Pedigree B have been genotyped.
CNVs are widely distributed in the genome; these CNVs might be the consequence of recurrent events via homologous recombination. Examining the sequences of UGT 2B family genes revealed ~95% sequence identity and also found a major concentration of repeat sequences which are thought to mediate homologous recombination. Therefore, we strongly believe the role of transposition activity to be the cause of deletion and duplication polymorphism. A typical example can be seen in the twins from the first family (Figure 1). Weighted protein network analysis started with UGT2B17, identified modules of physically interacting proteins, colocalized proteins, coexpressed proteins, and proteins falling under the common pathway, and related these modules to clinical variables and gene ontology information. In addition, identification and establishment of network pathways could help us to further understand the molecular mechanism in more refined manner than the existing one. The present study identified de novo CNV events in the UGT 2B region. Since CNVs in UGT 2B region were identified in normal subjects and since this gene is specifically expressed...
Figure 3: Schematic representation of log$R$ Ratios of the UGT 2B gene regions showing normal panel and deletion and duplication polymorphisms. A common representative image of the log$R$ Ratios which indicates the quantitative assessments of genotyping used to determine CNV of 4q13.2/UGT 2B gene region. log$R$ Ratios in the first panel are of a member without polymorphism, deletion polymorphism, and duplication polymorphism in the second and third panels, respectively. Arrows indicate the gain and loss status. Enriched log$R$ Ratios below the median line indicate deletion polymorphism and enriched log$R$ Ratios above the median indicate duplication polymorphism.

Figure 4: Network of UGT 2B genes involved in uridine diphospho-glucuronosyltransferase activity.
to metabolize the steroids, identifying the changes at protein level as well as small sample size was a possible limitation of this study.

Competing Interests
The authors declare no conflict of interests.

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References

[1] J. L. Freeman, G. H. Perry, L. Feuk et al., “Copy number variation: new insights in genome diversity,” Genome Research, vol. 16, no. 8, pp. 949–961, 2006.

[2] R. Redon, S. Ishikawa, K. R. Fitch et al., “Global variation in copy number in the human genome,” Nature, vol. 444, no. 7118, pp. 444–454, 2006.

[3] T. J. Aitman, R. Dong, T. J. Vyse et al., “Copy number polymorphism in Fcg3 predisposes to glomerulonephritis in rats and humans,” Nature, vol. 439, no. 7078, pp. 851–855, 2006.

[4] E. Gonzalez, H. Kulkarni, H. Bolivar et al., “The influence of CCL3L1 gene-containing segmental duplications on HIV-1/AIDS susceptibility,” Science, vol. 307, no. 5714, pp. 1434–1440, 2005.

[5] J. M. Young, R. M. Endicott, S. S. Parghi, M. Walker, J. M. Kidd, and B. J. Trask, “Extensive copy-number variation of the human olfactory receptor gene family,” The American Journal of Human Genetics, vol. 83, no. 2, pp. 228–242, 2008.

[6] T. R. Tephly and B. Burchell, “UDP-glucuronosyltransferases: a family of detoxifying enzymes,” Trends in Pharmacological Sciences, vol. 11, no. 7, pp. 276–279, 1990.

[7] C. Jin, J. O. Miners, K. J. Lileywhite, and P. I. Mackenzie, “Complementary deoxyribonucleic acid cloning and expression of a human liver uridine diphasphate-glucuronosyltransferase glucuronidating carboxylic acid-containing drugs,” Journal of Pharmacology and Experimental Therapeutics, vol. 264, no. 1, pp. 475–479, 1993.

[8] C. J. Gallagher, F. F. Kadiubar, J. E. Muscat, C. B. Ambrosone, N. P. Lang, and P. Lazarus, “The UGT2B17 gene deletion polymorphism and risk of prostate cancer: A case-control study in Caucasians,” Cancer Detection and Prevention, vol. 31, no. 4, pp. 310–315, 2007.

[9] A.-H. Karypidis, M. Olsson, S.-O. Andersson, A. Rane, and L. Ekström, “Deletion polymorphism of the UGT2B17 gene is associated with increased risk for prostate cancer and correlated to gene expression in the prostate,” The Pharmacogenomics Journal, vol. 8, no. 2, pp. 147–151, 2008.

[10] Y. Xue, D. Sun, A. Daly et al., “Adaptive evolution of UGT2B17 copy-number variation,” American Journal of Human Genetics, vol. 83, no. 3, pp. 337–346, 2008.