The Genetics of IgA Nephropathy: An Overview from China

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Abstract

Background: IgA nephropathy (IgAN) is the most common type of primary glomerulonephritis worldwide. Highly variable data for disease prevalence and reports of familial clustering suggest the involvement of genetic factors in IgAN. As China is an area with a high prevalence of IgAN, Chinese scholars have made a considerable effort to reveal the underlying genetic architecture of IgAN. Summary: In this review, we summarize recent achievements in the genetic studies of IgAN, focusing mainly on studies undertaken in China. Early association studies followed a population-based design and focused on a single variant or single gene. Subsequently, family-based designs and genetic interactions applied by Chinese scholars revealed an association of variants in MEGSIN and glycosyltransferase genes with IgAN. Recently, genome-wide association studies (GWAS) have been used to identify multiple susceptibility loci for IgAN, and they have, for the most part, been validated in Chinese populations. Key Messages: More efforts should be made to explore the underlying genetic mechanisms of GWAS-identified variants. In future studies in IgAN, the application of a systems genetics approach would be helpful and productive. Facts from East and West: The reported prevalence of IgAN is higher in Asia than in Europe and North America. However, differences in use of biopsy for the diagnosis of IgAN should be taken into account in analyzing data from both East and West. In Europe, IgAN affects men more frequently than women; this is not the case in Asia. Familial IgAN has been more frequently reported in Europe than in Asia. Within Europe, familial IgAN is more evident in southern than in northern populations. Changes in the pattern of serum IgA1 O-glycosylation is a common finding in IgAN patients in the East and West. SNPs within the gene coding for the enzyme C1GALT1 have been reported in Chinese and European patients. However, there is no evidence for a role of gene polymorphism of the C1GALT1 chaperone cosmc in Europeans. Genetic variants in the HLA gene family have been observed in populations from the East and West. Associations between IgAN and variants of the TAP1/PSMB and DEFA genes were observed in Asian but not in Western patients. Association with the angiotensin-converting enzyme gene was seen only in Asian patients.

Key Words
IgA nephropathy · Genetics · Linkage analyses · Association analyses

For an overview of the genetics of IgA nephropathy in Western countries, see Feehally and Barratt, Kidney Diseases 2015, DOI: 10.1159/000381738, www.karger.com/doi/10.1159/000381738.
IgA nephropathy (IgAN) is the most common type of primary glomerulonephritis worldwide [1]. In China, it accounts for 37–58% of biopsy-proven primary glomerulonephritis [2–4]. Patients with IgAN bear a broad spectrum of clinical and pathologic changes, and 30% of them progress to end-stage renal disease within 10 years after the diagnosis [5, 6]. The high prevalence and chronic progression of IgAN result in a considerable disease burden.

The exact pathogenesis of IgAN is unknown. However, increasing evidence suggests the involvement of genetic factors in the development and progression of IgAN [7]. Genetic studies have led to considerable improvement in knowledge of the pathogenesis of IgAN [8].

As China is an area with a high prevalence of IgAN, Chinese scholars have made considerable efforts to reveal the underlying genetic architecture of IgAN. In this review, after a brief description of the genetic characteristics of IgAN, we summarize recent achievements in the genetic studies of IgAN, focusing mainly on studies undertaken in China.

Genetic Characteristics of IgAN

The prevalence of IgAN shows considerable differences among geographic and racial populations. The Asian population has a prevalence of 30–60% [2–4, 9, 10], the European population one of 20–30% [11, 12], and the North American population one of 10% [13]. The influence of polices regarding renal biopsy on the reported prevalence of IgAN should be considered, but genetic factors also contribute to this difference: indigenous North Americans have a IgAN prevalence of 38% [14], whereas that of African-Americans is 2% [15].

Moreover, after the first report of two affected siblings with IgAN by Tolkoff-Rubin et al. in 1978 [16], studies about the familial clustering of IgAN from all over the world were published [17–19]. In a study from Italy, patients with IgAN were classified into ‘sporadic’ (only 1 family member was affected), ‘suspected familial’ (1 member had biopsy-proven IgAN and others showed persistent microscopic hematuria and refused renal biopsy) and ‘familial’ (at least 2 individuals had biopsy-proven IgAN) [20]. The investigators observed that >50% of IgAN patients had familial IgAN or suspected familial IgAN [20]. However, it has been reported that the proportion of individuals in China with familial IgAN and suspected familial IgAN is 1.3 and 7.4%, respectively, which seems to be much lower than that in European subjects. Reports of these pedigrees suggest the genetic disease features of IgAN. However, the striking differences in the prevalence of sporadic IgAN and familial IgAN around the world also imply a role for genetic factors in IgAN.

Genetic Studies of IgAN

Thanks to the Human Genome Project, hundreds and thousands of genetic markers that span the entire human genome have been discovered. Based on these markers, genetic studies have been conducted widely in recent years. Linkage analyses and genetic association studies for IgAN have been carried out. Linkage studies are based on the theory of genetic linkage and can thus be conducted in genetic pedigrees. The logarithm of the odds score is used in linkage analyses as a statistical test to evaluate the probability that a gene important for a disease is linked to a genetic marker. However, recruiting subjects with the familial forms of IgAN is quite difficult. Therefore, along with the rapid development of genotyping technology, genetic association studies designed for patients with familial and sporadic forms of IgAN have been conducted widely.

Linkage Analyses

Linkage analyses in IgAN have been reported from three research teams [21–23]. As early as 2000, Gharavi et al. [21] demonstrated linkage of IgAN to 6q22–23 under a dominant model of transmission with incomplete penetrance, and this locus was termed ‘IGAN1’. Later, scholars from Europe reported two additional loci linked to IgAN: 4q26–31 (IGAN2) and 17q12–22 (IGAN3) [22]. Another locus identified from a large IgAN pedigree in Canada was located at 2q36 [23], which is a region that also contains COL4A3 and COL4A4, the causal genes for thin basement membrane disease (TBMD). Patients with IgAN and TBMD share many clinical and pathologic features [24], so whether 2q36 is linked to IgAN or just to TBMD is unknown [25].

After identifying susceptible loci linked to IgAN, scholars have made considerable efforts to explore the causative genes in those regions, but unfortunately the causative genes have yet to be identified. Many reasons have been postulated regarding the failure of identification of the causative genes of IgAN by linkage analyses. The diagnosis of IgAN is entirely dependent on renal biopsy. Hence, the accurate identification of family mem-
bers as being ‘affected’ or ‘unaffected’ is difficult because of the lack of a reliable noninvasive test for screening. Moreover, the broadly variable clinical presentations of IgAN suggest that this disease may encompass multiple subsets, and that these subsets might result in genetic heterogeneity in patients, even in the familial form. Karnib et al. [26] found no evidence of linkage to the reported IgAN loci on chromosomes 6q22–23, 2q36 and 4q22–31 in a large Lebanese family with IgAN (five generations and 16 affected individuals). Those findings suggested genetic heterogeneity and called for distinction based on genetic/biomarker data before linkage analyses. In addition, there is uncertainty regarding the genetic model for IgAN. A dominant model with incomplete penetrance has been proposed [21], but the complexity of IgAN suggests that it is a multi-factorial and polygenic disease involving environmental and genetic factors.

**Association Studies**

Compared with linkage analyses, association studies have been conducted more widely in China. In the early days, association studies had a population-based design and focused on a single variant or single gene. Then, when association studies were closely associated with a low rate of replication, family-based designs were introduced. With the rapid progress of bioinformatic technology, subsequent association studies have paid more attention to multiple variants in a single gene (haplotype) and genetic interactions in multiple genes. In recent years, rapid improvements in single nucleotide polymorphism (SNP) genotyping technology have led to genome-wide association studies (GWAS). Two GWAS in IgAN based on Chinese populations have been conducted [27, 28]. Multiple loci identified by GWAS are now awaiting functional studies to reveal the underlying genetic pathogenesis.

**Early Genetic Association Studies**

Human leukocyte antigen (HLA) gene family [18, 29, 30], renin-angiotensin system-related [31–37], TRAC [38, 39] and cytokine-related genes (interleukin-1, transforming growth factor-β1) [40, 41] have undergone association studies to test the development and/or progression of IgAN.

Reports from Hong Kong, Taiwan and Sichuan have revealed genetic variants in the HLA gene family to be associated with the genetic susceptibility of IgAN, or a special subgroup of IgAN [18, 29, 30]. Renin-angiotensin system-related genes have been postulated as candidate genes for association studies in IgAN but have yielded contradictory results [31–37]. Among variants in the renin-angiotensin system, insertion/deletion (I/D) polymorphisms in angiotensin-converting enzyme (ACE) genes have been observed. Systematic reviews and meta-analyses have demonstrated an association between I/D polymorphisms in ACE genes and IgAN in Asian populations, but genetic association was not so important in Caucasian populations, which might explain the controversial results in previous genetic association studies [31, 33, 34]. Actually, most early genetic association studies based on a candidate-gene approach have not been replicated. Besides genetic heterogeneity between ethnic groups, methodological problems (e.g., small sample size, poorly matched controls, inadequate coverage of variants in candidate genes) and a lack of statistical power have made genetic association studies for IgAN using a candidate-gene approach rather unrevealing.

**Family-Based Association Studies**

When hundreds of candidate gene-based association studies for IgAN were conducted, many positive genetic associations were noted, but, simultaneously, the number of replications of these associations was low. This phenomenon arose from an intrinsic disadvantage of candidate-based association studies, population stratification, in which mismatched populations between cases and controls elicit spurious associations that could not be validated.

Family-based association studies can be applied to overcome the weakness of population stratification. Unlike a traditional population-based design, the simplest study design for a family-based association, called a ‘transmission disequilibrium test’ (TDT), uses genotype data from trios. Then, through comparison of the observed number of alleles transmitted with those expected in Mendelian transmissions, the excess of alleles transmitted to affected individuals can be identified, which should be the disease-susceptible alleles. Therefore, a TDT design is not sensitive to population structures, and significant findings suggest susceptible alleles.

Using a TDT design, Li et al. [42] revealed that MEGSIN C2093T and C2180T confer susceptibility to IgAN. In subsequent studies, involvement of genetic variations of MEGSIN in IgAN progression was demonstrated in Chinese populations [43, 44]. A recent meta-analysis on seven reports focusing on genetic variations in MEGSIN showed a genetic association between MEGSIN variants and risk of IgAN in an Asian population [45].
Recent Association Studies Focusing on Genetic Interactions

In the last decade, a key breakthrough in the study of the pathogenesis of IgAN has been the identification of an abnormality in O-glycosylation in IgA1 molecules [46]. Several studies on pathogenic mechanisms have reported that defects in IgA1 glycosylation lead to the formation of immune complexes and initiate the development of IgAN. More recently, research teams from the USA and China have independently reported that defects in IgA1 glycosylation are an inherited phenotype in patients with familial and sporadic IgAN and their relatives [47, 48] and constitute a heritable risk factor for IgAN.

The O-glycan of IgA1 molecules is synthesized by a series of glycosyltransferases. Thus, we conducted genetic association studies with four key glycosyltransferase genes for IgA1 O-glycosylation (CIGALT1, ST6GALNAC2, CIGALT1C1, MUC20) as candidates, and tested the genetic association of variants in these genes in a large population of Chinese subjects with IgAN [49–53]. Our study identified one protective (YATIG) and two risky (YAGDA, YATDG) regulatory haplotypes in the CIGALT1 gene and proposed a genetic association between the CIGALT1 gene and IgAN [49], which was validated independently in three subsequent studies in European and Chinese populations [54–56]. Besides CIGALT1, we also demonstrated that the ADG haplotype, in the promoter region of ST6GALNAC2 is a functional regulatory polymorphism that results in a predisposition to IgAN [50]. Furthermore, using two independent multiple-locus analysis algorithms, we demonstrated the genetic interaction between CIGALT1 and ST6GALNAC2 and IgA1 glycosylation, as well as the predisposition and severity of IgAN [52]. This observation suggested that variants in CIGALT1 and ST6GALNAC2, through their influence on IgA1 O-glycosylation, contribute to IgAN in a polygenic manner [52].

Genome-Wide Association Studies

Four GWAS have been conducted in IgAN populations [27, 28, 57, 58]. The first GWAS in IgAN detected about 300,000 SNPs in a European population [57]. In that study, when tested against a significance threshold in GWAS, HLA regions (across loci for HLA-B, DRB1, DQA, and DQB) showed significant associations with IgAN, which was in agreement with the results in earlier genetic association studies, in which a HLA gene family was identified.

The next GWAS used >2,000 Northern Chinese subjects as a discovery cohort and was validated in a Southern Chinese and a Caucasian population [27]. In that study, five associated loci were identified. Besides three independent loci in HLA regions, 1q32 (encompasses CFH-CFHRs) and 22q12 (encompasses OSM and LIF) attracted much attention. Based on novel identification by GWAS, our research team further explored the underlying mechanism of genes in 1q32 with IgAN susceptibility [59]. We found that genetic variants in CFH, CFHR3 and CFHR1 affected complement activation through their influence on CFH levels and thereby predisposed individuals to develop IgAN. A subsequent GWAS by a Chinese research team focused entirely on a Chinese population [28]. In that study, a Southern Chinese population was used as a discovery cohort, and identified loci were validated further in Northern and Southern Chinese populations. In addition to previous identification, two additional loci were reported: 17p13 (containing TNFSF13) and 8p23 (containing DEFA). Two subsequent functional studies focused on the DEFA gene. Xu et al. [60] revealed that polymorphisms within DEFA genes are involved in gene transcriptional regulation, and Qi et al. [61] reported the association of polymorphisms in the DEFA gene with the clinical phenotype of gross hematuria, which implied the role of mucosal immunity in pathogenesis of IgAN. The most recent GWAS in IgAN were carried out in 20,612 individuals of European and East Asian ancestry [58]. The enlarged population provided increased power for the identification of new loci. Actually, six new genome-wide significant associations, four in ITGAM-ITGAX, VAV3 and CARD9 and two new independent signals at HLA-DQB1 and DEFA were reported. When evaluating the association with pathogen diversity and upon application of a pathway-based GWAS approach to the identified IgAN-susceptible variants, a possible role for interactions between hosts and intestinal pathogens were suggested to shape the genetic landscape of IgAN. However, the exact functions of these newly identified loci have yet to be revealed.

Future Perspectives on Genetic Studies in IgAN

Chinese scholars have participated actively in genetic studies of IgAN, especially in recent GWAS. Cumulatively, we have identified (by GWAS) 15 loci for IgAN susceptibility, but the underlying genetic mechanism is unknown. There is an urgent need to understand how these genetic variants contribute to disease susceptibility. Therefore, in the search for novel IgAN-susceptible loci, more efforts should be made to explore the underlying function of the associated genetic variants. Typically,
most of the loci identified by GWAS for common diseases show mild-to-modest effects. Hence, definition of the mechanistic effects of genetic variants on clinical traits using classical molecular biology approaches will be difficult. The application of a systems genetics approach, which uses a range of experimental and statistical methods to integrate multiple levels of information (transcription as well as protein and metabolite levels) [62], would be a helpful and productive strategy in future genetic studies in IgAN.

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Disclosure Statement

The authors have no competing financial interests to declare.
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