IgA Nephropathy: Actually an Inherited Disease?

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Abstract
IgA Nephropathy (IgAN) is the most common primary glomerulonephritis worldwide. Over the last years strong evidence suggest the existence of genetic factors in the development and/or progression of IgAN. IgAN does not exhibit classic single-gene Mendelian but rather a complex genetic inheritance pattern. Until now, two basic approaches have been used in genetic studies of IgAN: Genome-Wide linkage Analysis Studies (GWAS) and candidate-gene association studies. Many candidates have been proposed, but most were studied in the context of IgAN progression rather than causality. In addition, in part due to our lack of knowledge about disease pathogenesis, most candidates were predicated on sparse a priori evidence for involvement in IgAN. Overall, candidate-gene studies for IgAN have been largely unrevealing. The advantage of GWAS, on the other hand, is that they do not require a priori assumptions about disease pathogenesis. To date six GWAS of familial IgAN have identified several loci located on chromosomes 2, 4, 6, 7, 12 and 17. The functional mapping of genes involved in the disease is anticipated to proceed further from the identification of susceptibility loci identified by linkage analysis to the isolation of candidate genes within gene disease-susceptibility loci. In conclusion, inheritance of IgAN follows a complex genetic pattern, most probably autosomal dominant inheritance with incomplete penetrance. It seems that several genetic loci contribute significantly to the disease susceptibility that underlies the primary immunologic defects observed in IgAN.

Keywords: IgA nephropathy; Genetic studies; Inherited disease; Susceptibility; Familial disease

Introduction
IgAN has been considered for long a benign disease. Today it is clear that as many as 30-40% of the patients develop renal disease, with ensuing significant social and financial impact. In western societies, IgAN is responsible of as much as 7-10% of End-Stage Renal Disease (ESRD) patients on substitution treatment. IgAN occurs at any age, but most commonly the age of onset is in the second or third decade of life. Males are more often affected than females, with a male-female ratio of 2:1. Most patients with IgAN present microscopic hematuria with or without mild proteinuria. About 40% of patients have episodes of macroscopic hematuria. This is sometimes preceded by infections, most commonly upper respiratory tract infections, a phenomenon known as “syphymyngitic” hematuria. Other infections like gastrointestinal or urinary tract infections have also been reported to precede macroscopic hematuria. Proteinuria is common and can vary from mild proteinuria to nephrotic syndrome. Recurrence of IgAN after renal transplantation is common. This finding, along with the observation that IgA deposits disappear from a kidney of an IgAN patient, after transplantation of this kidney to a non IgAN patient, are suggestive of a rather systemic disease.

Genetic Basis of IgAN
The strongest evidence for the existence of genetic factors in the development and/or progression of IgAN comes from descriptions of familial IgAN [1] whereas, interestingly, recent studies suggest that familial and sporadic IgAN may share a common pathogenic mechanism and similar outcomes [2]. The genetic susceptibility may be independent of environmental factors and may reflect an inherited susceptibility to develop mesangial glomerulonephritis. IgAN does not exhibit classic single-gene Mendelian inheritance pattern [3]. The complex genetic pattern of IgAN is reflected by the multiple pathways involved in its immunopathogenesis, namely multiple discrete immunologic abnormalities related to the abnormal overproduction and release of mucosal polymeric IgA1 (pIgA1) in the systemic compartment and possibly other protein functional abnormalities related to a propensity for mesangial deposition of pIgA1. It is therefore probable that the disease is actually caused by specific Single-Nucleotide Polymorphism (SNP) alleles in non-coding regions or synonymous SNPs in coding regions at identified IGAN loci rather than by “classic” nonsense/missense/splice site mutations and deletions/insertions. Therefore, the actual problem is probably the alterations by SNPs of the transcriptional activity of a disease gene and/or messenger RNA stability and, therefore, of the expression level of the encoded protein. It is interesting that recent studies indicate that 30% to 50% of human genes with coding SNPs can present allelic variation in gene expression [4,5]. To this context, the most comprehensive assumption is that several genetic loci convey the primary immunologic defects observed in IgAN and, therefore, contribute significantly to the disease susceptibility. Each locus may occur at a different prevalence rate in different racial/ethnic groups. Variations at these major genetic loci may not be sufficient for the development and progression of IgAN and the contribution from a potentially large number of modifying genes with modest genetic effects but high prevalence is probably needed as well. The various allelic combinations of these loci may underlie the different disease phenotypes (disease development and progression, nephritic vs. nephrotic clinical presentation, histopathologic subclass, severity of disease, responsiveness of proteinuria to Angiotensin-Converting Enzyme (ACE) inhibitors and/or Angiotensin II Receptor Blockers (ARBs), etc.) observed in IgAN. For diseases with complex genetic pattern, it has been shown that the optimal analysis approach is the combination of linkage, association and sequence approaches. Until now, two basic approaches have been used in genetic studies of IgAN: a) genome-wide linkage analysis study, a methodology that has
been used successfully to identify major disease/susceptibility loci, b) candidate-gene association study, mainly used to identify altered genes with modest genetic effects but high prevalence.

However, it is important to realize that genetic studies of IgAN are faced with significant challenges. Thus, familial forms of IgAN are frequently underrecognized because the associated urinary abnormalities in affected family members are often mild or intermittent. Moreover, once familial disease is documented, systematic screening by renal biopsy cannot be justified among asymptomatic at-risk relatives, necessitating reliance on less accurate phenotypes, such as microscopic hematuria, to diagnose affection. Additionally, IgAN has been observed to co-occur in families with Thin Basement Membrane Disease (TBMD), an autosomal dominant disease caused by heterozygous mutations in the collagen type IV genes (COL4A3/ COL4A4) [6]. Short of kidney biopsy or direct sequencing of the very large collagen genes, TBMD cannot be reliably excluded among relatives of IgAN patients. Finally, because urinary abnormalities may manifest intermittently, one also cannot unequivocally classify at-risk relatives as unaffected, necessitating affected-only linkage analysis. The inability to classify affected and unaffected individuals accurately is commonly encountered in linkage studies of complex traits, leading to decreased study power. Increasing sample size by including additional families is also not necessarily helpful in these situations because the diagnosis of IgAN likely encompasses several disease subsets, such that expansion to larger sample size can paradoxically reduce analytic power due to increased heterogeneity [7,8].

Genome-wide linkage analysis studies

Genome-wide linkage analyses involve recruitment of families with multiple affected individuals and are used successfully to identify major disease/susceptibility genes but have limited power to detect genes of modest effect. The advantage of genome-wide linkage studies is that they do not require a priori assumption about disease pathogenesis. These studies are very sensitive to phenotype misspecification; however their power is limited to detecting rare genetic variants with a relatively large effect on the risk of disease. To date, six genome-wide linkage studies of familial IgAN have been reported [9-15]. Families in these studies have all been ascertained via at least two cases with biopsy-documented IgAN, with additional family members diagnosed as affected based on clinical evidence (renal failure or multiple documentation of hematuria/proteinuria). In the first study, 30 families with two or more affected members were examined [9]; multipoint linkage analysis under the assumption of genetic heterogeneity yielded a peak LOD score of 5.6 on chromosome 6q22-23 (locus named IGAN1), with 60% of families linked. The remainder of families linked to chromosome 3p24-23 with a LOD score of 3.47. Most reported linkage intervals did not contain obvious candidate genes, but the 2q36 locus encompasses the COL4A3 and COL4A4 genes, which are mutated in TBMD. Together with the high penetrance of hematuria, this finding suggests that affected individuals in the 2q36-linked family may belong to an IgAN subtype that overlaps with TBMD. The fourth genome-wide analysis, carried out in a cohort of IgAN patients selected from the UK Glomerulonephritis DNA Bank, the region of the MHC (Major Histocompatibility Complex) [13]. The strongest association signal included a combination of DQ loci and HLA-B. This study suggests that HLA region contains some alleles that predispose to the disease in the UK population. In conclusion, four genome-wide linkage studies, carried out in four different IgAN patient populations, demonstrate different chromosomal traits linked to the disease. Recently, in a GWAS conducted by Ghairavi et al. [14] in Chinese and European IgAN patients five loci were traced (3 loci in the Major Histocompatibility Complex (MHC) region on chromosome 6p21, a common deletion CCHGR1 and CFHR3 on chromosome 1q32 and a locus on chromosome 22q12 to be related with IgAN and account for the 4.7% of the genetic heterogeneity of the disease. Finally, the most recent GWAS was conducted by Yu et al. [15], who performed a two-stage genome-wide association study of IgA nephropathy (IgAN) in Han Chinese, with 1,434 affected individuals and 4,270 controls in the discovery phase and follow-up of the top 61 SNPs in an additional 2,703 cases and 3,464 controls. They identified associations at 17p13 and 8p23, that implicated the genes encoding tumor necrosis factor (TNFSF13) and a-defensin (DEFA1) as susceptibility genes. In addition, they found multiple associations in the HMC region and confirmed a previously reported association at 22q12. They also found that rs660895 was associated with clinical subtypes of IgAN, proteinuria and IgA levels. These findings are consistent with the hypothesis that IgAN is associated with variants near genes involved in innate immunity and inflammation.

Candidate-gene association studies

Candidate-gene association studies examine polymorphisms in only specific genes that are selected based on a priori assumption about their involvement in the disease pathogenesis, and they are highly sensitive to population stratification, multiple testing, and...
the Megsin 23167G SNP variant is associated with both susceptibility [26]. The same group using a similar approach recently reported that these same SNP alleles were more often transmitted to patients

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In general, most of these studies were of poor quality and severely underpowered and, therefore, negative findings were almost universally inconclusive. Thus far, only one group attempted to survey the entire genome, yet the results have not been replicated, are inconclusive and difficult to interpret, as long as an underpowered cohort was studied with inadequate coverage of ~80,000 SNPs [24,25]. Moreover, 77% of all published candidate-gene studies reported positive findings, an observation that is likely explained by a combination of high rate of false positives and a strong publication bias. Most findings were not reproduced in other populations. Overall, to date, very few studies examining candidate genes have employed the family-based Transmission Disequilibrium Test (TDT) methodology, or the analysis of “at-risk” haplotypes or the Family-Based Association Test (FBAT), reflecting the emergence of the first studies to attempt to satisfy minimum criteria for a valid association study. A family- and haplotype-based association study employing the TDT methodology has shown that 2093C and 2180T SNP variants in the 3'-untranslated region of the Megsin gene were significantly more frequently transmitted from heterozygous parents to patients than expected in the extended TDT analysis (increased co-transmission in 232 Chinese families). In addition, Haplotype Relative Risk (HRR) analyses showed that these same SNP alleles were more often transmitted to patients [26]. The same group using a similar approach recently reported that the Megsin 23167G SNP variant is associated with both susceptibility and progression of IgAN in 435 Chinese patients and their family members using TDT and HRR analyses [27]. The GG genotype was found to be associated with severe histologic lesions and disease progression. Megsin is a member of the serpin (serine protease inhibitor) superfamily that is upregulated in the context of mesangial proliferation and extracellular matrix expansion in IgAN, and therefore represents a strong candidate gene for susceptibility to IgAN. Lately, the gene encoding Cosnc, a C1Gal-T1 chaperone protein which also mediates IgA O-galactosylation was studied as a candidate gene involved in the pathogenesis of IgAN but no evidence for a role for Cosnc mutations was reported [28].

A family-based association study performed by IgAN Consortium, including 53 patients, 45 complete trios, 4 incomplete trios and 36 discordant siblings, evaluated the impact of some Th1/Th2/Th3/ TR-type lymphocyte and monocyte/macrophage cytokines on IgAN susceptibility [29]. Cytokine gene polymorphisms with a potential regulatory role on their production were investigated using the FBAT: IFNγ in intron-1 CA-repeats at position 1349–1373; IL-13 -1055C/T; TGFβ 915G/C; IL-10 5’-proximal and distal microsatellites; TNFα –308G/A, –238G/A. The FBAT multi-allelic analysis showed an association between IFNγ polymorphism and susceptibility to IgAN, whereas the bi-allelic analysis showed that the 13-CA repeat allele was preferentially transmitted to the affected individuals. The in vitro production of IFNγ evaluated in PBMC from 10 genotyped patients demonstrated a correlation between the +874A allele and a lower production of IFNγ.

Furthermore, consistent with the known role of NF-xB in the transcriptional regulation of the INFγ gene, the +874A variant was associated with transcriptional downregulation of INFγ gene promoter activity. The occurrence of the +874A variant is responsible for the low production of IFNγ and predisposes to a preferential Th2-mediated immune response. The predominance of this variant in individuals with IgAN may be responsible for the onset of the disease. This unbalanced Th2 cytokine production in response to upper respiratory tract infections may be a significant pathogenic factor in human IgAN. The prevalence of the Th2 cytokines may also explain the abnormality in IgA1 glycosylation occurring in IgAN patients and the concomitant formation of circulating IgA1-IgG immune complexes. Hyperfunction of the Th2 cells and cytokine polarity are linked to a more nephrinotropic pattern of IgA1 glycosylation in the animal model, and the decreased glycosylation of IgA1 elicited by Th2 cytokines is blunted in vitro by the addition of IFNγ [30]. The core 1 β1,3-galactosyltransferase (C1Gal-T1) is suspected to be involved in the abnormal glycosylation process of IgA1 in IgAN. With the genetic characterization of the enzymes responsible for O-glycosylation of IgA1, it has been possible to study changes in the O-glycosylation of IgA1 at a genetic level. Most recently two groups [31,32] have independently found that SNPs in the C1Gal-T1 gene (genotype 1365G/G) confer genetic susceptibility to IgAN, yet not to the progression of the disease as well, in Chinese and Italian populations, albeit it is not clear how these polymorphisms relate to changes in the functional activity of C1Gal-T1.

Clinical Implications

The above results provide further evidence for genetic heterogeneity among IgAN families. Evidence of linkage to multiple chromosomal regions is consistent with both an oligo/polygenic and a multiple susceptibility gene model for familial IgAN with small/moderate effects in determining the pathological phenotype. Susceptibility to IgA nephropathy has a genetic component to varying degrees, with significant diversity between different geographical areas. Familial forms of IgAN may derive from a mutation of a specific gene occurring
in a founder several generations ago. Although several genetic loci are strongly associated with IgAN, the implication of the "guilty" genes is yet far from being well documented. It seems probable that the disease is inherited as an autosomal-dominant trait with highly variable penetrance.

But what about the vast majority of IgAN cases which seem to be sporadic? Although many of these cases may actually have a less apparent genetic basis, the existing evidence is rather deficient on this issue. To what extent genetic and environmental factors contribute to the pathogenesis of IgAN is still not known with accuracy. At present, genetic testing does not appear to have much diagnostic value except in clear-cut familial cases, because many loci are involved, and renal biopsy remains the golden diagnostic standard. It is noteworthy that most genetic studies actually examined susceptibility to the clinical expression of disease rather than susceptibility to the mesangial IgA deposition itself. On the basis of these preliminary findings, however, the extended genetic studies under way are anticipated to shed light to the pathogenetic mechanisms and allow for the earlier diagnosis of the disease or even the genetic susceptibility for the disease, more efficient preventive policies and more targeted and fruitful treatment in the near future.

Conclusions

Despite the fact that clear-cut genetic susceptibility can be found in the minority of IgAN cases, this might be attributed at least in part to its complex genetic pattern, most probably autosomal dominant pattern with incomplete penetrance. Incomplete penetrance might be attributed to the need for the presence of additional genetic and/or environmental factors for the clinical manifestation of the disease. To date several loci have been associated with IgAN by different groups, although there is still long road ahead to document definitely the implication of the respective candidate genes. These preliminary findings, however, have created great expectations that the extended genetic studies under way with the use of high technology high throughput assays will shed light to pathogenetic mechanisms of the disease and allow the earlier diagnosis of the disease or even the genetic susceptibility for the disease, more efficient preventive policies and more targeted and fruitful treatment in the near future.

References


