The importance of the polymorphisms of the ABCB1 gene in disease susceptibility, behavior and response to treatment in inflammatory bowel disease: A literature review

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A – research concept and design; B – collection and/or assembly of data; C – data analysis and interpretation; D – writing the article; E – critical revision of the article; F – final approval of the article

Abstract

Crohn's disease (CD) and ulcerative colitis (UD) are the 2 common clinical subtypes of idiopathic inflammatory bowel disease (IBD) characterized by chronic inflammation of the gastrointestinal tract. The multifactorial etiology and pathogenesis of IBD is still unknown; however, the interaction between genetic, environmental and immunological factors seems to be crucial. A member of the adenosine triphosphate (ATP)-binding cassette family, P-glycoprotein, encoded by the human ABCB1 gene, is among the most extensively studied transporters involved in drug disposition and effects. Single nucleotide polymorphisms (SNPs) located in exons 21, 26 and 12, i.e., G2677T/A, C3435T and C1236T, are of the greatest clinical importance. Functional defects of the intestinal epithelial barrier due to the lack of P-glycoprotein expression may constitute possible reasons for the development of colitis. Given that several drugs central to the therapy of IBD are also P-glycoprotein substrates, it has been hypothesized that its altered expression in IBD patients could modify the response to medical treatment. Nevertheless, there are conflicting reports of an association between these 3 SNPs and IBD. This article aims to review all relevant studies investigating the role of the polymorphisms of the ABCB1 gene in disease susceptibility, behavior and response to treatment in IBD.

Key words: inflammatory bowel disease, MDR, P-glycoprotein

DOI

10.17219/acem/92936

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Introduction

Crohn’s disease (CD) and ulcerative colitis (UC) are the 2 common clinical subtypes of idiopathic inflammatory bowel disease (IBD) characterized by chronic inflammation of the gastrointestinal tract. They may occur in various geographic regions, however, with different frequency. They affect mainly citizens of developed countries from Europe and North America. A large multicenter study conducted in 1991–1993 in 12 European countries estimated the incidence rate of UC and CD to be 10.4 and 5.6 per 100,000 person-years, respectively. The incidence of IBD may have been changing over time, while the incidence of UC appears to remain stable, and a rise in incidence of CD has been observed. The peak incidence of IBD occurs between 15 and 40 years of age, with a possible 2nd peak between 50 and 80 years of age.

Ulcerative colitis is characterized by recurring episodes of inflammation limited to the mucosal layer of the colon. It involves the rectum and may spread proximally in a continuous fashion. Crohn’s disease may involve the entire gastrointestinal tract from mouth to perianal area and is characterized by transmural inflammation and skip lesions, i.e., lesions appearing in a place surrounded by healthy mucosa. The transmural inflammation may lead to fibrosis and strictures as well as to microperforations and fistulae. Patients with UC usually present with bloody diarrhea. Associated symptoms include colicky abdominal pain, urgency, tenesmus, and fecal incontinence. Crampy abdominal pain, prolonged diarrhea and weight loss are the hallmarks of CD. Gross bleeding is less frequent than in UC.

The underlying pathogenesis remains unclear but may involve persistent bacterial infection, defective mucosal barrier and imbalance in the regulation of the immune system. Epidemiological and family studies suggest that genetic factors play a significant role in determining susceptibility to IBD. Approximately 10–25% of individuals with IBD have a first degree relative with either CD or UC. In a study conducted among Danish twins with IBD, the concordance rate among monozygotic pairs was 58.3% for CD and 18.2% for UC. Inflammatory bowel disease appears to follow a non-Mendelian pattern of inheritance. It is likely that the aggregate effect at several loci contributes to the IBD phenotype. The first identified gene associated with CD was NOD2/CARD15 gene (nucleotide-binding oligomerization domain, caspase recruitment domain), localized on chromosome 16. Three mutations in the sequence of the NOD2/CARD15 gene, Arg702Trp, Gly908Arg and Leu1007fsinsC, appeared to be the factors strongly related to CD. Wild-type NOD2 protein activates nuclear factor kappa B, making it responsive to bacterial lipopolysaccharides; this induction is deficient in patients with the mutant form of NOD2. Patients with CD, who are the carriers of at least 1 of the 3 NOD2/CARD15 gene mutations, are at a higher risk for early onset and development of stenosis and small intestine involvement.

The functional properties of the proteins encoded by the genes implicated in the susceptibility to IBD have enabled the identification of specific pathways important in the pathogenesis of IBD: ATG16L1 (the autophagy pathway), interleukin (IL)-17 and IL-23 receptor genes (pathways regulating adaptive immunity), and OCTN2 (the pathway regulating the epithelial function).

P-glycoprotein has been the most extensively studied member of the adenosine triphosphate (ATP)-binding cassette superfamily encoded by the human ABCB1 gene (previously known as MDR1). P-glycoprotein is a phosphorylated and glycosylated protein that consists of 1280 amino acids and 2 homologous and symmetric sequences, each containing 6 transmembrane domains (TMDs), and a nucleotide-binding domain (NBD). P-glycoprotein functions as a transmembrane efflux pump, thereby moving drugs from the intracellular to the extracellular domain. Adenosine triphosphate hydrolysis provides the energy for active drug transport against steep concentration gradient.

P-glycoprotein is expressed in several human tissues, including peripheral blood lymphocytes, epithelial cells in the small and large bowel, hepatocytes, pancreatic ductile cells, kidneys, adrenal glands, the epithelium of the brain choroids plexus, the capillaries of the brain, placenta, etc. Thus, it plays an important role in the excretion of xenobiotics and endogenous substrates via the canalicular membrane of hepatocytes into the bile, via the brush border membrane of enterocytes into the gut lumen and via the brush border membrane of proximal tubules into the urine. In the endothelial cells of the blood-brain barrier, P-glycoprotein prevents the entry of substrates into the central nervous system.

P-glycoprotein transports a wide range of substances with diverse chemical structures, among them anticancer agents, cardiac drugs (e.g., digoxin, quinidine), HIV protease inhibitors, immunosuppressants (e.g., cyclosporine), and β-blockers. Localized in gastrointestinal tract, it decreases their absorption and bioavailability.

Interestingly, most substrates of P-glycoprotein are also metabolized by the isoenzyme CYP3A4. This is of particular importance, because P-glycoprotein and CYP3A4 are co-localized in the small intestine and liver, organs crucial for absorption, distribution and excretion of drugs.

ABCB1 gene polymorphisms and IBD

The ABCB1 (MDR1) gene is located on chromosomal region 7q21 and consists of 28 exons encoding a protein composed of 1280 amino acids. It has been shown to be highly polymorphic, with 50 single nucleotide polymorphisms (SNPs) and 3 insertion/deletion polymorphisms reported so far. Three SNPs are currently considered to be the most clinically relevant, i.e., in exons 21 (G2677T/A), 26 (C3435T) and 12 (C1236T). Hoffmeyer et al. were the first to demonstrate a 2-fold reduction in P-glycoprotein
expression in duodenal biopsy samples among healthy Caucasian subjects homozygous for the mutant 3435T allele in comparison to subjects homozygous for the C3435 allele (wild-type). TT genotype was also shown to be associated with higher digoxin plasma concentrations after oral administration, suggesting greater drug absorption in individuals with low intestinal P-glycoprotein levels.18 These observations were only in apparent contradiction with the fact that 3435C→T mutation in exon 26 is a synonymous single-nucleotide polymorphism (i.e., does not alter the amino acid encoded). There is a linkage disequilibrium between SNPs in exon 26 (C3435T) and exon 21 (G2677T/A), suggesting that the observed differences in P-glycoprotein, initially attributed to the exon 26 SNP, may be the result of the associated tri-allelic polymorphism in exon 21. The latter is a nonsynonymous single nucleotide polymorphism (i.e., one causing an amino acid change, Ala893Ser/Thr).19 It has also been recently shown that a synonymous SNP in exon 12 (C1236T) is linked to the C3435T and G2677T/A SNPs.20

However, there are conflicting reports on an association between these 3 SNPs and IBD. The ABCB1 gene is an attractive candidate for the pathogenesis of IBD and, perhaps, it may also condition response to therapy. Firstly, it is located in a region of the human genome (7q21) that was found to possibly harbor a disease gene involved in susceptibility to IBD.21 Its role in IBD was then investigated in a mouse knockout model (mdr1a−/−) in which developmentally normal mice spontaneously developed a colitis resembling UC in humans. They also developed a spontaneous colitis when maintained under specific pathogen-free conditions. The colitis was prevented and reversed with the administration of antibiotics, suggesting that the intestinal flora may be the result of the associated tri-allelic polymorphism in exon 21. All this suggests that functional defects of the intestinal epithelial barrier (in terms of both loss of the xenobiotic efflux mechanism and host–bacteria interaction) due to the lack of P-glycoprotein expression are possible reasons for the pathogenesis of colitis.

C3435T polymorphism

In the initial case-control study conducted in Germany by Schwab et al., investigating the C3435T polymorphism, an increase of the T allele and TT genotype frequencies was identified in 149 patients with UC, but not with CD, compared with controls (p = 0.049, odds ratio (OR) = 1.4; p = 0.005, OR = 2.1, respectively).22 This finding has been replicated in several studies but in a handful of other works such observation was not evident. Ho et al. in a study conducted in Scotland reported a statistically significant association between UC and a higher frequency of the mutant T allele (p = 0.02, OR = 1.28) and of TT genotype (p = 0.04, OR = 1.60).23 Similarly, Farnood et al. observed in the Iranian population a higher risk of UC development for the 3435T allele carriers and 3435TT homozygotes.25 In contrast, negative findings have been reported in large studies from Germany and the UK,26 North America,27 Slovenia,28 and Italy.29 Paradoxically, Urcelay et al. found a significant association between the wild CC3435 genotype and CD in Spanish patients (p = 0.007), recognizing the 3435T allele as a risk factor for UC, and the 3435C allele as a risk factor for CD.30 The meta-analysis of 9 studies (1743 UC cases, 2311 CD cases and 2931 controls in total) has confirmed an association between C3435T and UC, with an OR = 1.12 (95% confidence interval (CI): 1.02–1.23). For CD, the pooled ORs were not significant for either the fixed-effect or the random-effects model.31 Similar results, i.e., a significant association of UC with T allele (OR = 1.17, 95% CI: 1.06–1.31) and TT genotype (OR = 1.36, 95% CI: 1.05–1.76), were obtained in another meta-analysis.32

It is worth mentioning that Ho et al. undertook a genotype–phenotype analysis and estimated the risk rate of UC with the proximally spreading lesions (i.e., extensive colitis) as 1.70 (p = 0.009) for the 3435T allele carriers and as 2.64 (p = 0.003) for the 3435TT homozygotes.24 Likewise, Ardizzone et al. found Italians carrying the mutant 3435T allele to present a 3-fold increased risk for developing CD with ileocolonic localization as compared to individuals with the wild-type allele.33

G2677T/A polymorphism

Investigating another polymorphism of ABCB1 gene, the tri-allelic G2677T/A SNP (Ala893Ser/Thr) in a multicenter North American cohort study, Brant et al. found a significant association of the G2677 allele (Ala893), known to decrease P-glycoprotein function, with IBD.27 An association with another allele of this SNP, i.e., 2677T (893Ser), was identified in UC patients (p = 0.029) by Potočnik et al. in Slovenia.28 Conversely, Ho et al. did not find any relation between this polymorphism and IBD.24 In the study by Onnie et al., the 2677T allele was significantly increased in British UC cases compared with controls (45.2% vs 39.6%; p = 0.034). In particular, the TT genotype was significantly associated with severe UC (OR = 1.90; 95% CI: 1.01–3.55) and the use of steroids in UC (OR = 1.77; 95% CI: 1.08–2.88).31 After pooling data from the available studies, the meta-analysis performed by Annese et al. found no significant relation between allele and genotype frequencies of the G2677T/A SNP and UC, as well as between CD and IBD as a whole.32

Linkage disequilibrium and heterogeneity considerations

As significant linkage disequilibrium has been observed between 3 SNPs (G2677T/A, C3435T and C1236T) in different populations, a haplotype analysis seems to be
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found a trend towards an increased frequency of 2-locus 2677T/C3435 haplotype in CD patients. Onnie et al. did not discover any significant C3435T-G2677T/A haplotype association with either CD or UC. The advantage of haplotype analysis in comparison to analysis of single polymorphisms in complex diseases has been confirmed in the study by Potočnik et al. They found haplotype defined by T–T–T (1236T–2677T–3435T) alleles to be significantly associated with higher risk for refractory CD (p = 0.044, OR = 3.1) and UC (p = 0.026, OR = 1.6), although using each of these SNPs separately resulted in association only at the border of significance.

Reasons for the discrepancy among the abovementioned studies may lie in population heterogeneity, sample size, selection of control population, incomplete phenotype description, and the lack of statistical power to detect the moderate effect size. It has been proven that there is a significant heterogeneity of the allele frequencies in various populations. For example, the frequency of the C3435 allele has been reported as 43–54% in Caucasians, 34–63% in Asians and 73–90% in Africans. The incidence of C/T and C/C3435 genotypes in Africans is much higher than in other racial populations.

Response to pharmacotherapy

Of interest, several drugs central to the therapy of IBD are also P-glycoprotein substrates like glucocorticoids, cyclosporine and methotrexate. The hypothesis that altered P-glycoprotein expression in IBD patients could modify the response to medical therapy was verified by Farrell et al. Compared with controls, the expression of P-glycoprotein in peripheral blood lymphocytes was significantly elevated in patients with CD who required bowel resection and patients with UC who required proctocolectomy for failed medical therapy. However, Farrell et al. did not investigate the role of genetic variants of the ABCB1 gene.

Conversely, Palmieri et al., who compared allele and genotype frequencies in 594 patients using systemic steroids and 297 patients taking immunosuppressive drugs, did not find any influence of both C3435T and G2677T/A polymorphisms on the response to therapy.

Polish population

Two studies have recently been published investigating the importance of C3435T and C1236T polymorphisms in determining IBD susceptibility in a population from central Poland. The 1st, evaluating the C3435T polymorphism, comprised 85 patients with IBD (45 with UC and 40 with CD) and 70 healthy volunteers. The 2nd, evaluating the C1236T polymorphism, comprised 85 patients with IBD (45 with UC and 40 with CD) and 70 healthy volunteers. The identification of the polymorphisms in the ABCB1 gene was carried out using the polymerase chain reaction – restriction fragment length polymorphism (PCR-RFLP) method. In both studies, the observed differences in genotype and allele frequencies were not significant. However, 3435CC genotype and 3435CC allele carriers were present more frequently among IBD patients than in controls (OR = 1.72, 95% CI: 0.95–3.12 and OR = 1.35, 95% CI: 0.94–1.93, respectively).

In parallel, 1236CT genotype and 1236T allele carriers were more frequent in IBD patients compared to controls (OR = 1.26, 95% CI: 0.66–2.42 and OR = 1.08, 95% CI: 0.41–2.14, respectively). Neither G2677T/A polymorphism nor the impact of G2677T/A, C3435T and C1236T polymorphisms on disease behavior and response to therapy in IBD have been explored so far in Poland. It is worth noting that even in our country a substantial heterogeneity of the allele frequencies in relation to the ABCB1 gene has been observed. For example, the frequency of the 3435CC, 3435CT and 3435TT genotype was 23.3%, 56.3% and 20.4%, and 42.0% and 17.0%, respectively, in subjects originating from Western Pomerania and Łódź region. There has been so far no study determining the importance of the polymorphisms of the ABCB1 gene in pediatric onset IBD. It seems equally challenging to evaluate the ABCB1 gene expression in the tissues of the gastrointestinal tract, because, as was stated by Yacyshyn et al., intra-epithelial, lamina propria and peripheral blood lymphocytes may demonstrate different gene expression and activity of its product, P-glycoprotein.

References


