Immunogenetic phenotypes in inflammatory bowel disease

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Abstract
The currently accepted etiopathogenic hypothesis suggests that the chronic intestinal inflammation and related systemic manifestations characteristic of inflammatory bowel disease (IBD) are due to an overly aggressive or pathologic immune response to resident luminal bacterial constituents. Predisposing factors are genetic dysregulation of mucosal immune responses and/or barrier function, with onset triggered by environmental stimuli. These factors and their interactions may also be important determinants of disease phenotype and disease progression. The interaction between genetic susceptibility and luminal bacteria deserves particular attention. The results from animal studies indicate that not all commensal bacterial species are identical in their abilities to induce disease in the face of a common genetic defect. For instance, HLA B27/β2 microglobulin transgenic rats raised under sterile conditions do not develop colitis. However, when colonized with normal specific pathogen-free commensal cecal bacteria, they develop aggressive disease involving the colon and gastro-duodenal area within one month. On the one hand, when colonized with a single bacterial strain, Bacteroides vulgatus, these same genetically susceptible rats develop moderate colitis but no gastro-duodenal disease. When selectively colonized with Escherichia coli, they exhibit neither disease nor T cell activation. This data suggests that not all commensal bacterial strains trigger an abnormal immune response for a given genetic susceptibility. The same non-pathogenic bacteria, however, may induce inflammation in a different genetically susceptible host. More specifically, further research from the same group demonstrated that Escherichia coli induced only mild cecal inflammation after 3 wk of monoassociation in interleukin 10-/- mice. In contrast, Enterococcus faecalis-monoassociated interleukin 10-/- mice developed distal colitis at 10-12 wk that was progressively more severe and associated with duodenal inflammation and obstruction by 30 wk. Their results suggest that different commensal bacterial species selectively initiate immune-mediated intestinal inflammation with distinctly different kinetics and anatomic distribution in the same host. These findings highlight the fact that both genetic susceptibility and luminal antigenic drive are important determinants of disease susceptibility and modification. Immune reactivity as measured by the serological expression of immune responses to specific bacteria may indeed be representative of this host gene luminal bacterial interaction. If indeed these immune responses represent the sum of a genetic and environmental predisposition to IBD, quantitative and qualitative expression of these immune responses may serve as an immunologic risk marker for IBD phenotypes.

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Key words: Inflammatory bowel disease; Immune reactivity; Disease phenotype

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INTRODUCTION
The currently accepted etiopathogenic hypothesis suggests that the chronic intestinal inflammation and related systemic manifestations characteristic of inflammatory bowel disease (IBD) are due to an overly aggressive or pathologic immune response to resident luminal bacterial constituents. Predisposing factors are genetic dysregulation of mucosal immune responses and/or barrier function, with onset triggered by environmental stimuli. These factors and their interactions may also be important determinants of disease phenotype and disease progression. The interaction between genetic susceptibility and luminal bacteria deserves particular attention. The results from animal studies indicate that not all commensal bacterial species are identical in their abilities to induce disease in the face of a common genetic defect. For instance, HLA B27/β2 microglobulin transgenic rats raised under sterile conditions do not develop colitis. However, when colonized with normal specific pathogen-free commensal cecal bacteria, they develop aggressive disease involving the colon and gastro-duodenal area within one month. On the one hand, when colonized with a single bacterial strain, Bacteroides vulgatus, these same genetically susceptible rats develop moderate colitis but no gastro-duodenal disease. When selectively colonized with Escherichia coli, they exhibit neither disease nor T cell activation. This data suggests that not all commensal bacterial strains trigger an abnormal immune response for a given genetic susceptibility. The same non-pathogenic bacteria, however, may induce inflammation in a different genetically susceptible host. More specifically, further research from the same group demonstrated that Escherichia coli induced only mild cecal inflammation after 3 wk of monoassociation in interleukin 10-/- mice. In contrast, Enterococcus faecalis-monoassociated interleukin 10-/- mice developed distal colitis at 10-12 wk that was progressively more severe and associated with duodenal inflammation and obstruction by 30 wk. Their results suggest that different commensal bacterial species selectively initiate immune-mediated intestinal inflammation with distinctly different kinetics and anatomic distribution in the same host. These findings highlight the fact that both genetic susceptibility and luminal antigenic drive are important determinants of disease susceptibility and modification. Immune reactivity as measured by the serological expression of immune responses to specific bacteria may indeed be representative of this host gene luminal bacterial interaction. If indeed these immune responses represent the sum of a genetic and environmental predisposition to IBD, quantitative and qualitative expression of these immune responses may serve as an immunologic risk marker for IBD phenotypes.
Immune responses to resident intestinal flora in humans have been reported. Duchmann et al demonstrated that CD patients boast reactivity to hundreds of bacterial antigens created from sonification of multiple bacterial species including enterobacteria, bacteroides and bifidobacterium. Immune reactivity to more specific microbial antigens has been reported in the sera of patients with IBD (Table 1).

Antibodies to *Saccharomyces cerevisiae* (ASCA) was the first CD specific immune response thought to be targeted towards microbial antigens. IgA and IgG antibodies are directed against a specific oligomannosidic epitope present on the cell wall of the yeast *Saccharomyces* [9]. To date it remains unknown as to what the specific microbial antigen ASCA is cross reacting with and giving rise to seropositivity specifically in the sera of patients with CD. ASCA is present in approximately 60% of CD patients, yet less than 5% in UC and non-IBD patients [6,7]. The specificity of ASCA renders a positive test result accurate in differentiating CD from UC and IBD from non-IBD in cases of diagnostic uncertainty. ASCA also remains an important marker of disease severity as defined by the development of complicating disease. More recent research has resulted in the identification of 3 additional markers representative of microbial driven immune responses, antibodies to the E.coli outer-membrane porin C (OmpC), the Pseudomonas fluorescentes CD related protein (anti-CD related bacterial sequence {I2}) and the CBir1 flagellin.

Antibodies to OmpC, whose antigen is purified from commensal *E.coli* [9,10], are present in 37%-55% of patients with CD and 2%-11% of patients with UC, while no more than 5% of non-IBD individuals express anti-OmpC [11-13]. I2 was isolated from affected colonic mucosa yet not in the unaffected segments and thus thought to be specific to CD [5]. Immune responses to this antigen are present in up to 55% of CD patients; however the specificity of this antibody has been questioned as it has been detected in the serum of UC patients (10%) and more impressively in up to 55% of CD patients; however the specificity of this antibody has been questioned as it has been detected in the serum of UC patients (10%) and more impressively in up to 20% of non-IBD patients (9, data on file Prometheus Labs). Given its limitation as a diagnostic marker, it is unclear as to whether this immune response will provide additional prognostic information and further studies are needed. Serologic expression cloning was used to identify an immunodominant antigen, CBir1 flagellin, to which strong immune responses (B cell and CD4T cell) occurred in colitic mice [14]. These findings were translated from the bench to the bedside and approximately 50% of CD patients had serum reactivity to CBir1 whereas UC, inflammatory and healthy controls exhibited little to no reactivity to this flagellin [15].

All of the prior immune responses are specific to CD. In contrast perinuclear anti-neutrophil antibody (pANCA) is noted for its association with UC or a UC like phenotype. This IBD-specific ANCA displays a unique perinuclear highlighting (pANCA) on immunoflourence staining and is DNAse sensitive [10]. Although it remains undefined, it has been suggested that the antigen to which pANCA is directed is a nuclear histone (H1) [17]. This antigen is clearly distinct from the proteasine 3 or the myeloperoxidase reactivity observed in those patients with vasculitic disorders. pANCA is likely an autoantibody that is representative of a cross-reactivity with a luminal bacterial antigen [10,18,19]. Despite epidemiological and methodological differences, pANCA has been shown repeatedly to be prevalent in the sera of approximately 60% and 20% of UC and CD patients, respectively [8,20-22].

Typically, < 5% of non-IBD patients are pANCA positive.

As seen with the genetic and clinical heterogeneity of CD, studies have shown immune response (immune phenotype) heterogeneity exists among CD patients. Landers et al analyzed immune response heterogeneity in 330 patients and found that ASCA was detected in 56% of patients; 55% were seroreactive to OmpC. 50% were seroreactive to I2, and 23% were pANCA [9]. Eighty-five percent responded to at least 1 antigen; only 4% responded to all 4. Among microbial antigens (ASCA, OmpC, I2), 78% responded to at least 1, and 57% were double positive, but only 26% responded to all 3. The level of response was stable over time and with change in disease activity. Among patients with the same qualitative antigen-response profiles, quantitative response differed. Moreover this study demonstrated that CD patients could be clustered into 4 distinct groups depending on their immune response patterns to microbial or autoantigens. One cluster was ASCA, a second was antibodies to OmpC and I2, the third pANCA and the fourth was low or no immune response to any tested antigens.

Subsequent analyses incorporating CBir1 demonstrated that antibodies to CBir1 are present in approximately 40% of CD patients negative for antibodies to specific microbial antigens (ASCA, OmpC and I2) which suggests a unique immune phenotype [14]. Immune reactivity to CBir1 may further define CD phenotypes in that anti-CBir1 expression is present in 40%-44% of pANCA positive CD patients only 4% in pANCA positive UC patients. This difference may denote a unique etiopathogenic mechanism of disease that helps to further stratify patients based on immunogenetic phenotypes.

### CLINICAL AND IMMUNE PHENOTYPES

Disease phenotype is not always a static phenomenon. Retrospective studies have examined the stability of disease phenotypes over the course of disease from time of diagnosis until point of last follow-up. It appears that disease location essentially remains stable over time, yet disease behavior evolves, such that after 20-yr of follow up, at least 80% of patients with originally non-complicating disease progress to complication, either
The frequency of disease behavior. The test for trend demonstrated a positive linear trend in the frequency of patients with IP and/or S disease as the number of positive immune responses toward I2, OmpC, ASCA and CBir1 increased ($P = 0.002$). The odds ratios (OR) reflect the odds of having internal penetrating and/or stricture disease when positive for any 1, combination of 2, 3 or all 4 immune responses, as compared to those patients negative for all immune responses (baseline group).

penetrating or stricture in nature. These findings suggest that non-complicating (non-penetrating, non-stricture) disease behavior may not be a stable phenotype but just a temporary state that evolves into one of the two complicating disease states over time.

Immune responses were first investigated as tools to differentiate UC from CD given the specificity of ASCA for CD and pANCA for UC. Advances in the sensitivity of the test characteristics lead to studies evaluating antibodies as diagnostic tools to differentiate IBD from non-IBD. Although conflicting, studies do support the use of these markers, particularly in children, to guide clinicians in cases of diagnostic uncertainty. As new markers are identified and the test characteristics improve, the notion of immune responses optimizing diagnostic accuracy may become more clearer and clinically valid. It has become clear, however, that immune responses may also have perhaps a more important mechanistic implication in the pathogenesis of IBD. As alluded to above, these immune reactivities, as measured by the serological expression of immune responses to specific bacteria, may be representative of the host gene luminal bacterial interaction characteristic of IBD. Moreover if these immune responses represent the sum of a genetic and environmental predisposition to IBD, quantitative and qualitative expression of these immune responses may serve as an immunologic risk marker for IBD phenotypes. The initial immune clinical phenotype studies demonstrated that although pANCA has been established as a UC-specific marker, approximately 25% of all CD patients also express pANCA. These CD patients are described as "UC-like" and tend to have an uncomplicated disease course. In contrast, higher ASCA levels were also shown to be associated with earlier age of disease onset, both strictureting and internal penetrating disease behaviors and need for small bowel surgery. Further reports have found that patients with Crohn's disease who are positive for ASCA IgA, IgG, or both, may define a subset of patients with Crohn's disease at increased risk for early surgery. ASCA has also been shown to be associated with a more aggressive disease course among a cohort of pediatric CD patients. Anti-CBir1 has also been shown to be independently associated with complicated disease behaviors in adult CD patients. Of interest is that these associations were not found in a pediatric CD cohort suggesting potential differences in the mechanistic influence of CBir1 in the pediatric age group.

These studies also demonstrated that both the number of immune responses to the different microbial antigens expressed by a given individual as well as the magnitude (titer level) of these immune responses correlated most significantly with the presence of complicated CD phenotypes. Of interest is that these associations were not found in a pediatric CD cohort suggesting potential differences in the mechanistic influence of CBir1 in the pediatric age group.

IMMUNOGENETIC PHENOTYPES AND NATURAL HISTORY OF DISEASE

The prospective acquisition and determination of immune reactivity prior to the development of a disease complication is important to be able to truly evaluate the potential predictive value of immune responses in defining the natural history of IBD. It is generally agreed upon
that the presence and level of immune responses do not change in a given CD patient. We recently published the first prospective evaluation of the prognostic value of immune responses in patients presenting with non complicated CD. This study included only those patients not presenting with internal penetrating and or strictureting disease at diagnosis and continued to be uncomplicated at the time sera was collected for immune response measurement so that it could be ascertained that the complication occurred after the antibodies were measured. Among those who developed a complication during the follow-up, the median time from diagnosis to the onset of the complication was 48 months. As of the last follow up of this cohort, 8% of the group seropositive to at least one serological marker had developed a complication at only 3% in the seronegative group. Survival analysis (Figure 3) demonstrated that among those patients positive for at least one serology, more progressed to internal penetrating and or strictrueing disease than those negative for all serologies. Saying it differently, those patients positive for at least one serology progressed faster that those negative for all serologies. Determining the factors that can predict the progression from uncomplicated to complicated disease states may stratify patients into at risk populations and impact the ultimate therapeutic management of patients with the goal of halting or more importantly preventing progression to complicated behaviors.

**FAMILIAL EXPRESSION OF IMMUNE PHENOTYPES**

There has been interest in evaluating whether immune responses are familial traits due to genetic factors. ANCA expression was the first marker to be studied. Several studies have observed an increased frequency of pANCA in unaffected relatives of UC patients but not environmental controls. This relationship however has not been observed in all studies. Sutton et al demonstrated that the quantitative and qualitative expression of ASCA was familial. Another study confirmed that ASCA occur particularly frequently in CD patients, especially with the presence of a positive family history. However, they are also significantly increased in UC patients with a family history and in a considerable number of unaffected relatives of inflammatory bowel disease families, irrespective of the characteristics of their families (UC, CD, mixed, ASCA positive, and ASCA negative). The presence of ASCA in unaffected relatives might point towards a genetic predisposition to either CD or UC. Studies in a twin population demonstrated an agreement in ASCA titers within concordant monzygotic twin pairs with Crohn’s disease and suggested that the level of increase in ASCA reactivity(magnitude) is genetically determined. None of these studies evaluated the influence of ASCA on the natural history of the affected offspring. A very recent study has shown that antibodies to OmpC have a strong familial aggregation pattern. In this study, expression of anti-OmpC in unaffected family members of multiplex IBD families were compared with controls. As shown in figure 4, unaffected family members in CD only families had a similar frequency of anti-OmpC to those in mixed families. However, when comparing these two groups with healthy controls, both of them showed a much higher frequency of anti-OmpC expression. In addition, unaffected family members in these two groups displayed a greater prevalence of anti-OmpC than those unaffected from UC only families. This study also addressed whether ASCA and OmpC were co-segregating in patients and unaffected relatives. The results however suggest that anti-OmpC and ASCA may represent related but distinct immune responses in CD patients. More importantly the distinct immune phenotypes may translate into distinct clinical phenotypes.

**SUMMARY**

In summary, the emergence of immunogenetic phenotypes lends support to the proposed hypothesis that susceptibility genes regulate distinct immune processes, driven by luminal antigens, expressed as specific immune phenotypes which in turn influence clinical phenotypes in IBD patients. Immune responses to various microbial antigens among patients with CD may be related to different pathophysiological mechanisms of disease and a genetically susceptible host may yield distinct disease characteristics as a result of the immune responsiveness expressed by the at risk individual. Further research is necessary to validate the associations of immune responses to disease behavior and prognosis and for novel
immune responses to be identified so to provide more information on the underlying etiopathogenic mechanisms of characteristic of IBD. Advances in this field will afford clinicians the opportunity to create and implement appropriate and timely therapeutic management regimes based on the aggressiveness of the IBD subtype in order to alter and thus improve the long-term prognosis.

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