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**Role of *TNFSF15* in the intestinal inflammatory response**

Kadiyska T *et al*. *TNFSF15* and intestinal inflammation

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**Abstract**

Gastrointestinal diseases, specifically Crohn’s disease (CD), ulcerative colitis (UC), diverticular disease (DD) and primary biliary cirrhosis (PBC) are all characterized by complicated inflammation of the digestive tract. Their pathology is multifactorial, and risk factors encompass both genetic and environmental factors. Recent advances in the genetic component of inflammatory bowel diseases (IBDs) have revealed that the tumor necrosis factor superfamily member 15 (*TNFSF15*) contains a number of risk alleles, associated not only with IBD but also with other diseases such as DD and PBC. These risk alleles in *TNFSF15* and the altered expression of its gene product can serve as the common ground between these disorders and bridge the gap between them by explaining if not all, then at least some of the underlying processes that lead to a dysregulated immune response and subsequent chronic inflammation. Here, we aim to outline how the *TNFSF15* gene is involved in the proliferation and cell fate of different populations of T cells and subsequently, in the control of both pro- and anti-inflammatory cytokines. Furthermore, we summarize what is currently known of *TNFSF15* control region variants, how they are associated with each mentioned disease and how these variants can explain the autoimmune pathology of said diseases through altered *TNFSF15* expression.

**Key words:** Tumor necrosis factor superfamily member 15; Death receptor 3; Crohn’s disease; Ulcerative colitis; Diverticular disease; Primary biliary cirrhosis

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**Core tip:** Tumor necrosis factor superfamily member 15 (*TNFSF15*) and the protein it encodes, tumor necrosis factor ligand-related molecule 1 (TL1A, full transcript), play a vital role in the mucosal immunity. Expression of TL1A, and death receptor 3 (DR-3) mediated signaling, both exert their effects in Crohn’s disease (CD), ulcerative colitis (UC), diverticular disease (DD) and primary biliary cirrhosis (PBC), which can serve to bridge the gap of knowledge regarding the genetic components of this group of inflammatory diseases as well as provide common ground for a putative targeted treatment.

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**INTRODUCTION**

Inflammatory bowel disease (IBD) is a chronic disorder that constitute an important worldwide health problem. This group of diseases is multifactorial and characterized by chronic relapsing intestinal inflammation[1]. The two major subtypes of IBD are ulcerative colitis (UC) and Crohn’s disease (CD). Nowadays, IBD is a global disease with the highest prevalence in Western countries (North America, Europe and Oceania) although recently there has been an accelerated incidence rate in the newly industrialized countries of Asia, South America and Africa, where societies have become more westernized[1]. Although the exact etiology of IBDs is still unknown, numerous studies have attempted to shed light on the subject, and have revealed that the multifactorial nature of IBDs encompasses genetic susceptibility, environmental factors, intestinal microbiota and the immune response system as factors involved in the pathogenesis of IBDs[2].

 Another common gastrointestinal disorder, similar in its prevalence amongst western populations is colonic diverticulosis. The term “diverticulosis” refers to the occurrence of diverticula due to the formation of pouches by the mucosal wall of the intestine[3]. Colonic diverticulosis, or diverticular disease (DD), is a broad-spectrum term, as the condition involves a number of clinical manifestations that can range from the presence of constant abdominal systems without inflammation [symptomatic uncomplicated diverticular disease (SUDD)] to a significant and symptomatic inflammatory process (segmental colitis associated with diverticulosis and diverticulitis)[4].

 Both diverticulitis and IBDs share overlapping characteristics and symptoms including, but not limited to: clinical presentation involving diarrhea, mucus in the stool, abdominal pain, weight loss, fistulae, bowel structuring and inflammation[5,6]. This overlap can make diagnosis difficult for the attending clinician, although distinction can be achieved by endoscopical examination[5]. Despite this difficulty and in order to improve our understanding of the relation between inflammation and gastrointestinal disorders, we have to ask the question, what is the driving factor behind these shared attributes of DD, UC and CD?

 The common ground for the pathological signs of IBDs and DD appears to be a dysregulated mucosal immune response[7,8]. This dysregulation often results in impaired epithelial barrier function and damage to the surrounding epithelial tissue. Both pro- and anti- inflammatory cell lines and their respective secreted cytokines are involved in this response. In CD, T helper 1 (Th1)/Th17 cells and interleukin (IL)-12 as well as IL-23 are characteristic, whereas in UC the major factor is natural killer T (NKT) cells secreting IL-13 and IL-5[9].

 Tumor necrosis factor superfamily member 15 (*TNFSF15*), also known as tumor necrosis factor ligand-related molecule 1 (TL1A) and vascular endothelial growth inhibitor (VEGI) is a tumor necrosis factor (TNF) family member, is a gene, encoding a ligand produced by a variety of cell lines, including endothelial cells, macrophages, dendritic cells (DCs) and T cells[10]. First described in 2002 as a T-cell stimulatory cytokine[11], studies have discovered that it affects cell lines related to both the innate and adaptive immune responses by its receptor, death receptor 3 (DR3)[12]. Since then, the role of this cytokine-receptor pair has been linked to the immunomodulation and vascular endothelial function observed in IBDs[6].

***TNFSF15* FUNCTION AND EXPRESSION**

The gene product of *TNFSF15*, TL1A, is a TNF-like factor, which is expressed in endothelial cells (human umbilical vein endothelial cells, adult dermal microvascular endothelial cells and uterus myometrial endothelial cells), gut lamina propria lymphocytes and macrophages.

 TL1A is a longer splicing variant of the coding gene *TNFSF15* compared to the firstly described protein TL1/VEGI. The difference between the two variants is that TL1A is encoded by all four coding exons, whereas TL1 is encoded by a continuous DNA containing the forth exon and its 5’ adjacent intron. As a result, the two variants have identical C-terminal regions, while the N-terminal regions are different for the two proteins. TL1A is a type II transmembrane protein, containing 251 amino acids and has a molecular weight of 28 kDa. The transmembrane form of TL1A can be cleaved by enzymes and exists as a functional soluble protein[11,13]. This cleavage can vary depending on the cell of origin[14], with the soluble form being more abundantly synthesized by DCs; the membrane-bound protein was found to be expressed by both T cells and DCs. The different forms, like other members of the TNF superfamily, have different functions; soluble TL1A can be detected after DC and monocyte stimulation *in vitro*, and increased levels have been detected in serum samples from patients with rheumatoid arthritis, an autoreactive disease[15].

 The receptor for TL1A, DR3, was identified in the 1990s[16], and was later discovered to be highly homologous to TNF receptor 1 (TNFR1)[12]. Signaling by DR3 is facilitated primarily through the use of TNFR-associated death domain protein (TRADD), which contains a TNF receptor associated factors (TRAF)-binding domain as well as a death domain. This combination allows DR3 to activate nuclear factor kappa B (NF-kB) and mitogen-activated protein kinase (MAPK)[17], which allows it to play a role in both apoptosis and anti-apoptosis, cell survival and proliferation[18].

 The expression of TL1A is closely linked to the levels of inflammation over the course of IBD and is also correlated to areas affected by the disease[10]. While TL1A baseline expression can be low[19], proinflammatory stimulation seems to be the switch that increases TL1A expression[20]. Both TL1A and DR3 are expressed across all members of the T cell family[21,22], despite the original discovery of TL1A in endothelial cells. The action of TL1A-DR3 signaling is most profound in the differentiation and stimulation of T cell subtypes. Co-stimulation with TL1A increases IL-2 signaling[11,20,23], whereas TL1A itself stimulates the proliferation of T cells. Specific CD4+ T cells can up-regulate DR3 and produce interferon gamma (IFN-γ) in response to TL1A combined with IL-12 and IL-18 can be found in the intestinal mucosa[23], suggesting a putative mechanism for TL1A gut signaling and expression[19]. TL1A also affects Th17 cells, as DR3 expression is highly upregulated on this cell subset[24], although TL1A-mediated Th17 proliferation is achieved in a DR-3 independent manner[20]. Furthermore, TL1A plays a role in the development of regulatory T cells (Treg), as stimulation by the soluble form of TL1A increases Treg proliferation[25]. However, *in vitro* assays have shown that the increased numbers of Tregs also show reduced suppressive capability[26].

***TNFSF15* AND INFLAMMATION**

Genetic studies attempting to evaluate the role of *TNFSF15* have only begun recently, following previously suggested hints of genetic factors involved in IBD[27,28]. The first genome-wide association studies (GWAS) conducted in 2005 discovered an association between TL1A and CD in a Japanese cohort of patients[29]. Subsequent studies have replicated and confirmed the association of TNFSF15 in European populations, both for patients with CD and UC[30]. Further investigation on specific patient subsets confirmed the protective haplotype[31] and revealed that TL1A expression is increased in carriers of the risk haplotype in a Jewish cohort of patients with CD and *Escherichia coli* (*E.coli*)exposure[32].

 The findings of the previously mentioned studies and the data obtained allowed for the further investigation of *TNFSF15* single nucleotide polymorphisms (SNPs) and their role not only in UC and CD, but also in DD. The first case study aiming to investigate how these SNPs can exert an effect revealed that the SNP rs7848647 and specifically, the risk allele G conferred a higher risk, in additive mode, towards DD requiring surgical intervention[33]. As a follow-up, a further study aimed to increase the number of participants and to include six other SNPs, four of which had been previously associated with CD[29] as a risk haplotype, and to reveal if such an association could be found for DD as well. Results demonstrated not only that the CD risk haplotype was associated with DD, but two protective haplotypes emerged as well[6]. Although both studies provided hopeful results, they also suggest that there might be further undiscovered SNPs in DD pathology.

 *TNFSF15* variants have also been associated with primary biliary cirrhosis (PBC), a chronic and progressive liver disease, leading to hepatic failure and liver cirrhosis. One of the hallmarks of PBC is an autoimmune reaction towards biliary epithelial cells. Combined with data from twin studies, this has driven research into a possible genetic component of PBC. The first GWAS studies demonstrated the association of *TNFSF15* rs4979462 with PBC in Asian populations[34] as the second strongest susceptibility gene. Specifically, rs4979462 has been found to be one of the main causal variants in the gene, due to a creation of a novel nuclear factor 1 (NF-1) binding site, a finding further strengthened by the increased *TNFSF15* mRNA expression[35]. Other large-scale GWAS studies have also demonstrated that *TNFSF15* is a part of a multitude of PBC risk loci involved in T cell, B cell and natural killer (NK) cell stimulation and proliferation[36].

 TL1A expression has also proven to be the one of the key hallmarks of IBD. It was first observed in UC and CD[10,21,37] with both protein and mRNA levels increased compared to healthy controls[19]. This expression is regulated in a two-fold manner in gastrointestinal disorders: first, single-nucleotide polymorphisms have been found to correlate with TL1A expression[32,38,39] in a variety of immune cells. Second, TL1A can be induced by a number of gut-specific bacteria[22], with expression levels adjusting accordingly to the presence or absence of bacteria. Localized increased inflammation correlating with increased TL1A expression and exposure to bacteria in patients with CD, DD and UC appears to be the common ground between these gastrointestinal disorders (Figure 1).

 Further proof for the involvement of *TNFSF15* comes in the form of studies, investigating the association of risk variants within the gene and the requirement of surgical intervention as part of the treatment plan[33,40]. Cases that do not respond to medical treatment and present themselves with severe colonic inflammation ultimately require surgical resection, which represents a higher risk for the patient.

 Because of its mode of action, and by having a very specific niche of activity and expression, *TNFSF15* can be considered as a putative therapeutic target. Studies have investigated the effect of anti-TL1A antibodies on sodium-sulfate induced colitis[41] and T-cell transfer models[42]; some have even successfully managed to reverse fibrosis in these models[43]. As TL1A expression can vary, depending on genetic variations in its control region, it remains to be seen whether a targeted anti-TL1A therapy can be applied in a clinical setting.

**CONCLUSION**

*TNFSF15* and the protein it encodes share a unique position amongst other genetic factors for gastrointestinal disorders. It enhances the T cell responses in CD, UC and DD acting as a bridge between these disorders. It also plays a role in the immune response of mucosal tissue against bacterial infections, another common factor in gastrointestinal disorders. Furthermore, genetic variations in this gene have been associated as risk factors for all of the diseases examined here. All of these characteristics point towards the further study of *TNFSF15* and the TL1A-DR3 interaction among patients with IBD and DD both as a putative therapeutic target and a risk prediction factor.

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**Figure 1 Effect of tumor necrosis factor ligand-related molecule 1 signaling across different subtypes of leukocytes.** Tumor necrosis factor ligand-related molecule 1 (TL1A) acts by promoting cytokine signaling, cell differentiation and proliferation. These effects vary between different subtypes. While TL1A promotes natural killer T cells (NKT) and T helper Th1/17 proliferation, it can also suppress regulatory T cell (Treg) functions and forkhead box P3 (Foxp3)+ expression. Regardless, TL1A is not associated exclusively with any form. CD: Crohn’s disease; DC: Dendritic cell; DD: Diverticular disease; IFN-γ: Interferon gamma; IL: Interleukin; NKT: Natural killer T cells; TL1A: Tumor necrosis factor Ligand-related molecule 1; Th: T helper; Treg: Regulatory T cell; Foxp3: Forkhead box P3; UC: Ulcerative colitis.