Tripartite motif family proteins in inflammatory bowel disease: Mechanisms and potential for interventions

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Abstract
Inflammatory bowel disease (IBD) is a chronic recurrent gastrointestinal inflammatory disease that poses a heavy burden to the global healthcare system. However, the current paucity of mechanistic understanding of IBD pathogenesis hampers the development of aetiology-directed therapies. Novel therapeutic options based on IBD pathogenesis are urgently needed for attaining better long-term prognosis for IBD patients. The tripartite motif (TRIM) family is a large protein family including more than 70 structurally conservative members, typically characterized by their RBCC structure, which primarily function as E3 ubiquitin ligases in post-translational modification. They have emerged as regulators of a broad range of cellular mechanisms, including proliferation, differentiation, transcription and immune regulation. TRIM family proteins are involved in multiple diseases, such as viral infection, cancer and autoimmune disorders, including inflammatory bowel disease. This review provides a comprehensive perspective on TRIM proteins' involvement in the pathophysiology and progression of IBD, particular, on intestinal mucosal barriers, gene susceptibility and opportunistic infections, thus providing novel therapeutic targets for this complicated disease. However, the exact mechanisms of TRIM proteins in IBD pathogenesis and IBD-related carcinogenesis are still unknown, and more studies are warranted to explore potential therapeutic targets of TRIM proteins in IBD.

1 | INTRODUCTION

Inflammatory bowel disease (IBD) is a chronic relapsing inflammatory disease of the gastrointestinal tract that can be subdivided into Crohn's disease (CD) and ulcerative colitis (UC).¹ CD is distinguished by transmural inflammation and skip lesions distributed throughout the gastrointestinal tract, usually complicated by perianal lesions. UC associated inflammation is generally confined to rectum and colon and afflicts only the superficial mucosa, which usually demonstrates mucopurulent bloody stool.²,³ The prevalence of IBD is as high as 0.5% in western countries, which has brought huge expenditure and heavy burdens to the healthcare system.⁴ In 2020, the annual direct expenditure of every UC patient in Europe was estimated as €2000, and that of CD was €3500.⁵ Moreover, in newly industrialized countries with lower IBD prevalence, such as China and India, the large populations and rising incidence rates bring remarkably heavy burdens to society.⁶ At present, the mainstream therapy for IBD focuses on immunosuppression, in which biological agents have significantly improved prognosis.⁷ However, despite these advances, over 40% IBD patients still require at least one surgery during the course of the disease; moreover,
The tripartite motif (TRIM) protein family contains more than 70 members characterized by the RBCC structure, in which one or two variable C-terminal domains show high structural diversity that determines the functional specificity of each protein. TRIM proteins mainly function as E3 ubiquitin ligases, thus participating in the ubiquitination process and regulating many important post-translational protein modifications. TRIM family proteins not only play essential roles in many biological processes, including proliferation, differentiation, transcription and apoptosis, along with the regulation of immune responses, but also participate in many diseases, including cancer, infectious diseases, neurodegeneration and developmental diseases. Recently, TRIM family proteins, such as TRIM20 and TRIM27, were demonstrated to regulate intestinal barrier function and get involved in the pathophysiology of IBD. This review summarizes the structures and functions of TRIM proteins and discusses the mechanisms underlying their role in IBD pathophysiology to provide a novel approach for the exploration of potential therapeutic targets for IBD.

2 | TRIM FAMILY

2.1 | Structure of TRIM family proteins

In general, TRIM family members share a highly conservative and typical structure of “RBCC”, which means that the order of TRIM proteins, from N- to C-terminus, is a really interesting new gene (RING) domain, one or two B-box domains and a coiled-coil (CC) domain. The RBCC domain is usually followed by more divergent C-terminal domains, which determine the specificity of each TRIM protein. Moreover, the subcellular distributions of TRIM proteins show high variability, with their presence being reported in the cytoplasm, nucleus and plasma membrane.

The RBCC motif, which generally consists of three specific domains, is present in almost all members of the TRIM family. The RING domain is the foremost structure consisting of 40–60 amino acid residues with the zinc-finger structure that binds two zinc ions. Some investigations have found that they specifically bind to E2 ubiquitin-conjugating enzymes to exert E3 ubiquitin ligase activity, thus mediating the conjugation of specific proteins with one of the most widely recognized post-translational modifiers, ubiquitin. The B-box domain consists of 32–42 residues located behind the RING domain, which can combine with one or two zinc atoms as well. Depending on the number of residues, it can be divided into B-Box 1 and B-Box 2. Nevertheless, not every TRIM protein contains both simultaneously; some, such as TRIM36, TRIM46 and TRIM54, only have B-Box 2 domains. B-Box 1 invariably locates ahead of B-Box 2 when both exist. However, the exact functions of B-box domains remain unclear. Since B-box shares similar structural characteristics with the RING domain, it is speculated that B-box may function as E3 ubiquitin ligase, similar to the RING domain. Several previous studies have attempted to support this hypothesis. For example, TRIM16 and TRIM29 still function as E3 ubiquitin ligases even though they lack the RING domain, and further analysis suggested that the B-box domain shares similar folding pattern and zinc-binding characteristics with the RING domain, which are important for E3 ligase activity. Moreover, some researchers hypothesize that B-box enhance the recognition of target proteins. The coiled-coil (CC) domain is the third characteristic structure of TRIM proteins, which is known to mediate the homologous or heterogeneous oligomerization of TRIM proteins and has a specific subcellular localization function.

Unlike the highly conserved RBCC sequences, the C-terminal domain behind the RBCC domain is highly variable; they have been implicated in substrate recognition and serve as a binding site for different targets. So far, a total of ten types of C-terminal domains have been identified by structural analysis, and different combinations of them allow TRIM proteins to be categorized into 11 distinct subclasses (C-I to C-XI in Table 1), which may contain no or as many as three C-terminal domains. In addition, a specific set of TRIM proteins which lack the RING domain are classified as the uncategorized group (Table 1). The most prevalent C-terminal domain supposes to be SPRY, which exists in more than half of the TRIM family members and sometimes coexists with the PRY domain. The SPRY domain mediates the recognition of and interaction with target proteins or RNA and regulates host-pathogen interactions and innate immune responses. The C-terminal subgroup one signature (COS) domain is another frequently encountered C-terminal domain, which can bind to the microtubule cytoskeleton and participate in homodimerization or heterodimerization. The fibronectin type 3 (FN3) domain interacts with DNA and heparin, and plant homeodomain (PHD) mediates binding to histones to regulate transcription. The meprin and tumour necrosis factor (TNF) receptor-associated factor homology (MATH) domain interacts with TNF receptors and regulates the function of important transcription factors, such as NF-κB. Less common C-terminal structures include ADP-ribosylation factor-like (ARF), filamin-type immunoglobulin (FIL), transmembrane (TM), and NHL (named after three proteins, with “H” representing HT2A [TRIM32]) domains. In summary, the high variability of C-terminal domains determines TRIM's specificity and functional diversity by binding to different substrates, thus exerting diverse regulatory effects.

2.2 | Functions of TRIM family proteins

TRIM proteins share E3 ubiquitin ligase activity; therefore, they participate in the ubiquitin–proteasome pathway, which is an important post-translational modification process for regulating the homeostasis and degradation of proteins.
Ubiquitin, a highly conserved protein of 76 residues, is expressed in all eukaryotic cells.\(^\text{14}\) Ubiquitination is an ATP-dependent process in which the C-terminal glycine of ubiquitin is covalently bound to a lysine residue of a target protein via the sequential catalysis by an E1 ubiquitin-activating enzyme, E2 ubiquitin-conjugating enzyme and E3 ubiquitin ligase.\(^\text{42}\) Afterwards, the ubiquitinated targets are subjected to proteasome degradation, wherein the fate of the substrate is determined by the amount of modified lysine and conjugated ubiquitin.\(^\text{43}\) Depending on the functions of target proteins, ubiquitination participates in many biological processes, such as autophagy, innate immune signalling pathways (e.g., JAK-STAT3 pathway and TLR-mediated pathway) and carcinogenesis.\(^\text{14}\) Most TRIMs function as E3 ubiquitin ligases to regulate these biological processes, which are dependent on the specific substrates recognized by them. Moreover, some TRIMs regulate these processes by binding to ubiquitin-like proteins including SUMO\(^\text{44}\) and dNEDD8.\(^\text{45}\) The nine unclassified TRIM proteins lacking the RING domain may indirectly promote ubiquitination or act in conjunction with other TRIM proteins.\(^\text{15}\) As a family characterized by specific structural features rather than functional purposes, TRIM proteins are described as modulators of multiple cellular and physiological activities related to many diseases (e.g., cancer, viral infection and autoimmune disorders) by regulating the activity, stability, degradation, distribution and interaction process of some key proteins.\(^\text{31,46}\)

**TABLE 1** Classification and structure of TRIM proteins in humans

| Family | RBCC structure | Members | Functions |
|--------|----------------|---------|-----------|
| C-I    | N R B1 B2 CC COS FN3 PRY SPY | TRIM1, TRIM9, TRIM18, TRIM36, TRIM46, TRIM57, TRIM63 | Interaction with microtubule cytoskeleton\(^\text{29}\) |
| C-II   | N R B1 B2 CC COS | TRIM54, TRIM55, TRIM63 | Muscle protein turnover\(^\text{30}\) |
| C-III  | N R B1 B2 CC COS FN3 | TRIM42 | Interaction with DNA and heparin\(^\text{17}\) |
| C-IV   | N R B1 B2 CC PRY SPY | TRIM4, TRIM5, TRIM6, TRIM7, TRIM10, TRIM11, TRIM15, TRIM17, TRIM21, TRIM22, TRIM25, TRIM26, TRIM27, TRIM34, TRIM35, TRIM38, TRIM39, TRIM41, TRIM43, TRIM47, TRIM48, TRIM49, TRIM50, TRIM51, TRIM53, TRIM58, TRIM60, TRIM62, TRIM64, TRIM65, TRIM68, TRIM69, TRIM72, TRIM75, TRIM77, TRIM71 | Interaction with various proteins or RNA\(^\text{31}\) |
| C-V    | N R B1 B2 CC | TRIM8, TRIM19, TRIM31, TRIM40, TRIM56, TRIM61, TRIM73, TRIM74 | Transcriptional regulation\(^\text{19}\) |
| C-VI   | N R B1 B2 CC PHD BR | TRIM24, TRIM28, TRIM33 | Interaction with histones and regulation of gene transcription\(^\text{32}\) |
| C-VII  | N R B1 B2 CC FIL NHL | TRIM2, TRIM3, TRIM32, TRIM71 | Transcriptional and post-transcriptional regulation of RNA\(^\text{33}\) |
| C-VIII | N R B2 CC MATH | TRIM37 | Interaction with TNF receptors and regulation of transcription factors such as NF-kB.\(^\text{34}\) |
| C-IX   | N R B1 B2 CC ARF | TRIM23 | GTPase function\(^\text{31}\) |
| C-X    | N R B1 B2 CC FIL | TRIM45 | Repressor of multiple signalling pathway\(^\text{35}\) |
| C-XI   | N R B1 B2 CC TM | TRIM13, TRIM59 | Autophagy modulation\(^\text{36}\) |
| Uncategorized | N R B1 B2 CC | TRIM14, TRIM16, TRIM20, TRIM29, TRIM44, TRIM51, TRIM66, TRIM70, TRIM76 | Indirect role in assisting ubiquitination\(^\text{15}\) |

Note: The TRIM proteins associated with IBD or IBD-related opportunistic infections are shown in bold.

Abbreviations: ARF, ADP-ribosylation factor-like domain; B1, B-Box 1; B2, B-Box 2; BR, bromodomain; C, C-terminal; CC, Coiled-coil Domain; COS, C-terminal subgroup one signature domain; FIL, filamin-type immunoglobulin domain; FN3, fibronectin type 3 domain; MATH, meprin, and TNF receptor-associated factor homology domain; N, N-terminal; PHD plant homeodomain; R, RING Domain; TM, transmembrane domain.
| TRIM | Referred barrier functions | Expression in inflammation tissue | Tested cell organizations | Experimental model | Function | Reference |
|------|---------------------------|----------------------------------|--------------------------|-------------------|----------|-----------|
| 11   | Mechanical barrier        | —                                | TRIM11<sup>+/−</sup> HT-29 cells and wild type cells; Mice intestinal epithelial cell; active CD patients’ colonic mucosa | DSS-induced colitis mice | TRIM11 participates in TSC1/mTOR’s role in restraining epithelial necroptosis and intestinal inflammation by mediating the ubiquitination of RIPK3 | 48        |
| 19   | Chemical barrier          | —                                | HeLa cells and HEK293 cells | —                 | TRIM 19 stimulates human beta-defensin 2 secretion by regulating myeloid elf-1-like factor transcription | 49        |
| 20   | Mechanical barrier        | Overexpression                   | Colon tissues of MEFV<sup>+/−</sup> and wild type mice | DSS-induced colitis mice | MEFV regulates epithelial junctional integrity (occluding and claudin-2), cytokines level (IL-6, IL-18) as well as cell markers expression (STAT3) | 50        |
|      | Immune barrier            | Overexpression                   | Colon tissue of MefV<sup>V726A/V726A</sup>, MefV<sup>V726A/V726A</sup>, TNF<sup>−/−</sup>, TNFR1<sup>−/−</sup>, TNFR2<sup>−/−</sup> and wild type mice | MefV<sup>V726A/V726A</sup> FMF mice model | MefV<sup>V726A/V726A</sup> mutation induces substantial colitis with immune cell infiltration in the intestine through regulating the TNF/TNFR axis and TRIM20 (pyrin) activation process | 51        |
|      |                           | Mutation                         | IBD patient DNA FMF patient DNA | IBD patients | The prevalence of FMF in IBD patients is higher than normal people, and prevalence of IBD in FMF patients is higher as well | 52–57     |
| 21   | Mechanical barrier; Immune barrier | Downexpression                   | Intestinal tissue from TRIM21<sup>+/−</sup> and wild-type mice; Colon tissue from UC-associated cancer patient | Azoxymethane-<sup>-</sup>DSS mice model | TRIM21 can participate in modulating gene expression of Ki67, E-cadherin, β-catenin+ cells, matrix metalloproteinase 10, cyclooxygenase 2, hypoxia-inducible factor-1α, angiogenin 4, IL-1β, IL-6, TNF-, TGF-β, Foxp3, IL-10, IFN-γ | 58        |
|      | Immune barrier            | Downexpression                   | TRIM21<sup>+/−</sup> and wild-type mice. Inflamed mucosa of patients with IBD | Trinitrobenzene sulfonic induced colitis mice model; CD45RB<sup>hi</sup> CD<sup>+</sup> T cell-induced colitis mice model | TRIM21 restricts intestinal inflammation by inhibiting the differentiation of Th1 and Th17, and its downstream includes interferon regulatory factor 3 | 59        |
| 22   | Immune barrier            | Mutation                         | Paediatric-onset IBD patient gene; | —                 | Nearly 1% of patients with paediatric-onset IBD were diagnosed with TRIM22-related disease | 60        |
| TRIM | Referred barrier functions | Expression in inflammation tissue | Tested cell organizations | Experimental model | Function | Reference |
|------|-----------------------------|-----------------------------------|---------------------------|-------------------|----------|-----------|
| Immune barrier | Mutation | HEK 293 cells; Inflamed and non-inflamed intestinal and rectal tissues from patients with very early onset IBD; | -- | -- | TRIM22 can interact with NOD2 and regulate NOD2-dependent signalling pathways for various anti-pathogenic processes (muramyl dipeptide or Respiratory Syncytial Virus) | 61 |
| 27 Immune barrier | -- | Trim27<sup>−/−</sup> and wild type mice; | DSS-induced colitis mice | TRIM27 positively regulates activation of STAT3 by promoting JAK1–STAT3 complex formation after IL-6 stimulation | 62 |
| Immune barrier | Overexpression | HEK293T cells, SW480 cells; CD patients’ sigmoid colon tissue | -- | -- | TRIM27 restrains NOD2-mediated inflammatory responses by ubiquitinating and degrading NOD2 | 63 |
| 28 Immune barrier | -- | Bone marrow and colon tissues of TRIM28<sup>−/−</sup> and wild-type mice | T cell transfer colitis model | TRIM28 could modulate the epigenetic silencing of Treg-characteristic genes, thus regulating helper and regulatory T cell differentiation and activation | 64 |
| 29 Mechanical barrier | -- | GHR<sup>−/−</sup> and wild type mice | Xenograft model | Growth hormone could inhibit DNA repair of epithelial cells by inducing TRIM29 | 65 |
| 30α Immune barrier | -- | LTβ<sup>−/−</sup>, LTβ<sup>−/−</sup> and wild type mice | DSS-induced colitis mice | Lymphotoxin αβ (LTβR) derived from CD<sup>+</sup> T cells binds to lymphotoxin-β receptor (LTβR) of macrophages to restrict inflammation through TRIM30α-dependent signal pathway | 66 |
| 31 Biological barrier | Downexpression | Intestinal tissue of CD patients and controls | -- | -- | TRIM31 can enhance the autophagy process in intestinal cells, and this process has no dependencies on Atg5 or Atg7 | 28 |
| Immune barrier | -- | TRIM31<sup>−/−</sup> and wild-type mice | DDS-induced colitis mice | TRIM31 could ubiquitinate and degenerate NLRP3 inflammasome in macrophages, and TRIM31 deficiency attenuates the colitis severity in symptoms, colonic length, and intestinal histology | 67 |
| 33 Immune barrier | Downexpression | Trm33<sup>−/−</sup> and wild-type mice | DDS-induced colitis mice | TRIM33 exerts essential roles in intestinal | 68 |

(Continues)
Intestinal barrier dysfunction is an important hallmark of IBD. TRIM family proteins regulate the function of the four parts of intestinal barrier (mechanical, biological, chemical and immune). These functions are summarized in Table 2 and Figure 1.

### 3.1 TRIM and intestinal barriers

#### 3.1.1 Mechanical barrier

The mechanical barrier, also known as the physical barrier, is an essential defensive line of the intestinal barrier. Its purpose is to separate...
the luminal contents from the intestinal immune system, whose structural basis is the mucus layer, intact epithelial cells and the junctional complexes between epithelial cells.\textsuperscript{73} Under physiological conditions, the epithelial surface of the intestinal mucosa is covered with mucus, which contributes to both the mechanical and chemical barrier function. The intestinal monolayer epithelium, together with the intercellular junctional complexes (e.g., tight junctions), exerts a significant impact in regulating intestinal permeability and preventing harmful substances (such as bacteria, toxins or other inflammatory mediators) migrating from the intestinal cavity to the epithelium, whose impairments have been reported in the pathogenesis of IBD.\textsuperscript{74,75} TRIM20, encoded by the \textit{MEFV} gene, is widely considered to be associated with the development of familial Mediterranean fever (FMF).\textsuperscript{76} However, it is frequently reported that FMF patients have a significantly higher incidence of IBD and the incidence of FMF in IBD is also the case in which \textit{MEFV} mutations are often detected (e.g., M694V, M680I and V726A mutation).\textsuperscript{52,53} Very little is known concerning the underlying mechanism for that phenomenon, although some information has surfaced in the past few years. Sharma et al. found that TRIM20 regulates tight junction (occludin and claudin-2) integrity, thereby modulating the permeability of the intestinal epithelium in a murine model of dextran sulphate sodium (DSS)-induced colitis.\textsuperscript{50} Moreover, they found that experimental colitis in homozygous \textit{MEFV} \textit{V726A/V726A} mutation mice presented much severe colitis, marked by immunocyte infiltration in the intestinal tract, contributing to the immune barrier.\textsuperscript{51} Apart from tight junctions, several TRIM proteins regulate the epithelial integrity to modulate the mechanical barrier function. For example, TRIM11 was reported to be an important modulator of RIPK3-dependent epithelial necroptosis, whose decline could be observed in the colonic mucosa of CD patients.\textsuperscript{48} In addition, some studies have suggested a protective role of TRIM21 and TRIM58 in IBD pathogenesis via modulating epithelial regeneration, tissue repair and angiogenesis.\textsuperscript{58,70} In addition, Chesnokova et al. observed that growth hormone inhibits epithelial DNA repair by inducing TRIM29.\textsuperscript{65}

3.1.2 | Biological barrier

The biological barrier is composed of trillions of microorganisms that colonize the intestine. These microorganisms are referred to as the intestinal microflora and could assist in nutrition absorption, host defence and the function of the immune system.\textsuperscript{77} Changes in the
composition of intestinal microflora and alterations of interactions between intestinal microflora and intestinal immunity have notable impacts on IBD. A recent study reported decreased TRIM31 expression in the intestinal tracts of CD patients. Subsequent experiments revealed that TRIM31 usually accumulates around one type of ubiquitin-coated invasive bacteria, such as Shigella or Salmonella. This is attributed to the specific interaction between TRIM31 and the bacterial receptor NDP52, which induces protective autophagy against invasive bacterial infection or other normal commensal bacteria. Therefore, it was inferred that TRIM31 plays essential roles in restricting invasive bacterial infection in intestinal epithelial cells by promoting autophagy, which can affect gut microbes and lead to IBD in pathological cases.

3.1.3 | Chemical barrier

The impairment of intestinal chemical barrier, primarily mucus and antimicrobial peptides, can contribute to IBD. The mucus produced by goblet cells, is rich in mucin glycoprotein, and Mucin-2 is one of its most abundant components. Antimicrobial peptides from Paneth cells could promote the permeability of target bacterial membranes and induce bacterial killing. The chemical barrier is essential for the segregation function and antimicrobial activities in the intestine to implicate the pathogenesis of IBD.

A recent study revealed TRIM34’s crucial role in the release of Mucin 2 from goblet cells, to the integrity of the inner mucus layer and attenuates colitis induced by DSS. TRIM19 can upregulate the expression of endogenous human β-defensin 2, which is an important antimicrobial peptide involved in IBD pathogenesis and has been reported to exert protective effects against a variety of pathogens, such as Escherichia coli and Candida albicans.

3.1.4 | Immune barrier: adaptive/innate immune

The intestinal microflora, mucosal barrier, and immune system work together to maintain intestinal homeostasis in healthy individuals. However, this balance is disrupted in IBD patients, characterized by dysregulated intestinal flora, destruction of the mucosal barrier, and overactivation of the immune system. Studies have shown that both innate and adaptive immunity are engaged in the pathophysiology of IBD. The former is likely to have an effect on IBD through cytokines and complement production, microbial recognition and autophagy, whereas the latter involves the processes of humoral and cellular immunity. In the following section, we review the association between the immune barrier and TRIM proteins in the pathogenesis of IBD.

3.1.5 | Innate immunity

Innate immunity serves as the first line of defence against pathogens and acts by mediating intestinal microorganism recognition and quickly inducing inflammatory responses. NOD2, CARD9 and TLR2 are some of the crucial molecules involved in innate microbial sensing. The NOD2 gene, located in the genomic region imparting IBD susceptibility as determined by genome-wide association studies, encodes NOD2, an intracellular pattern recognition receptor that targets peptidoglycans present in gram-positive or gram-negative bacteria, and subsequently activates the NF-κB signalling pathway. Both TRIM22 and TRIM27 have been identified as regulators of NOD2 and contribute to intestinal inflammation in IBD.

CARD9 is another important component of innate microbial sensing pathways identified by exome sequencing for its involvement in IBD, which also lead to the activation of the NF-κB signalling pathway and cytokine production, especially in fungal infection. Cao et al. found that TRIM62 only ubiquitinates intact CARD9, while the C-terminal truncation of CARD9 disrupted this interaction. As a result, C-terminal truncated CARD9 variant exhibits protective roles in intestinal inflammation through inhibiting cytokine production and immune response. This study also demonstrated that TRIM62-mediated ubiquitinoylation can regulate CARD9-dependent NF-κB signalling pathways to alter intestinal immunity in murine models of Candida albicans infection and DSS-induced colitis. Subsequently, this research team discovered a small molecule targeting for disrupting the interaction between CARD9 and TRIM62, which imitates the protective role of CARD9 variants, to illustrate a novel potential therapy for IBD treatment. Toll-like receptor 2 (TLR2) is another pattern recognition receptor, and it has been reported that TRIM58 downregulates innate immune response in myeloid cells via TLR2 signalling pathway, such that severe mucosal injury was observed in TRIM58 knockout mice during DDS induction. Furthermore, TRIM40 serves as a negative modulator of inflammation that inhibits NF-κB activity in the gastrointestinal tract.

In addition to microbial recognition, TRIM proteins also participate in IBD pathogenesis by regulating innate immune cells, such as macrophages and myeloid cells. TRIM33 is essential for monocyte recruitment and differentiation along with macrophage M1/M2 switch and membrane-bound TNF expression, whose impairment may result in an increase in monocyte and decrease in macrophage counts in the colon accompanied by upregulated colonic inflammation. In DSS-induced murine models, TRIM31 was observed to promote the ubiquitin-proteasome pathway of the NLRP3 inflammasome in macrophages, whose defects have been implicated in IBD pathogenesis.

3.1.6 | Adaptive immunity

TRIM family proteins participate in the cellular immunity process of CD+ T lymphocyte differentiation into helper cell types (Th1, Th2 and Th17) and regulatory T (Treg) cells. TRIM21 was observed to have protective roles in IBD by downregulating some pro-inflammatory cytokines (e.g., IL-6 and TNF-α) and inhibiting CD4+ T cell differentiation into Th1 and Th17 cells. The IL-6/JAK/STAT3 signalling pathway has been reported to regulate the differentiation of multiple immune cell types, such as Th2 and Th17 cells. TRIM27 promotes the interaction between STAT3 and JAK1 in this pathway to trigger STAT3 activation in the serum or colonic mucosa, thereby inducing inflammation.
TRIM27 knockout mice were observed to gain stronger resistance to DSS-induced colitis, and histopathological examination revealed that their colonic mucosa was more intact, with less disruption.\textsuperscript{62} TRIM28 influences the expansion and differentiation of T cells by silencing Treg-characteristic genes, thus influencing effector, regulatory and helper T cell phenotypes and autoimmunity in the intestinal tract.\textsuperscript{64} In addition, a TRIM30α-dependent signalling pathway initiated by the lymphotoxin-β receptor on macrophages was observed to downregulate the inflammatory response during DSS induction.\textsuperscript{66}

4 | TRIMS AND GENETIC SUSCEPTIBILITY TO IBD

TRIM20, encoded by \textit{MEFV}, is the most-studied member of the TRIM family that contributes to the genetic susceptibility of IBD.\textsuperscript{54–57,89–91} Recently, it was reported that TRIM20 not only affects the prevalence of IBD but also influences phenotypic characteristics or complications of IBD.\textsuperscript{89–91} Several studies have illustrated a correlation between human \textit{MEFV} mutation and paediatric colitis or IBD, among which IBD complicated with FMF is quite common.\textsuperscript{54–57} Recently, two cases of paediatric IBD were reported in gorillas’ model with typical clinical signs, higher levels of C reactive protein and calprotectin, and a clear response to steroids.\textsuperscript{92} Sequencing analysis showed that both of them shared a variant of \textit{MEFV}, which indicates the involvement of TRIM20 in paediatric IBD.\textsuperscript{92} With respect to the association between TRIM proteins and intestinal manifestations, Kasamaki et al. reported one case of \textit{MEFV} mutation with small bowel stenosis, although IBD had not been diagnosed.\textsuperscript{93} Moreover, Fidder et al. found that \textit{MEFV} mutations may be associated with the increased frequency of strictureing patterns and extraintestinal complications in patients with CD.\textsuperscript{90} The E148Q variant of \textit{MEFV} is associated with higher susceptibility to

| TRIM protein | Opportunistic infections | Tested cell organizations | Function | Reference |
|--------------|--------------------------|---------------------------|----------|-----------|
| 14 | MTB | RAW 264.7 macrophages, HEK293T, murine embryonic fibroblasts, and Lenti-X cells | TRIM14 negatively regulates macrophages' type I IFN response against MTB infection through a cGAS-dependent signalling pathway | 99 |
| 19 | HCMV | Human dermal fibroblast and HEK293 and HEK293T cells | HCMV IE1 Protein could bind to and disrupt TRIM19 to gain resistance towards type I interferon-related immune responses | 100 |
| 20 | C. diff | TNF\textsuperscript{−/−}, and TNFR1\textsuperscript{−/−}, TNFR2\textsuperscript{−/−} and wild type BM-derived macrophages | TRIM20 upregulation induced by TcdB can be observed on the basis of TNF/TNFRI axis | 51 |
| 22 | MTB | HEK 293 cells; Inflamed and non-inflamed intestinal and rectal tissues from patients with very early onset IBD; | TRIM22 influences the infection of MTB by modulating the NOD2 pathway, NF-κB pathway, apoptosis, and autophagy in macrophages or monocytes | 61 |
| 28 | HCMV | Human CD34\textsuperset{+} cells | The mTOR-mediated phosphorylation switch of TRIM28 contributes to the eruption of HCMV from the latency period by recruiting HPIα and SETDB1 to the viral genome and inducing the following transcriptional modulation process | 103 |
| 31 | HCMV | Intestinal tissue of CD patients and controls | HCMV-infected intestinal cells own a lower expression of TRIM31 amount and a higher bacterial load under the suppression of an Atg5/Atg7 independent autophagy pathway | 28 |
| 46 | C. diff | FHC human normal colon epithelial cells; C57/BL6J mice | TcdB induced colonic inflammation through the DUSP1/MAPKs and NF-κB signalling pathway, in which TRIM46 exerts essential regulatory roles | 104 |
| 62 | Fungus | TRIM62\textsuperset{−/−}, CARD9\textsuperset{−/−} and wide type mice | TRIM62 ubiquitinates CARD9, modulates the CARD9-Mediated signalling pathway, and regulates anti-fungal immunity in the intestine (C. albicans) | 72 |

Note: C. diff, Clostridium difficile; HCMV, human cytomegalovirus; MTB, mycobacterium tuberculosis; TcdB, Clostridium difficile toxin B.
perianal lesions in CD.\textsuperscript{91} Regarding UC, it was reported that \textit{MEFV} mutations may aggravate its clinical progress, and increase the possibility of colectomy.\textsuperscript{94} In addition, Stittich et al. performed sequence analysis on 38 IBD patients from five families and found that a missense mutation in the PRY/SPRY domain of TRIM11 (p.H414Y) was associated with an increased risk of IBD. The researchers also found that the TRIM11 mutation p.H414Y may increase the activity of the NF-κB promoter.\textsuperscript{95} NF-κB is a crucial downstream target in the microbe-sensing pathways mediated by CARD9 or NOD2, and its activation can promote the expression of multiple proinflammatory cytokines, so as to exacerbate intestinal inflammation and IBD participate in pathophysiology.\textsuperscript{87} Furthermore, TRIM family proteins are critical regulators of the NF-κB signalling pathway and act by functioning as ubiquitin E3 ligases.\textsuperscript{96} Therefore, the increased risk of IBD in patients with mutant TRIM11 may be related to the activation of the NF-κB signalling pathway.

5 | TRIMS AND OPPORTUNISTIC INFECTION IN IBD

A higher risk of severe infections as well as opportunistic infections in IBD patients has been reported, which is generally believed to be a result of the nature of the disease itself or the adverse events associated with the administration of immunosuppressants.\textsuperscript{97} Opportunistic infection can aggravate intestinal inflammation and enhance the prevalence of refractory IBD; thus, some opportunistic pathogens can contribute to IBD pathogenesis or progression.\textsuperscript{98} Human cytomegalovirus (HCMV), Clostridium difficile (C. diff), Mycobacterium tuberculosis (MTB) and \textit{Candida albicans} are some of the implicated pathogens, due to their substantial prevalence and incidence.\textsuperscript{98} TRIM family proteins have also been shown to be involved in opportunistic infections (Table 3).

HCMV usually erupts during the immunosuppressive period of IBD, which is reportedly induced by the mTOR-mediated TRIM28-related phosphorylation switch.\textsuperscript{103} Moreover, Ra et al. observed that intestinal cells showed relatively low TRIM31 expression after HCMV infection accompanied by a remarkable bacterial load, which was reversed by TRIM31 reintroduction.\textsuperscript{28} In addition, the HCMV Iκ protein could improve its resistance to antiviral immune responses by binding to and disrupting TRIM19.\textsuperscript{100} Another commonly observed opportunistic infection is caused by \textit{Clostridium difficile} toxin B (TcdB); one of its main virulence factors was reported to induce an inflammatory response in the colon via the TRIM46/DUSP1/MAPKs pathway, and TRIM46 deficiency can restrict colon inflammation.\textsuperscript{104} Conversely, Xu et al. found that TRIM20 mediates the innate immune response to TcdB as a protective role and Sharma et al. further illustrated that TcdB led to the upregulation of TRIM20 expression through the TNF/TNF pathway.\textsuperscript{51,101} For MTB, TRIM22 was shown to modulate the immune response through NOD2 or the NF-κB pathway, as well as through autophagy.\textsuperscript{63} MEVF mutations have been reported to protect against tuberculosis infection,\textsuperscript{102} while TRIM14 functions as an inhibitory modulator of the type I IFN response.\textsuperscript{99} As noted above, TRIM62 regulates anti-fungal immunity in the intestine by promoting CARD9-mediated signalling pathway, and its deficiency may increase susceptibility to fungal infections, such as those caused by \textit{Candida albicans}.\textsuperscript{72}

6 | THERAPEUTIC APPLICATIONS OF TRIM FAMILY PROTEINS IN IBD

As noted above, a plethora of TRIM family members have been shown to have protective or detrimental roles in the genetic susceptibility, pathogenesis and complications of IBD.\textsuperscript{105} Efforts have been made to exploit these functions of TRIM proteins in clinical practice.

One group reported that TRIM29 mediates the ubiquitination of stimulator of interferon genes (STING) and the STING-TBK1-IRF3 signalling pathway, thereby inhibiting interferon-I and pro-inflammatory cytokine production, which contributes to the essential roles of TRIM29 in many autoimmune diseases, including IBD.\textsuperscript{105} Subsequently, this group tried to develop a novel therapy by targeting the inhibition of TRIM29 via gene silencing for diseases such as IBD.\textsuperscript{105}

As noted above, TRIM31 downregulates intestinal inflammation in CD by inducing autophagy.\textsuperscript{28} Therefore, compounds comprising TRIM31 and its activator were developed by the same team for treating IBD.\textsuperscript{106} They also proposed the use of TRIM31 for the diagnosis of IBD or screening of IBD therapeutics.\textsuperscript{106} Diagnosis of IBD was achieved by measuring levels of the TRIM31 protein or mRNA in patient intestinal epithelial samples and comparing with control samples, while screening drugs would focus on comparing the expression of TRIM31 when administered or not administered.\textsuperscript{106}

7 | CONCLUSIONS

E3 ubiquitin ligase, which recognizes specific protein substrate and mediates its conjugation with ubiquitin, is a crucial molecule in ubiquitin–proteasome pathway for post-translational modification, so as to regulate the activity of different proteins in various biological process. The TRIM family is a group of highly conserved proteins with E3 ubiquitin ligase activity, which has been shown to be involved in multiple autoinflammatory diseases, including systemic lupus erythematosus, rheumatoid arthritis and, in particular, inflammatory bowel disease.\textsuperscript{107} A growing number of studies have demonstrated significant differences in TRIM protein expression between IBD patients and the general population, either in the gut or other tissues, suggesting their involvement in IBD. In this review, the relationships between TRIM proteins and IBD were discussed from three aspects: intestinal mucosal barrier function, gene susceptibility and IBD-related opportunistic infection. TRIM proteins have been shown to modulate the expression of many key molecules in the intestinal mucosal barrier, including tight junction proteins, antimicrobial peptides and mucin. Moreover, TRIM proteins participate in multiple IBD-related signalling pathways, including microbial recognition (e.g., NOD2, CARD9 and TLR2 pathways), immune cell differentiation (e.g., IL-6/
Nevertheless, few studies on families have been implicated in IBD (Table 1). It is difficult to identify specific regularity between TRIM subfamilies and their roles in IBD, despite the structural and functional similarities within each subfamily. Moreover, most studies evaluating the role of TRIM in IBD have centred on TRIM20—for which retrospective studies usually predominates—while underlying mechanisms remain unclear. Furthermore, the TRIM family is widely accepted as a regulator of tumour oncogenesis or suppression, and its role in colorectal tumours has been reported by several studies. Nevertheless, few studies on TRIM proteins and IBD-related carcinogenesis have been reported, so efforts should be made to explore this critical issue. Further clarification of these questions will enable the development of emerging IBD therapies based on the modulation of TRIM family proteins.

CONFLICT OF INTERESTS

The authors declare that they have no conflicts of interest.

AUTHOR CONTRIBUTIONS

R.C. and Y.T. wrote and revised the manuscript. J.L. drafted the figures. L.L., Z.Z., M.C. and S.Z. revised the paper. All authors approved the final vision of the manuscript and the submission.

DATA AVAILABILITY STATEMENT

DATA AVAILABILITY STATEMENT Data sharing is not applicable to this article, as no new data were created or analyzed in this study.

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REFERENCES

1. Ng SC, Shi HY, Hamidi N, et al. World incidence and prevalence of inflammatory bowel disease in the 21st century: a systematic review of population-based studies. Lancet. 2017;390(10114):2769-2778.
2. Roda G, Chien NS, Kotze PG, et al. Crohn’s disease. Nat Rev Dis Primers. 2020;6(1):22.
3. Ungaro R, Mehandru S, Allen PB, Peyrin-Biroulet L, Colombel JF. Ulcerative colitis. Lancet. 2017;389(10080):1756-1770.
4. Malodecky NA, Soon IS, Rabi DM, et al. Increasing incidence and prevalence of the inflammatory bowel disease with time, based on systematic review. Gastroenterology. 2012;142(1):46-54.e42. quiz e30.
5. Zhao M, Gönczi L, Lakatos PL, Burisch J. The burden of inflammatory bowel disease in Europe in 2020. J Crohns Colitis. 2021;15(9):1573-1587.
6. Kaplan GG. The global burden of IBD: from 2015 to 2025. Nat Rev Gastroenterol Hepatol. 2015;12(12):720-727.
7. Baumgart DC, Le Berre C. Newer biologic and small-molecule therapies for inflammatory bowel disease. N Engl J Med. 2021;385(14):1302-1315.
8. Singh S, George J, Boland BS, Vande CN, Sandborn WJ. Primary non-response to tumor necrosis factor antagonists is associated with inferior response to second-line biologics in patients with inflammatory bowel diseases: a systematic review and meta-analysis. J Crohns Colitis. 2018;12(6):635-643.
9. Frolikis AD, Dykeman J, Negrón ME, et al. Risk of surgery for inflammatory bowel diseases has decreased over time: a systematic review and meta-analysis of population-based studies. Gastroenterology. 2013;145(5):996-1006.
10. Chang JT. Pathophysiology of inflammatory bowel diseases. N Engl J Med. 2020;383(27):2652-2664.
11. Schoutz I, Keita ÅV. Cellular and molecular therapeutic targets in inflammatory bowel disease-focusing on intestinal barrier function. Cells-Basel. 2019;8(2):193.
12. Vancamelbeke M, Vermeire S. The intestinal barrier: a fundamental role in health and disease. Expert Rev Gastroenterol Hepatol. 2017;11(9):821-834.
13. Martini E, Krug SM, Siegmund B, Neurath MF, Becker C. Mend your fences: the epithelial barrier and its relationship with mucosal immunity in inflammatory bowel disease. Cell Mol Gastroenterol Hepatol. 2017;4(1):33-46.
14. Hatakeyama S. TRIM family proteins: roles in autophagy, immunity, and carcinogenesis. Trends Biochem Sci. 2017;42(4):297-311.
15. Di Rienzo M, Romagnoli A, Antonioli M, Piacentini M, Fixia GM. TRIM proteins in autophagy: selective sensors in cell damage and innate immune response. Cell Death Differ. 2020;27(3):887-902.
16. Zhao G, Liu C, Wen X, et al. The translational values of TRIM family in pan-cancers: from functions and mechanisms to clinics. Pharmacol Ther. 2021;227:107881.
17. Gushchina LV, Kwiatkowski TA, Bhattacharya S, Weisleder NL. Conserved structural and functional aspects of the tripartite motif gene family point towards therapeutic applications in multiple diseases. Pharmacol Ther. 2018;185:12-25.
18. Shen Z, Wei L, Yu ZB, et al. The roles of TRIMs in antiviral innate immune signaling. Front Cell Infect Microbiol. 2021;11:628275.
19. Ozato K, Shin DM, Chang TH, Morse HR. TRIM family proteins and their emerging roles in innate immunity. Nat Rev Immunol. 2008;8(11):849-860.
20. Jin Z, Zhu Z. The role of TRIM proteins in PRR signaling pathways and immune-related diseases. Int Immunopharmacol. 2021;98:107813.
21. Massiah MA, Simmons BN, Short KM, Cox TC. Solution structure of the RBCC/TRIM B-box1 domain of human MID1: B-box with a RING. J Mol Biol. 2006;358(2):532-545.
22. Khan R, Khan A, Ali A, Idrees M. The interplay between viruses and TRIM family proteins. Rev Med Virol. 2019;29(2):e2028.
23. Bell JL, Maluykova A, Holien JK, et al. TRIM16 acts as an E3 ubiquitin ligase and can heterodimerize with other TRIM family proteins. J Mol Biol. 2006;358(2):532-545.
24. Xing J, Weng L, Yuan B, et al. Identification of a role for TRIM29 in the control of innate immunity in the respiratory tract. Nat Immunol. 2016;17(12):1373-1380.
25. Pauletto E, Eickhoff N, Padró NA, Blattner C, Zwart W. TRIMming down hormone-driven cancers: the biological impact of TRIM proteins on tumor development, progression and prognostication. Cells-Basel. 2021;10(6):1517.
26. Li X, Yeung DF, Fiegen AM, Sodroski J. Determinants of the higher order association of the restriction factor TRIM5alpha and other tripartite motif (TRIM) proteins. J Biol Chem. 2011;286(32):27959-27970.
90. Fidder H, Chowers Y, Ackerman Z, et al. The familial Mediterranean fever (MEVF) gene as a modifier of Crohn’s disease. *J Immunol*. 2019;203(6):1636-1649.

91. Petersen BS, Bokemeyer B, Wenker C, et al. First known case of paediatric inflammatory bowel disease in a western lowland gorilla may be linked to a familial mutation in the MEFV gene. *Gut*. 2020; 69(6):1153-1154.

92. Kasamaki K, Kusano C, Ikehara H, et al. Familial Mediterranean fever with small bowel stenosis. *Intern Med*. 2019;58(14):2025-2028.

93. Petersen BS, Bokemeyer B, Wenker C, et al. Tripartite motif family proteins in inflammatory bowel disease: Mechanisms and potential for interventions. *Cell Prolif*. 2022;55(5):e13222. doi:10.1111/cpr.13222.