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Authors
Shih, David Q
Barrett, Robert
Zhang, Xiaolan
et al.

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Constitutive TL1A (TNFSF15) Expression on Lymphoid or Myeloid Cells Leads to Mild Intestinal Inflammation and Fibrosis

David Q. Shih1, Robert Barrett1, Xiaolan Zhang1,2, Nicole Yeager1, Hon Wai Koon3, Piangwarin Phaosawasdi1, Yahu Song3, Brian Ko1, Michelle H. Wong1, Kathrin S. Michelsen1, Gislaine Martins1, Charalabos Pothoulakis3, Stephan R. Targan1

1Inflammatory Bowel and Immunobiology Research Institute, Cedars-Sinai Medical Center, Los Angeles, California, United States of America, 2Department of Gastroenterology, The Second Hospital of Hebei Medical University, Shijiazhuang, China, 3Division of Digestive Diseases, Inflammatory Bowel Disease Center, David Geffen School of Medicine, University of California Los Angeles, Los Angeles, California, United States of America

Abstract

TL1A is a member of the TNF superfamily and its expression is increased in the mucosa of inflammatory bowel disease patients. Moreover, a subset of Crohn’s disease (CD) patients with the risk TL1A haplotype is associated with elevated TL1A expression and a more severe disease course. To investigate the in vivo role of elevated TL1A expression, we generated two transgenic (Tg) mice models with constitutive Tl1a expression in either lymphoid or myeloid cells. Compared to wildtype (WT) mice, constitutive expression of Tl1a in either lymphoid or myeloid cells showed mild patchy inflammation in the small intestine, which was more prominent in the ileum. In addition, mice with constitutive Tl1a expression exhibited enhanced intestinal and colonic fibrosis compared to WT littermates. The percentage of T cells expressing the gut homing chemokine receptors CCR9 and CCR10 was higher in the Tl1a Tg mice compared to WT littermates. Sustained expression of T11a in T cells also lead to increased Foxp3+ Treg cells. T cells or antigen presenting cells (APC) with constitutive expression of Tl1a were found to have a more activated phenotype and mucosal mononuclear cells exhibit enhanced Th1 cytokine activity. These results indicated an important role of TL1A in mucosal T cells and APC function and showed that up-regulation of TL1A expression can promote mucosal inflammation and gut fibrosis.

Introduction

Inflammatory bowel disease (IBD), encompassing CD and ulcerative colitis (UC), is a chronic inflammatory disorder caused by dysregulated immune responses in a genetically predisposed individual (reviewed in [1]). CD is a chronic inflammatory condition that predominately affects the gut with distinctive pathological features such as patchy transmural intestinal inflammation, relative sparing of inflammation in the rectum, intestinal fibrostenosis and a dysregulated T helper (Th) 1 and Th17 immune response [1,2]. Accumulating data, including genome-wide association studies (GWAS), demonstrate that more than 80 distinct genetic loci confer CD susceptibility, and are being used to define critical molecules and pathways that converge in physiologic processes that lead to mucosal inflammation [2,3]. Among the several recently discovered IBD associated gene variants, only those in the tumor necrosis factor superfamily member 15 (TNFSF15) have been shown to be associated with CD in all ethnic and age groups [1].

TL1A plays an important role in modulating the adaptive immune response. In the Th1 effector immune response, binding of TL1A to its receptor (death domain receptor 3, DR3, TNFRSF25) enhances IFN-γ production from peripheral and mucosal T cells [4,5]. Studies have shown that TL1A can be induced in APC by FcγR signaling [6,7] and microbial antigen/organisms [8] suggesting that augmentation of the Th1 immune response by TL1A may occur through APC-T cell interactions. In murine models of chronic mucosal inflammation, TL1A enhanced Th1 and Th17 effector function by up-regulating IFN-γ and IL17 production, respectively, in gut-associated lymphoid tissue (GALT) CD4+ cells under Th1/Th17 polarizing conditions, indicating that TL1A is an important modulator in the development of gut mucosal inflammation [9]. In addition to mediating Th1 responses, TL1A also promotes Th2 and Th17 effector cell function [10,11,12].

Several studies implicate the TL1A/DR3 signaling pathway in mucosal inflammation. The expression of TL1A and DR3 is up-regulated in the inflamed gut mucosa of two distinct murine models of ileal inflammation [13]. Neutralizing anti-mouse TL1A Ab attenuated inflammation in both the dextran sulfate sodium induced chronic colitis and a G protein β2-/- (Gα2ζ–/–) T cell transfer colitis model [9]. In humans, TL1A can enhance IFN-γ production in CD4+ T cells expressing the gut-homing receptor CCR9 [14]. Increased expression of both TL1A and DR3 was found in gut mucosal biopsies and lamina propria (LP) T cells of

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* E-mail: targans@cshs.org
both UC and CD patients [4,15]. Furthermore, TL1A and DR3 expression are correlated with severity of gut mucosal inflammation as their transcripts were several times more abundant in RNA from mucosal biopsies taken from inflamed CD lesions than in those taken from uninvolved areas [4,15].

To determine the in vivo consequence of increased TL1A expression, we generated two Tg murine models that constitutively express TL1A in either lymphoid or myeloid cells. We found that constitutive expression of TL1A in either lymphoid or myeloid cells induced mild spontaneous patchy intestinal inflammation by 10 months of age. We showed that a higher percentage of T cells and APC from the mesenteric lymph nodes (MLN) have an activated phenotype and express the gut homing chemokine receptors CCR9 and CCR10, associated with increased production of IFN-γ. Consistent with previous reports [16,17], we observed goblet cell hyperplasia and an increased number of Paneth cells in mice that constitutively express TL1A. These 2 novel murine TL1A Tg models have patterns of site directed mucosal inflammation and fibrosis seen in human CD and may be useful models to study the pathogenesis of IBD.

Results

Generation of in vivo constitutive TL1a expression in the myeloid and T cell lineage

To investigate the contribution of sustained APC or T cell TL1a expression on gut mucosal homeostasis and inflammation, we generated Tg mice that constitutively express TL1A in either T cells or myeloid cells. We used the proximal lck promoter and CD2 enhancer to drive T cell lineage-specific expression and the c-fms promoter to mediate myeloid specific expression in APC such as macrophages and DC [18,19]. We also cloned an IRES-GFP element downstream of the murine TL1A so that Tg TL1A expressing cells could be identified by GFP. The cloning strategies and schematic of the Tg construct are described in Materials and Methods section and figure 1A, respectively. The Tg constructs were injected into C57BL/6 pronuclei to ensure genetic homogeneity except for the TL1A transgene.

Expression of the transgene was determined in myeloid Tg mice (named FMS-Tl1a-GFP Tg) and T cell lineage-specific expression of TL1A (named LCK-CD2-Tl1a-GFP Tg) by the expression of GFP. We found that in the FMS-Tl1a-GFP Tg mice, GFP was present in over 70% of CD11c and F4/80 positive cells, and the expression of the TL1A transgene persisted as the mice aged (Fig. 2A). The specificity of the c-fms promoter was illustrated by the fact that we did not detect GFP expression in CD3, CD4 or CD8 positive T cells (Fig. 1C). In the LCK-CD2-Tl1a-GFP Tg mice, GFP was present in over 90% of CD3, CD4 and CD8 positive T cells (Fig. 1C). Similar to the FMS-Tl1a-GFP Tg, the expression of the transgene in the lymphoid Tl1a Tg mice also persisted as the mice got older (Fig. 1C). Less than 10% GFP expression was detected in F4/80 or CD11c positive cells (Fig. 1B), indicating that the Lck promoter and cd2 enhancer elements drive T cell lineage-specific expression. We directly showed that TL1A mRNA is higher in the spleen, mesenteric lymph nodes (MLN), colon and ileum of both Tg mice compared to WT mice (Fig. 1D). Together, these data demonstrate that we generated tissue specific constitutive in vivo expression of TL1A in APC and T cells.

Mice with constitutive TL1a expression do not develop gross tissue inflammation

Both FMS-Tl1a-GFP and LCK-CD2-Tl1a-GFP Tg mice are fertile, but are born at less than Mendelian frequency. Mendelian expectation is 50% transmission of the transgene. When Tg hemizygous mice are mated to WT mice, the frequency of FMS-Tl1a-GFP Tg mice born is 42% (58/138) and in LCK-CD2-Tl1a Tg mice, the frequency is 40% (51/126). Both the FMS and LCK-Tl1a Tg mice appeared healthy and gained weight at similar rates (Fig. 2A). There were no differences in the disease activity index (DAI) [20] between Tg and WT for up to 10 months (Fig. 2B). We also did not observe differences in the splenic cell number, MLN cell number, lamina propria mononuclear cell (LPMC) number in the small bowel or colon, colon length, small bowel (SB) length or spleen size between WT and TL1A Tg mice at 2 or 10 months (Fig. 2C and data not shown). There was a trend toward higher cell numbers in the MLN and SB LPMC in Tl1a Tg mice but they did not reach statistical significance (Fig. 2B and data not shown).

TL1a expression in APC and T cells induce mild histologic small bowel inflammation

Elevated TL1a expression is implicated in gut mucosal inflammation. We therefore investigated whether mice with elevated TL1a expression develop spontaneous colitis at 2 months and 10 months of age. The colon and small intestine did not show gross inflammation between WT and Tg mice using a standard macroscopic scoring system (Fig. 3A) [21]. Another measure for gut inflammation is to determine myeloperoxidase (MPO) activity [22]. We found significantly increased MPO activity in the small intestine of LCK-CD2-Tl1a Tg mice than WT mice (Fig. 3B). MPO activity was similar in the colon of WT and Tl1a Tg mice (Fig. 3B).

Histological examination of the colon did not reveal increased inflammatory infiltrates, mucin depletion, epithelial cell hyperplasia, abnormal crypt architecture, crypt abscess or erosions in either WT or Tg mice at 2 or 10 months of age (Fig. 3C and data not shown). As elevated TL1A production in IBD patients is associated with fibrostenotic disease [23,24], we assessed whether the TL1a Tg mice had increased histologic fibrosis. Notably, we observed increased fibrosis in the colonic mucosa and submucosa of both LCK-CD2- and FMS-Tl1a Tg mice as compared to WT littermate mice by 10 months of age using the Masson Trichrome stain (Fig. 3D).

At 2 months of age, histologic examination of the small intestine revealed a significant increase in the number of goblet cells and Paneth cells (Fig. 4A and 4B). There was blunting of the villi and increased LPMC in the ileum of both the LCK-CD2- and FMS-Tl1a Tg mice as compared to WT mice (Fig. 4A). These histological changes were reflected by a significant increase in the inflammatory score in the ileum of Tl1a Tg mice using a standard quantitative scoring system (Fig. 4B) [21]. We did not observe villus blunting nor increased mononuclear cells in the LP of the duodenum and jejunum between WT and Tl1a Tg mice at 2 months of age (Fig. 4A).

At 10 months, we similarly observed Paneth cell hyperplasia in both LCK-CD2- and FMS-Tl1a Tg compared to WT mice (Fig. 5A and 5B). In contrast to younger mice (2 months of age), there were no detectable differences in the number of goblet cells between WT and Tg mice at 10 months of age (Fig. 5A and 5B). The inflammatory changes in the small intestine such as increased mononuclear cell infiltrate of the LP and blunting of the villi was more prominent and progressive, involving the duodenum, jejunum and ileum (Fig. 5A and 5B). The increased inflammation in both LCK-CD2- and FMS-Tl1a Tg mice was associated with increased histologic fibrosis by the more extensive Masson Trichrome stain in the small intestine (Fig. 6). Together, these results indicated that constitutive expression of TL1A in T cells and myeloid cells lead to progressive spontaneous intestinal inflammation and fibrosis.
Mice with constitutive T11a expression develop extra-intestinal pathology

At a low frequency, we observed extra-intestinal pathology in both the FMS-T11a- and LCK-CD2-T11a Tg mice. One such feature was an erythematous ulcerated skin lesion that was observed in 1 out of 58 FMS-T11a Tg, 2 out of 51 LCK-CD2-T11a Tg and 1 out of 155 WT littermate mice (Fig. 7). Another observed pathology was joint erythema and swelling (Fig. 7) that caused movement difficulties and resulted in mice not able to feed. Arthropathy was observed in 2 out of 38 FMS Tg, 2 out of 51 LCK-CD2 T11a Tg mice and 0 out of 155 WT mice. Three out of 4 Tg mice with arthropathy had monoarticular and 1 LCK-CD2-T11a Tg mice had polyarticular

Figure 1. Generation of constitutive in vivo expression of T11a in T- and antigen presenting cells. (A) Schematic of LCK-CD2-T11a and FMS-T11a transgenic construct. An internal ribosomal entry site (IRES) element is used for both transgenic constructs so that a bicistronic message can be made from the transgene and the T11a expressing cells are tagged by GFP. (B) Flow cytometric analysis of the transgene marker GFP on either CD11c or F4/80 gated splenocytes from WT (black filled), LCK-CD2-T11a Tg mice (L-Tg, dotted line) or FMS-T11a Tg mice (M-Tg, solid grey line). Representative histograms are shown. (C) Representative analysis of the transgene marker GFP on either CD3, CD4 or CD8 gated splenocytes from WT (black filled), LCK-CD2-T11a Tg mice (L-Tg, dotted line) or FMS-T11a Tg mice (solid grey line). (D) T11a mRNA expression was determined in the spleen, MLN, ileum or colon by real-time polymerase chain reaction. Data are expressed as mean percent of β-actin ± standard deviation (SD). *P<0.05, **P<0.01. n = 6 independent littermate mice per group were used for B-D.

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disease. Our data suggested these novel murine models with constitutive expression of Tl1a in APC and T cells develop extraintestinal pathology such as ulcerated skin lesion and arthropathy.

Accelerated T- and antigen presenting cell activation in LCK-CD2- and FMS-Tl1a Tg mice

To assess whether constitutively expressed Tl1a can co-stimulate T cells in vivo, we compared the expression of an activation marker on CD4⁺ and CD8⁺ cells between Tg and WT littermate controls. CD4⁺CD45RB⁺⁺CD25⁺ Treg cells were gated out in order to examine the expression of activation markers on conventional T cells. At 2 months, there were almost 2 fold higher CD4⁺CD44⁺ T cells from the spleen but not MLN of LCK-CD2-Tl1a Tg mice (Fig. 8A). There was also no difference in the expression of the activation marker CD44 on CD8⁺ T cells in the MLN and spleen at 2 months of age between Tl1a Tg mice and WT littermates (Fig. 8A). By 10 months of age, a higher percentage of Tl1a Tg CD4⁺ cells in both the spleen and MLN expressed the activation marker CD44, particularly in the LCK-CD2-Tl1a Tg mice (Fig. 8B). In contrast, only CD8⁺ cells from the LCK-CD2-Tl1a Tg (not FMS-Tl1a Tg) spleen and MLN exhibited an increased expression of the activation marker CD44 (Fig. 8B).

To assess the effect of constitutive Tl1a expression on the activation state of DC and macrophages, the expression of the
activation marker CD86 was compared between Tl1a Tg and WT littermate controls. Flow cytometric (FACS) analysis revealed increased CD86$^+$ expression on DC (CD11c$^+$) and macrophages (F4/80$^+$) in both the spleen and MLN of Tl1a Tg mice compared to WT mice (Fig. 9A). In contrast to LCK-CD2-Tl1a Tg mice, there was a negligible further increase in the percentage of activated DC and macrophages over time (compare 2 and 10 months, Fig. 9A and B). These data demonstrated that sustained Tl1a expression could result in enhanced activation of CD4$^+$, CD8$^+$, DC and macrophages in vivo.

Constitutive in vivo Tl1a expression leads to an increased number of T cells expressing Treg and gut homing markers

FACS analysis did not reveal any differences in the frequencies of CD3$^+$, CD4$^+$, CD8$^+$, MHCI$^+$, CD11c$^+$ or F4/80$^+$ cells in the spleen and MLN between FMS-Tl1a Tg, LCK-CD2-Tl1a Tg or WT littermate mice (data not shown). There was an increase in the frequency of Foxp3$^+$ cells in the spleen and MLN of LCK-CD2-Tl1a Tg mice (Fig. 10A and B). The percentage of Foxp3$^+$ cells further increased as the LCK-CD2-Tl1a Tg mice aged (Fig. 10A and B). In contrast, there was no difference in the frequency of Foxp3$^+$ cells in the spleen or MLN of FMS-Tl1a Tg mice at both 2 and 10 months (Fig. 10A and B).

A trend toward a higher cell number in the MLN and small bowel LPMC was noted in older (Fig. 2C and 5A). To determine whether this finding was due to increased trafficking to the gut immune compartment, we assessed the expression of the gut homing markers CCR9 and CCR10 in the MLN and spleen of Tl1a Tg and WT littermate mice. In the spleen, there was no difference in CCR9$^+$ or CCR10$^+$ cells at 2 or 10 months between either Tl1a Tg or WT mice (data not shown). Notably, we found an increase in the percentage of cells expressing CCR9 and CCR10 in the MLN of LCK-CD2-Tl1a Tg mice compared to WT mice.

Figure 3. Tl1a Tg mice do not develop gross intestinal inflammation but exhibit enhanced colonic fibrosis. (A) Gross appearance (wall thickening, hyperemia, rigidity or adhesions) of small intestine and colon are measured from 10 months old WT, LCK-CD2-Tl1a Tg (L-Tg) or FMS-Tl1a Tg (M-Tg) mice using a standard scoring system [21]. Data are expressed as mean ± SD. (B) Myeloperoxidase (MPO) activity is measured on the distal 3 cm of ilea and mid-colon and data are expressed as arbitrary unit (U) per gram (g) of protein. *P<0.05 (C) Representative hematoxylin and eosin (H&E) stained colon section obtained from mid-colon of 10 months old WT, LCK-CD2-Tl1a Tg or FMS-Tl1a Tg mice are shown. (D) Masson Trichrome staining of collagen deposition in tissue sections of mouse mid-colon. Collagen is stained blue versus red background. There are increased blue collagen stain in LCK-CD2-Tl1a Tg and FMS-Tl1a Tg compared to WT littermate mice. Magnification 200X. Results are representative of six mice per group for A–D.

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 Constitutive TIl1a expression leads to increased numbers of goblet and Paneth cells in the small intestine and ileal histological inflammation. (A) Representative H&E stained sections obtained from the indicated portions of small intestine from 2 months old WT, LCK-CD2-TIl1a Tg (L-Tg) or FMS-TIl1a Tg (M-Tg) mice are shown. Goblet cells are denoted by an open arrow. Paneth cells are denoted by a filled arrow. Results are representative of six mice per group. Magnification 200X. (B) The numbers of goblet (top panel) and Paneth cells (middle panel) were determined by examining at least 80 individual villi at the indicated portions of the small intestine from six mice (2 months old) per group by 2 observers blinded to mouse genotype. Data are expressed as mean (SD). Histologic scores (bottom panel) were determined by 2 observers blinded to mice using standard methods [21]. Data are expressed as mean (SD). At least 36 fields from 6 mice per group at 200x magnification were scored.

*P<0.05, **P<0.01.

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Figure 5. Persistent Paneth cell hyperplasia and worsened small intestinal inflammation as the Tl1a Tg mice aged. (A) Representative H&E stained section obtained from the indicated portions of small intestine from 10 months old WT, LCK-CD2-Tl1a Tg (L-Tg) or FMS-Tl1a Tg (M-Tg) mice are shown. Goblet cells are denoted by an open arrow. Paneth cells are denoted by a filled arrow. Results are representative of six mice per group. Magnification 200X. (B) The numbers of goblet (top panel) and Paneth cells (middle panel) were determined by 2 observers blinded to mice genotype. Histologic scores (bottom panel) were determined from 10 months old WT, LCK-CD2-Tl1a Tg (L-Tg), or FMS-Tl1a Tg (M-Tg) mice using standard methods [21]. Quantification of goblet and Paneth cells was determined by examining at least 80 individual villi and histological scores were determined by examining at least 36 fields at 200X magnification. Six independent mice per group were used (A–B). Data are expressed as mean (SD.

*P<0.05, **P<0.01.

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mice at 2 months and the difference became even greater as the mice aged (Fig. 11A). For the FMS-Tl1a Tg mice, a higher percentage of CCR9\(^+\) and CCR10\(^+\) cells was observed at 10 months (but not at 2 months) (Fig. 11A).

**Tl1a Tg mice have an enhanced proinflammatory cytokine profile in the MLN and small intestine**

To assess the molecular consequences of increased expression of activation and gut homing marker, we measured the expression of IFN-\(\gamma\), IL-13 and IL-17 by FACS analysis and ELISA. There was no difference in the expression of CD4\(^+\)IFN-\(\gamma\)^, CD4\(^+\)IL-13^, CD4\(^+\)IL-17^ T cells by either intracellular stain or ELISA at 2 months of age between FMS-Tl1a Tg, LCK-CD2-Tl1a Tg or WT littermate mice (data not shown). By 10 months of age, the frequency of CD4\(^+\)IFN-\(\gamma\)^ and CD4\(^+\)IL-17^ T cells increased by approximately 3- and 2- fold respectively, in the MLN of LCK-CD2-Tl1a Tg mice (Fig. 11B). In the FMS-Tl1a Tg mice, the percentage of CD4\(^+\)IFN-\(\gamma\)^ cells increased by approximately 2-fold (Fig. 11B). We did not observe any differences in the CD4\(^+\)IL-13^ T cells in both of the Tl1a Tg mice and only a negligible difference in the CD4\(^+\)IL-17^ T cells in FMS-Tl1a Tg mice (Fig. 11B).

To confirm the FACS findings, we isolated cells from the spleen, MLN and LPMC from the colon and small intestine and assessed their ability to produce cytokines following stimulation with anti-CD3 and anti-CD28. Similar to the intracellular stain, we found significantly higher IFN-\(\gamma\) production in the MLN and small intestine LPMC of LCK-CD2-Tl1a Tg mice (Fig. 12A). FMS-Tl1a Tg mice also exhibited significantly increased IFN-\(\gamma\) production in MLN and a trend toward a higher level in the LPMC from the small intestine (Fig. 12A). There was also a trend toward higher IL-17 and IL-13 production in the MLN and small intestine LPMC of both LCK-CD2-Tl1a and FMS-Tl1a Tg mice, but this did not reach statistical significance (Fig. 12B and C). Interestingly, the expression of the anti-inflammatory cytokine IL-10 appeared to be higher in both the Tl1a Tg mice and reached statistical significance in the splenic cells of LCK-CD2-Tl1a Tg as compared to WT mice (Fig. 12D). These data suggested that constitutive expression of Tl1a in vivo resulted in enhanced IFN-\(\gamma\) and IL-10 production and potentially enhanced IL-17 and IL-13 production.

**Discussion**

This study shows that constitutive expression of Tl1a on either T- or APC cells leads to progressive histological inflammation in the small intestine and worsened fibrosis in both the small intestine and colon over time. Even though sustained expression of Tl1a in T- or APC did not result in gross macroscopic gut inflammation or colitis symptoms as measured by DAI (weight loss, fecal occult blood or loose stool) for up to 1 year, there were histologically determined inflammation as evidenced by blunting of the small intestinal villi, hyperplasia of goblet and Paneth cells and increased LPMC infiltrate. These histological changes are markers of early gut inflammation that are also found in other murine models of colitis and ileitis [25,26]. As the small bowel inflammation worsened with age, there was no difference in the goblet cell number between Tl1a Tg and WT littermate mice (Fig. 4 and 5). This was likely due to the fact that goblet cell number decreased with worsened inflammation in older Tl1a Tg mice [26,27,28]. This histologic inflammation was only detected in the small intestine (not colon) and was particularly evident in the ileum. The degree of inflammation in the small intestine was similar between Tg mice with sustained Tl1a expression in T- and myeloid cells (Fig. 4).
Another histological finding was increased small intestinal and colonic fibrosis in T11a Tg compared to WT littermate mice by 10 months of age (Fig. 3D and 6). This could be due to the fact that there was higher transgene expression in the LCK-CD2-T11a Tg mouse (approximately 90% of T cells) compared to its expression in APC (approximately 70% of DC and macrophages). Alternatively, constitutive T11a expression in lymphocytes could be more fibrogenic than its expression on either DC or macrophages. The presence of colonic fibrosis in the absence of detectable histologic inflammation suggested that T11a may be a pro-fibrogenic factor in addition to its role in inflammation. Notably, TL1A haplotype B which is associated with increased secretion of soluble and

Figure 8. Sustained T11a expression leads to an increased percentage of activated T cells. FACS plot of 2 month (A) and 10 month (B) splenocytes and MLN cells showing expression of activation markers CD44. Either CD4⁺ or CD8⁺ cells are gated as indicated. Data shown are representative of 4 mice per group. WT = wildtype, L-Tg = LCK-CD2-T11a-GFP Tg mice, M-Tg = FMS-T11a Tg mice. doi:10.1371/journal.pone.0016090.g008
membrane TL1A is also characterized by increased need for surgery especially in Jewish patients [23,24]. Fibrostenotic disease was also more common in this group. It would be medically useful to determine whether downregulating the TL1A/DR3 pathway can prevent the progression of IBD-associated fibrosis since there is currently no effective medical therapy to treat established fibrostenotic disease in IBD patients. It would be interesting to study the interaction between TL1A and other known profibrogenic factors such as transforming growth factor beta 1 (TGF-β1) and insulin-like growth factor 1 (IGF-1) using the chronic 2,4,6-trinitrobenzensulfonic acid (TNBS) colonic injury model which causes both colitis and intestinal fibrosis [29,30,31].

Figure 9. Sustained Tl1a expression leads to an increased percentage of activated DC and macrophages. FACS plot of 2 month (A) and 10 month (B) splenocytes and MLN cells showing expression of activation markers CD86. Either F4/80$^+$ or CD11c$^+$ cells were gated as indicated. Data shown are representative of 4 mice per group. WT = wildtype, L-Tg = LCK-CD2-Tl1a-GFP Tg mice, M-Tg = FMS-Tl1a Tg mice.

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We observed a low frequency of extra-intestinal pathology in both LCK-CD2-Tl1a Tg and FMS-Tl1a Tg mice. These included ulcerated skin lesions and erythematous swollen joints (fig. 7), which are similar to known extra-intestinal manifestations associated with human IBD. The cause of the extra-intestinal pathology in Tl1a Tg mice remains to be investigated, but may be related to the differential expression of CCR10 homing molecules in the Tl1a Tg mice. In addition to its function as a gut mucosal homing receptor [32], CCR10 is important in lymphocyte trafficking to the skin [33,34].

The chemokine receptors CCR9 and CCR10 play a major role in in vivo lymphocyte trafficking to portals of microbial entry, such as gut mucosal tissues and the skin [33,34,35,36,37]. The higher LPMC infiltrate in the small intestine as compared to the colon in Tl1a Tg mice may be due to the fact that CCR9 is preferentially found in the small intestine [38] and its ligand CCL25/TECK is expressed in both lamina propria venules and small intestine enterocytes [39,40]. In contrast, colonic expression of CCR9 and its ligand CCL25/TECK is limited [39,40]. The clinical importance of dysregulated immune trafficking in IBD is highlighted by the efficacious therapies such as natalizumab that blocks gut homing in inducing and maintaining remission in CD [41].

We found that constitutive expression of Tl1a resulted in a progressive activated phenotype over time in CD4+ and CD8+ T cells (Fig. 8), macrophages and DC (Fig. 9). Increased T cell and APC activation in Tl1a Tg mice may be a direct consequence of constitutive TL1A costimulation of DR3 expressing T cells and macrophages [11,42,43] or indirectly through failure of immunological tolerance leading to enhanced activation. In fact, recent studies have shown that the TL1A-DR3 pathway can attenuate the suppressive effect of Tregs on effector T cells [16,17]. The activated phenotype of T and APC in combination with the constitutive costimulation of TL1A-DR3 signaling pathway may lead to the significantly higher IFN-\(\gamma\) expression (Fig. 11B and 12A) that we saw in the Tl1a Tg mice, which is consistent with the previous observation that TL1A is involved in mediating the Th1 response [4,5,8,9,13,14,15,44,45]. There was also a trend toward increased IL-13 and IL-17 expression in mice with constitutive Tl1a expression (Fig. 12B and C), which was consistent with the

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**Figure 10. Increased numbers of regulatory T (Treg) cells in Tl1a Tg mice.** FACS plot of CD4+ Foxp3+ splenocytes (A) or CD4+ Foxp3+ MLN cells (B) are shown. Data shown are representative of 4 mice per group at either 2 or 10 months of age. WT = wildtype, L-Tg = LCK-CD2-Tl1a-GFP Tg mice, M-Tg = FMS-Tl1a-GFP Tg mice.

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role of TL1A in Th2- and Th17-mediated functions in various mouse models [9,10,11,12,16,17]. Lastly, we found significantly increased IL-10 production in LCK-CD2-Tl1a Tg mice (Fig. 12D), which could be related to higher percentage of Foxp3+ Treg cells in LCK-CD2-Tl1a Tg as compared to FMS-Tl1a Tg or WT mice (Fig. 10A and B).

Recently, two groups independently reported the generation of transgenic mice with sustained TL1A expression in DC and T cells. Our results were similar to the constitutive Tl1a expressing Tg mice in Meylan et al. and Taraban et al., with the increased goblet, Paneth and Treg cells, ileitis and activated phenotype in T and APC cells [16,17]. However, a major difference was that the...
ileitis found in the recently published Tl1a Tg mice appeared to be mainly attributed to Th2/IL-13 effector pathway [16,17], whereas we observed a higher production of IFN-γ from the Tl1a Tg mice generated in this study (Fig. 1B and 12A). In addition, we observed extra-intestinal pathology and increased gut fibrosis in the Tl1a Tg mice, which were not reported in the other Tl1a Tg mice. Several reasons may account for the phenotypical differences between the Tl1a Tg mice. One explanation may be methodological differences. For example, different promoter/enhancer elements were used to drive tissue specific expression of the different Tl1a Tg mice. We used the LCK-CD2 promoter/enhancer element [19], whereas the CD2 promoter [46] was used to generate the T cell specific Tl1a Tg mice [16]. The c-fms promoter vector [18] was used to generate myeloid specific Tl1a Tg mice for this report whereas the CD11c promoter [47] was used for the recently published studies [16,17]. Differences in the phenotype of the transgenic mice using the different T cell promoter/enhancer element to drive transgene expression was observed previously in the T cell specific LIGHT Tg mice [19,48,49]. Another reason may lie in the mouse strain used. The Tl1a Tg mice used in this report were generated by direct microinjection into C57BL/6 pronuclei whereas the DC specific Tl1a Tg mice [17] was generated by microinjecting into FVB/N zygotes and subsequently backcrossed 4-5 generations to the C57BL/6 strain. Alternatively, these phenotypical differences may be due to differences in the gut microbiota between the animal housing facilities. Previous studies have demonstrated that genetically identical inbred mice strain raised in different facilities had different immune composition due to the different gut microbiota composition [50,51,52]. The potential immunomodulatory role of gut microbiota in the different Tl1a Tg mice highlights the previously described role of the TL1A-DR3 signaling pathway in bacteria recognition and microbial-host interactions [6,8,10,16].

In summary, we generated tissue specific, in vivo expression of Tl1a in T cells and APC and found that it lead to patchy small bowel inflammation and fibrosis in both the small intestine and the colon. Sustained expression of Tl1a leads to an activated immune phenotype, increased expression of gut homing molecules and Th1 (and possibly Th17 and Th2) responses. These novel Tl1a transgenic models of intestinal inflammation may be useful to study fibrotic response in inflammation and the pathogenesis of various disorders of immune dysregulation such as IBD.

Materials and Methods

Transgenic mice and genotype

Murine Tl1a cDNA (Open Biosystems, Clone ID 30740802) was digested by EcoRI/Smal and inserted into the EcoRI/Smal site of pIRE2-EGFP (Clontech). To generate lymphoid specific Tl1a transgenic mice, DNA fragment containing murine Tl1A-IRE2-EGFP was amplified by PCR using sense primer 5’-AATGGGGGGCGGGGGGCTCTGCTGAGAAGGGAT-CAG-3’ and antisense primer 5’-TTTACGGGCGGCGCCCT-TAAAGATAATGAGGTTGGG-3’, digested with Ascl, and cloned into Ascl site of plck.E2 (generous gift from T. Hettmann, The University of Chicago), which contains the proximal lck promoter, human growth hormone gene (polyadenylation site), and locus control region elements from the human CD2 gene to generate plasmid pLCK-Tl1a-IRE2-EGFP. The plck.E2 has been used to mediate T cell lineage specific expression [19]. The Tl1A-IRE2-EGFP fragment was sequenced to confirm that no mutations were generated during the cloning process. A 10-kb fragment was excised by NotI digest and used for microinjection into C57BL/6 pronuclei performed by Caliper Life Sciences. 8 independent murine lines containing genomic integration of the Tl1a transgene were identified by PCR using the following primers: 5’-GACCTAACAAAGGCTCCGTCTTGAGG-3’ and 5’-GCACAATCTCTCTGCTGTCCTTTGGA-3’. Only 2 of the 8 lines showed T-cell lineage specific expression of Tl1a Tg (95% of T cells and 34% of T cells). We used the founder transgenic (Tg) mouse line with Tl1a transgene expression in 95% of T cells and called it LCK-CD2-Tl1a-GFP Tg mouse.

To generate myeloid specific Tl1a Tg mice, a Xhol/EcoRI fragment was blunt ended by klenow (NEB) and cloned to blunt ended MluI site of c-fms promoter vector (generous gift from D. Underhill, Cedars-Sinai Medical Center). Tl1A-IRE2-EGFP fragment was sequenced to confirm that no mutations were generated during the cloning process. The c-fms promoter vector has been used previously to drive expression for cells of the mononuclear phagocyte lineage including macrophages and dendritic cells [18]. A 10-kb fragment was excised by XhoI/Clai digest and used for microinjection into C57BL/6 pronuclei performed by Caliper Life Sciences. 6 independent murine lines containing genomic integration of the Tl1a transgene were identified by PCR using the following primers: 5’-TTTG-GAAAGCTGATTTGAGGGTCCA-3’ and 5’-AGCTCCTCT- TGCCATCCTCTTGCT-3’. Two of the six lines showed myeloid specific expression of the transgene in approximately 70% of macrophages and dendritic cells, but one has higher GFP level than the other founder line. We used the founder Tg line with the higher GFP expression and called this line FMS-Tl1a-GFP Tg mouse.

All mice were maintained under specific pathogen-free conditions in the Animal Care Facility at Cedars-Sinai Medical Center. Littermate control mice were used for all experiments. This study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the NIH. Mice used in all experiments were specifically approved by Cedars-Sinai Medical Center Animal Care and Use Committee. Approved protocol 2269 was used for this study.

Disease activity index (DAI), macroscopic and histopathological analysis

Mice were weighed and inspected for diarrhea and rectal bleeding once a week for the first 2 months after weaning, then once a month thereafter. The DAI (combined score of weight loss, presence of blood in stool, and stool consistency) was determined according to a standard scoring system previously described [20].

Colon and small intestine were scored for macroscopic evidence of inflammation using the established classification [21]. Tissue was fixed in 10% neutral buffered formalin (Sigma). Samples were embedded, sectioned, and stained with hematoxylin and eosin by the Histology Core at Cedars-Sinai Medical Center. Masson Trichrome staining was performed as described previously [29].

Histopathological scores of colons and small intestine were assigned in a blinded manner by at least 2 trained pathologists (DQS and HWK) using previously established methods [21]. The histologic score is calculated from observation of at least 36 different fields of stained sections at 200X from 6 mice in each group. The numbers of goblet cells and Paneth cells were determined by examining at least 80 individual villi and crypts from each group.

Real-time PCR analysis

Total RNA was isolated as previously described [8]. Two micrograms of total RNA was used in each RT reaction using the
Cytokine expression was measured by flow cytometry and intracellular staining. Cells were stained with antibodies against murine CD4 (RM-5), CD44 (IM7), CD86 (GL1) [BD Biosciences], CCR9 [242503], CCR10 [248918] [R&D systems], CD8 [53-67], CD11c [N418], F4/80 [BM4] [Biolegend], and FoxP3 [JFK-165] [eBiosciences]. CD16/CD32 (clone 2.4G2) was used to block nonspecific FcR binding [eBiosciences]. Intracellular staining was performed with antibodies against IFN-γ, IL-17, IL-13, and IL-10 [eBioscience]. For determination of intracellular cytokine production, cells were incubated for 5 hours with 37°C with BD Leukocyte Activation Cocktail and BD GolgIPlug [BD Biosciences]. For FoxP3 staining, fixation and permeabilization buffers provided from the manufacturer [eBiosciences] were used. Data of the stained cells were acquired on a Cytomax flow cytometer and analyzed using FlowJo analysis software [Tree Star, Inc].

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