Animal Model

Induction of Colitis in Mice Deficient of Peyer’s Patches and Mesenteric Lymph Nodes Is Associated with Increased Disease Severity and Formation of Colonic Lymphoid Patches

Thomas W. Spahn,* Hermann Herbst,† Paul D. Rennert,‡ Norbert Lüering,* Christian Maaser,* Mathias Kraft,* Adriano Fontana,§ Howard L. Weiner,¶ Wolfram Domschke,* and Torsten Kucharzik*

From the Department of Medicine B* and the Institute of Pathology,† Münster University Hospital, Münster, Germany; Biogen Incorporated,‡ Cambridge, Massachusetts; the Center for Neurologic Diseases,¶ Brigham and Women’s Hospital and Harvard Medical School, Boston, Massachusetts; and the Department of Medicine,§ Section of Clinical Immunology, Zürich University Hospital, Zürich, Switzerland

Inflammatory bowel disease is associated with immune activation in Peyer’s patches and mucosal lymph nodes. The role of these organs in dextran sodium sulfate (DSS)-induced colitis was investigated. We used mice lacking Peyer’s patches and/or lymph nodes because of lymphotoxin-α gene deficiency or treatment in utero with lymphotoxin-α/receptor IgG and tumor necrosis factor-receptor-I (55)-IgG fusion proteins. Mice lacking Peyer’s patches and lymph nodes because of lymphotoxin-α deficiency or in utero fusion protein treatment developed more severe colitis than control mice as indicated by more severe intestinal shrinking, longer colonic ulcers, and higher histological disease scores. Oral DSS triggered the formation of colonic submucosal lymphoid patches in these mice and caused an increase in the number of submucosal lymphoid patches in mice treated in utero with the fusion proteins. Mice lacking Peyer’s patches only showed more submucosal lymphoid patches whereas intestinal length and histological disease score were similar to control mice. In conclusion, more severe DSS-induced colitis correlates with the loss of the mesenteric lymph nodes. However, neither the absence of Peyer’s patches nor the presence of colonic lymphoid patches were correlated with increased disease severity. (Am J Pathol 2002, 161:2273–2282)

The physiological intestinal immune response toward intraluminal antigens include IgA secretion and the induction of systemic immune hyporesponsiveness oral tolerance.1 Inflammatory bowel disease is associated with a breakdown of tolerance toward the resident intestinal flora2,3 and immune activation in the gut-associated lymphatic tissue (GALT). The GALT consists of Peyer’s patches (PPs) and mesenteric lymph nodes (MLNs) as organized intestinal lymphoid follicles. PPs are lymphoid follicles in the intestinal wall and contain M cells that can uptake particulate intraluminal antigens.4 Although the role of PPs and MLNs in the induction of intestinal immune responses5–7 and of oral immune tolerance has recently been investigated,8–10 little is known about their role in the induction of inflammatory bowel disease.

Lymphotoxin-α (LTα) and LTβ are members of the tumor necrosis factor (TNF) cytokine family. LTαβ is critical for the induction of secondary lymphoid organs and the development of the spleen.11–14 LTα−/− mice do not develop PPs or LNs and have a disrupted splenic architecture. LTβ is required for the development of PPs but not of MLNs as LTβ (LTβ−/−) gene-deficient mice lack PPs but can develop at least some MLNs.14,15 Gestational treatment of mice with lymphotoxin-β-receptor-IgG-fusion protein (LTβRlgG) or LTβRlgG and TNF-receptor-I(55)-IgG-fusion protein (TNFRlgG) inhibits the formation of PPs or of PPs and MLNs depending on the treatment regimen.15,16

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Address reprint requests to Thomas W. Spahn, M.D., Department of Medicine B, Albert Schweizer-Str. 33, D-48129 Münster, Germany. E-mail: spahnth@t-online.de.
Little is known about the role of GALT organs in the induction and course of experimental colitis. We therefore used mice made deficient of either PPs or PPs and MLNs by fusion protein treatment (PP-null/LN+; PP/LN-null) or LTα deficiency (LTα−/−) to study the differential role of PPs and MLNs in the induction of colitis.

Materials and Methods

Mice

129xB6 wild-type (wt) and lymphotoxin-α gene-deficient (LTα−/−) mice were purchased from the Jackson Laboratories (Bar Harbor, ME). All mice were kept under sterile conditions in microisolator cages in the animal facilities of the Münster University Department of Dermatology with unlimited access to food and water according to federal animal protection regulations (permit G5/99 and G78/2000).

Abrogation of PPs or PP and MLN Development

Female 129xB6 mice were daily checked for vaginal plugs. To abrogate development of PPs alone, mice were intravenously injected with 200 μg of lymphotoxin-β-receptor IgG fusion protein (LTβRIgG; Biogen, Cambridge, MA) on days 16 and 18 after conception. Suppression of PP and MLN development was induced by intravenous injection of 100 μg of LTβRIgG and 100 μg of TNF-receptor-(l55kD)-IgG-fusion protein (TNFRIgG, Biogen) in pregnant mice on gestational days 11, 13, 15, and 17. Both regimens included treatment of the progeny with an intraperitoneal injection of 20 μg of LTβRIgG within 24 hours after birth. As treatment with human IgG does not induce changes in lymphoid organ development, we used age- and sex-matched 129xB6 mice as controls to progeny of LTβRIgG-treated animals. Mice were individually checked for the absence of the respective lymphatic organs by gross examination (LN) or soaking the intestines in 10% (v/v) acetic acid solution. Only mice devoid of PPs or PPs and LN, respectively, were included in the analysis. Progeny of treated mice were 7 to 10 weeks of age at the time of the experiment.

Induction of Colitis

Colitis was induced in all groups by the addition of dextran sodium sulfate (DSS) [molecular weight, 40,000; 4% (w/v); ICN, Biochemicals, Eschwege, Germany] to the drinking water. The mean DSS/water consumption was recorded. Mice were treated for 7 days with DSS or normal drinking water (NDW). Body weight was assessed before and after 7 days of oral DSS.

MPO Assay

This assay was performed as previously described.

Assessment of Colitis

All mice were euthanized on day 7 after colitis induction. The entire colon was removed and the length was recorded.

Histology

To assess the distribution of inflammatory changes in the colon the organ was cut into three sections of one-third of the total colon length and attached to a cork board before fixation. Tissues were subsequently fixed in 2% (v/v) paraformaldehyde solution and embedded in paraffin and subsequently sectioned. Hematoxylin and eosin (H&E) staining was used for general assessment of intestinal inflammation. Gomori’s silver staining of reticular fibers was performed as described.

Immunohistochemistry

Tissues were snap-frozen in OCT compound (Miles, Elkhart, IL). Frozen serial sections were prepared and incubated as described previously. Shortly, sections were fixed with acetone, washed, and preblocked with 5 μg/ml of anti-CD16/CD32 Fc block (Pharmingen, B&D, Heidelberg, Germany) in Tris-buffered saline with 0.25% bovine serum albumin, 0.05% Tween 20, and 10% heat-aggregated rabbit serum. Cells were stained with 5 μg/ml of biotinylated anti-B220 monoclonal antibody RA3–6B2 (Pharmingen) in the same buffer, followed by 10 μg/ml of fluorescein conjugated-neutrallite (fluorescein isothiocyanate-avidin; Southern Biotechnology Associates, Birmingham, AL). Staining of CD4 and CD8 expression was performed in a similar manner using PE-labeled monoclonal antibody L3T4–6B2 (Pharmingen) and biotin labeled antibody Ly-2 (CD8, clone 53-6.7). Staining of sections for MAdCAM-1 (Pharmingen) expression used monoclonal antibody MECA 367 (Pharmingen) followed by fluorescein isothiocyanate-conjugated goat anti-rat IgG (Pharmingen). Sections were visualized by a fluorescence microscope and photographed (Leica Microscan, Wetzlar, Germany) and photos were subsequently scanned for digital image processing.

Histological Scoring of Disease Severity

Tissue sections were examined by a gastrointestinal pathologist (HH) in a blinded manner. Each section was scored for severity and extent of ulceration, and the tissue thickness from the muscularis propria to the luminal border was determined. Lesion severity was graded using a modification of a previously defined scoring system with a scale of 0 to 4: 0; normal; 1, minimal; 2, mild; 3, moderate; 4, severe. Minimal lesions contained small, focal, or widely dispersed areas of inflammation and/or fibrosis above the muscularis mucosae. Mild lesions were multifocal or locally extensive and contained inflammation or fibrosis extending into the submucosa. Moderate lesions consisted of multifocal lesions with ulcers consuming <10% of the assessed mucosal surface. Severe
was defined similar to moderate lesions and/or ulcers consuming >10% of the assessed mucosal surface.

Percent length of colon ulcers was assessed as follows: the length of individual ulcers on longitudinal colon sections was measured, and the sum of all individual ulcers was multiplied \( \times \) 100 and divided by the length of the colonic surface on the respective slide of the histological section.

**Statistical Analysis**

Differences between body weights, bowel lengths, and number of intestinal ulcers were compared using Mann-Whitney U statistics supported by InStat software for Macintosh computers.

**Results**

**Colitis in LT\(\alpha\)−/− Mice and in Mice Deficient of PPs and MLNs after Gestational Fusion Protein Treatment**

We used LT\(\alpha\)−/− mice, which are congenitally devoid of secondary lymphoid tissues, to study the role of PPs and MLNs in the course of DSS colitis. To further assess the role of PPs and PPs and MLNs in the regulation of experimental colitis in a manner separable from the LT\(\alpha\) deficiency we also induced acute DSS colitis in mice made deficient of PPs and MLNs by treatment with LTβR/βG and TNFRI/RI G (PP/LN-null mice). Combined LTβR/βG and TNFRI/RI G treatment in mice from gestational day 11 through gestational day 17 inhibits PP and MLN formation without causing a life-long LT\(\alpha\) and LT\(\beta\) cytokine defect.

Acute colitis was more severe in LT\(\alpha\)−/− mice than in wt mice. The average disease score was higher in mice deficient of the LT\(\alpha\) gene and colonic ulcers covered <25% of the assessed mucosal surface (Figure 1, A and B). There was more severe shrinking of the inflamed colon than in wt mice (Figure 1C). To assess if the increased inflammatory activity was associated with an influx of neutrophil granulocytes, we also assessed colonic myeloperoxidase activity using the myeloperoxidase assay. However, myeloperoxidase activity was similar in diseased LT\(\alpha\)−/− and wt mice (data not shown). In addition, body weight during the course of colitis did not differ between wt and LT\(\alpha\)−/− mice.

Colitis was also more severe in PP/LN-null mice than in wt mice, as indicated by a higher average disease score (Figure 2A). Colonic ulcers covered up to 12% of the mucosal surface of PP/LN-null mice with colitis as compared to 3% in mice with PPs and MLNs (Figure 2B). Colitis in the absence of PPs and LNs was associated with more severe shrinking of the inflamed colon (Figure 2C) and more weight loss (Figure 2D). Leder staining of inflamed colon tissue from PP/LN-null mice for granulocytes did not show evidence of granulocyte infiltration in submucosal follicles (data not shown). In addition, MPO activity was similar in colonic tissue from PP/LN-null and wt mice undergoing colitis (data not shown).

As shown in Figure 3A, we could not detect any lymphoid follicles in the colon of LT\(\alpha\)−/− mice treated with NDW. Oral DSS induced the formation of lymphoid tissue in the absence of LT\(\alpha\) (Figure 3A). We found a higher number of lymphoid patches in the colon of LT\(\alpha\)−/− mice than in diseased wt mice.

Fusion protein treatment in utero did not suppress the formation of lymphoid patches in the colon. There were similar numbers of submucosal lymphoid follicles in untreated control and PP-null/LN-null mice (Figure 3B). As shown in Figure 3B, induction of colitis was associated with a dramatic increase of the number of submucosal lymphoid follicles in mice without GALT whereas the number of lymphoid follicles remained unchanged in control mice.
not clearly distinct from B-cell areas of the patch (Figure 5, G and H). We could detect only very few scattered CD4+ T cells in colonic follicles of untreated wt mice whereas CD4+ cell areas could be detected in cervical lymph nodes (CLNs) of wt mice (data not shown). We could also detect scattered CD4+ T cells in colonic lymphoid patches of diseased wt, LTα−/−, and PP/LN-null mice (not shown). We detected similar expression of the high endothelial venule marker MadCAM-1, which is characteristic for mucosal lymph nodes, in lymphoid follicles of wt LTα−/− and PP/LN-null mice (not shown).

To further characterize the architecture of inflammatory colonic aggregates in wt, LTα−/−, and PP/LN-null mice, we performed silver staining of reticular fibers. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates. Although organized lymphoid tissues contain a reticular fiber connective tissue structure, this is not expected in nonorganized leukocyte aggregates.
Figure 4. Microphotographs of representative sections of colon from wt (A, B), LTα−/− (C–E), and PP/LN-null (F–H) mice after oral treatment with NDW [wt (A), LTα−/− (C), PP/LN-null (F)] or 4% DSS in the drinking water for 7 days [wt (B), LTα−/− (D and E), PP/LN-null (G and H)]. Large arrows indicate lymphoid patches, small arrows indicate limits of epithelial ulcers. Areas of edema are indicated by asterisks. Original magnifications: ×5 (A–G); ×10 (H).
Figure 5. Immunohistochemical staining of colonic lymphoid aggregates from untreated wt mice (A and B), untreated PP/LN-null mice (C and D), PP/LN-null mice undergoing colitis (E and F), and LTα−/− mice undergoing colitis (G and H). **Left**: Staining for B220; **right**, staining for CD8. Original magnifications, ×10.
derive from the blood or expand from resident lamina propria cells. Colonic patches have been described in both mouse and human. To distinguish the organized colonic patches from inflammatory leukocyte aggregates we also performed staining for reticular fibers. We found similar reticular staining in both colonic lymphoid follicles and patches of wt mice, PP/LN-null, and LTα−/− mice undergoing colitis suggesting that the inflammatory lymphoid patches observed in our study have similar connective tissue structures as organized lymphoid tissues. Recently, cryptopatches have been described in the murine intestine, which give rise to T-cell

Figure 6. Gomori’s silver staining of colonic lymphoid patches of wt (A, B), LTα−/− (C, D), and PP/LN-null mice (E, F) undergoing colitis. Original magnifications: ×20 (A, C, E), ×40 (B, D, F).
receptor (TCR) \(\alpha/\beta^+\) and TCR\(\gamma/\delta^+\) intraepithelial lymphocytes.\(^{23,24}\)

We obtained very similar data results from studies using LT\(\alpha-/\) mice, which also lack PPs and MLNs. Using LT\(\alpha-/\) mice we again observed significantly longer colonic ulcers as well as higher disease scores as compared to diseased wt mice. We can therefore conclude that the absence of organized intestinal lymphoid tissue at the onset of disease promotes the formation of colonic ulcers after oral DSS and the de novo formation of lymphoid patches triggered by inflammation. Thus the presence of organized intestinal lymphoid tissue at the time of disease induction attenuates the severity of colitis whereas de novo-formed lymphoid tissue is associated with colitis induction. It is of note that weight loss was more severe in PP-null/LN-null mice undergoing colitis than in diseased LT\(\alpha-/\) mice. This observation suggests that the proinflammatory cytokine LT\(\alpha\)-R IgG might contribute to the wasting observed in DSS colitis.

Although we could not detect any lymphoid follicles in LT\(\alpha-/\) mice when treated with NDW, after oral DSS treatment we observed multiple colonic lymphoid patches in LT\(\alpha-/\) mice. It seems likely that the inability to detect the submucosal colonic lymphoid follicles in LT\(\alpha-/\) mice is because of a lack of a stimulus for cell migration to sites of lymph node development in the absence of LT\(\alpha\), which is partially reversed on inflammatory challenge.

This observation, and the observation that colonic follicles develop in wt mice treated with LT\(\beta\)-R-IgG and TNFR55-IgG \textit{in utero}, suggests that there is a lymphotoxin-independent pathway of secondary lymphoid tissue formation in the intestine. It is of note that this tissue consists primarily of B cells. Thus, B-cell follicle formation might be regulated in a LT\(\alpha\)-LT\(\beta\)-R interaction-independent manner. Hamada and colleagues\(^{26}\) have recently described follicle-like structures in the small intestine that mainly consist of B cells. These lymphoid aggregates were named “isolated lymphoid follicles” (ILFs). The lymphoid patches observed in our study resemble ILFs and might be a counterpart to them in the large intestine. The presence of MLNs was initially reported in a small percentage of LT\(\alpha-/\) mice.\(^{11,12}\) suggesting other signals could compensate for intestinal lymphoid tissue formation. Similar plasticity in mucosal LNs has been described in LT\(\beta-/\) mice.\(^{13}\) In the example of mucosal LN formation the plasticity in reliance of LN development on LT\(\beta\) was correlated with the immune status of the mother during gestation, that is, the LT\(\beta\)-independent pathway of mucosal LN development was made manifest after a colony was established from steriley rederived progeny. These observations on the development of various components of GALT suggest that a soluble factor produced by the mother can influence the development of lymphoid organs in the embryo. Although this novel observation makes evolutionary sense (the progeny are born equipped to deal with the immediate environment), and the influence of pathogens on GALT development is well appreciated\(^{27}\) this finding is nonetheless unexpected and in need of further study. At this time the precise mechanism and source of newly detectable colonic patches in LT\(\alpha-/\) mice that were treated with DSS remains unclear.

Mice without PPs but with MLNs also showed an increased number of colonic lymphoid patches but no differences in the loss of body weight or severity of intestinal shrinking. Thus, abrogation of LT\(\alpha\) formation alone is sufficient to induce the migration of lymphoid cells into the inflamed intestine. Thus, induction of lymphoid aggregates in the inflamed intestine after oral DSS is not necessarily associated with more severe disease as indicated by longer intestinal ulcers and increased weight loss.

While our manuscript was in preparation, Dohi and colleagues\(^{28}\) reported on the course of TNBS-induced colitis in mice made deficient of PPs by gestational fusion protein treatment. Using BALB/c mice, as compared to 129xB6 mice used in our study, they observed no organized lymphoid follicles in the colon after gestational fusion protein treatment. Furthermore, colitis induced by TNBS did not induce colonic lymphoid follicles. We suspect the differences in the mouse strains, disease models, or maternal environment are accountable for the differences observed in colonic lymphoid tissue and disease course.

In addition to LT\(\beta\) LIGHT can also bind to the LT\(\beta\)-R. Although the interaction between LIGHT and LT\(\beta\)R would be inhibited by continuous LT\(\beta\)R-IgG treatment during disease, as administered in the study of Dohi and colleagues,\(^{28}\) LIGHT could bind to the LT\(\beta\)-R in our PP-null/LN-null and LT\(\alpha-/\) mice as gestationally administered LT\(\beta\)-R-IgG is hardly detectable in the serum of treated pregnant mice at the age of 6 to 8 weeks\(^{17}\) when our mice were used for experiments. In addition, the \(rta\)-gene deficiency does not interfere with LIGHT binding to the LT\(\beta\)R.

An important similarity also emerges from the study of Dohi and colleagues\(^{28}\) and our studies. The disease that develops in response to oral DSS feeding or TNBS enema is much more severe in the absence of the normal array of mucosal secondary lymphoid organs such as PPs and draining mucosal LNs. It is therefore likely that
populations of cells normally present in PPs and mucosal LNs play a role in modulating the response to intestinal inflammation. One important candidate cell to fulfill this role are dendritic cells, whose complexity and function in PPs and LNs are only recently becoming well understood.29 For example, such dendritic cells have recently been shown to be sources of interleukin-10, a cytokine known to influence the outcome of gut inflammation.30

Despite the acute course that DSS-induced colitis takes in mice there is good evidence in the literature that the adaptive immune system can indeed regulate the disease. DSS-pulsed macrophages stimulate T cells from colitis mice for secretion of T helper 1 cytokines31 in vitro. Inflammatory infiltrates consist of plasma cells, neutrophils, and mononuclear cells.32 Severity of DSS-induced colitis can be modulated by cyclosporin-induced suppression of T-cell activation33 and intravenous immunoglobulin.34

There is evidence for preferential immune activation in regional lymphoid follicles in animal models of inflammatory bowel disease. In indomethacin-induced colitis, there is proliferation and apoptosis of M cells in the inflamed intestine.35 In TCR-α/−/− mice, there was more proliferation and a higher level of autoantibody production in appendix lymphoid follicle tissue than in PP tissue during colitis.36 In a Th-2 model of inflammatory bowel disease induced in interferon-γ−/− mice, colitis is associated with colonic patch hypertrophy.21 Lymphoid hyperplasia has been reported as a feature of chronic DSS-induced colitis.37 Secondary lymphoid follicles are also observed in patients with Crohn’s disease. Our results suggest an important role of the GALT in maintenance of mucosal integrity and control of mucosal inflammation. Indeed, development of colonic follicles during colitis may be an attempt to control intestinal inflammation.

It is reasonable to speculate that the severe colitis that develops in the absence of PPs and MLNs may be in part related to a failure of tolerance induction because of the loss of normally present regulatory cells in these organs such as dendritic cell populations mentioned above. Oral feeding of sonicates from anaerobic intestinal bacteria and of haptenized colonic protein down-modulates colitis in mice.38,39 Thus intraluminal and intestinal wall antigens have the capacity to induce tolerance toward inflammatory intestinal immune responses. Our previous study and those of other groups have implicated both PPs and MLNs in the development of intestinal oral tolerance8,9 with MLNs playing a pivotal role. There is evidence showing that lymph nodes draining mucosal sites are critical for the induction of immune tolerance. We have recently demonstrated that induction of MLN formation in the LTα−/− model restores the capacity to induce tolerance to oral antigen in the absence of PPs.10 Wolvers and colleagues40 have shown that CLNs are critical for the induction of nasal tolerance. Surgical removal of CLNs abrogated the capacity to induce tolerance by the nasal route and transplantation of CLNs but not of other peripheral LNs to mice without CLNs restored the capacity to induce nasal tolerance.

In this study we observed an increase in the disease score in mice deficient of PPs and MLNs whereas disease activity in mice lacking PPs only was comparable to control mice. We therefore assume that the lack of the capacity to induce oral tolerance in the absence of MLNs8 may be associated with an exacerbation of intestinal inflammation.

Recently, it has been reported that treatment of adult mice with LTβR-IgG could attenuate experimental colitis in mice.41 This observation is not in contradiction to our study as we used LTβR-IgG fusion protein for gestational inhibition of lymphotxin-receptor interaction to ablate the formation of secondary lymphoid organs, rather than treating them once formed. The LTαβ/LTβR pathway has varied and distinct biological roles to play during development and in the normal physiology of the adult.42 Because LTβR-IgG treatment of adult wt mice seems to influence the development of a variety of immune diseases, including inflammatory bowel disease, it might be anticipated that treatment of adult wt mice would have efficacy in the DSS model as well.

In summary we report more severe intestinal inflammation in the absence of MLNs whereas the absence of PPs alone did not affect disease severity. These observations suggest a regulatory function for MLNs but not for PPs in intestinal inflammation.

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