Intestinal epithelial cell-derived interleukin-7: a mechanism for the alteration of intraepithelial lymphocytes in a mouse model of total parenteral nutrition

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Yang H, Sun X, Haxhija EQ, Teitelbaum DH. Intestinal epithelial cell-derived interleukin-7: a mechanism for the alteration of intraepithelial lymphocytes in a mouse model of total parenteral nutrition. Am J Physiol Gastrointest Liver Physiol 292: G84–G91, 2007; doi:10.1152/ajpgi.00192.2006.—Total parenteral nutrition (TPN), with the absence of enteral nutrition, results in profound changes to both intestinal epithelial cells (EC) as well as the adjacent intraepithelial lymphocyte (IEL) population. Intestinal EC are a rich source of IL-7, a critical factor to support the maintenance of several lymphoid tissues, and TPN results in marked EC changes. On this basis, we hypothesized that TPN would diminish EC-derived IL-7 expression and that this would contribute to the observed changes in the IEL population. Mice received enteral food and intravenous crystalloid solution (control group) or TPN. TPN administration significantly decreased EC-derived IL-7 expression, along with significant changes in IEL phenotype; decreased IEL proliferation; and resulted in a marked decrease in IEL numbers. To better determine the relevance of TPN-related changes in IL-7, TPN mice supplemented with exogenous IL-7 or mice allowed ad libitum feeding and treated with exogenous administration of anti-IL-7 receptor (IL-7R) antibody were also studied. Exogenous IL-7 administration in TPN mice significantly attenuated TPN-associated IEL changes, whereas blocking IL-7R in normal mice resulted in several similar changes in IEL to those observed with TPN. These findings suggest that a decrease in EC-derived IL-7 expression may be a contributing mechanism to account for the observed TPN-associated IEL changes.

INTERLEUKIN 7 (IL-7) is a member of the gamma chain-dependent (γc) family of cytokines, which share a common receptor γc component and include IL-2, IL-7, IL-9, and IL-15. These cytokines influence T cell development and function (12, 36). IL-7 has a major effect on the phenotype and development of thymocytes as well as other lymphoid tissues (26, 35). In vivo only IL-7, and not other γc family members (e.g., IL-4 or IL-15), was found to be essential for homeostatic proliferation of naive peripheral T cells (34). IL-7 is produced by thymic and intestinal epithelial cells (EC) (9, 35, 37), and in turn IL-7 receptors (IL-7R) have been detected on the surface of thymocytes and intestinal intraepithelial lymphocytes (IEL) (32, 35). Additionally, IL-7 receptors have been identified on peripheral T cells, B lineage cells, and colonic lamina propria lymphocytes (32, 35). Administration of IL-7 has been demonstrated to enhance both peripheral T cells and IEL numbers and increase peripheral T cell and IEL function (9, 46).

Interactions between mucosal lymphocytes and intestinal EC are thought to be crucial for maintaining mucosal immunity. Several studies have indicated that EC may play an important role in mucosal immune responses by helping to regulate IEL phenotype and function (11, 35). IEL are a distinct population of T lymphocytes that reside above the basement membrane and lie between EC. IEL act as the initial lymphoid defense layer against intraluminal foreign antigens (7) and may be of critical importance for proper functioning of the mucosal immune system (35). Previous studies by our group have shown that IEL play an important role in the maintenance of the gut barrier function and support intestinal EC growth (42–45, 47). There is an average of 10–20 IEL per 100 villi EC in human small intestine (11). This interrelation is well demonstrated with IL-7. IL-7 knockout and IL-7R knockout mice show distinct declines in absolute numbers of thymocytes and in the intestine, IEL (24). In an IEL culture model, IL-7-supplemented media significantly prevented the spontaneous apoptosis of IEL by decreasing caspase activity and preventing the decline in Bcl-2 expression (40).

It is estimated that total parenteral nutrition (TPN), or the intravenous administration of nutrition, is essential for the sustenance of patients unable to tolerate enteral nutrition, and over 550,000 patients receive TPN in the United States on a yearly basis (1). Despite this, TPN administration results in a number of immunological problems, including an increase in systemic sepsis and perioperative infections. Many of these infections may well be due to aberrancies in the mucosal immune defense system, including marked changes in the number and function of mucosal lymphocytes, including IEL (16, 17, 42, 49). It is unknown what mechanism(s) lead to these TPN-associated IEL changes. Recently, we have shown that IL-7 administration in healthy wild-type mice led to significant changes in IEL phenotype and function, including an increase in the CD80β+ and mature (CD44+) IEL subpopulations. IL-7 administration also significantly changed IEL-derived cytokine expression (46). Furthermore, we also demonstrated a close physical communication between EC-derived IL-7 and IEL in a mouse model (46). On the basis of these findings, we hypothesized that TPN-induced mucosal changes will lead to a decline in EC-derived IL-7 expression and that this decline would be responsible for changes in the neighboring IEL phenotype and function. Additionally, we hypothesized that exogenous administration of IL-7 would prevent many of the observed TPN-induced changes to the IEL.

METHODS

Animals

Studies reported here conformed to the guidelines for the care and use of laboratory animals established by the University Committee on the Use of Laboratory Animals (1). All experiments were approved by the University Committee on the Use of Laboratory Animals. Animals were housed in separate rooms with specific pathogen-free conditions and allowed ad libitum access to food and water. All surgeries were performed on anesthetized animals. Animals were killed by cervical dislocation.
Use and Care of Animals at the University of Michigan, and protocols were approved by that committee (UCUCA no. 7703). Male, 2-mo-old (weighing 24–26 g), specific pathogen-free, adult C57BL/6J mice (Jackson Laboratories, Bar Harbor, ME) were maintained in temperature-, humidity-, and light-controlled conditions. During administration of intravenous solutions, mice were housed in metabolic cages.

TPN Model

Administration of TPN was performed as previously described (17). Mice were infused with a crystalloid solution at 4 ml/day. After 24 h, mice were randomized into two groups (n = 6 per group). The control group received the same intravenous saline solution at 7 ml/24 h, in addition to standard laboratory mouse chow and water ad libitum. The TPN groups received an intravenous TPN solution at 7 ml/24 h. The TPN solution has been described in detail previously (17) and contained a balanced mixture of amino acids, lipids, and dextrose in addition to electrolytes, trace elements, and vitamins. Caloric delivery was based on estimates of caloric intake by the control group and from previous investigators (23), so that caloric delivery was essentially the same in both groups. All animals were killed at 7 days by use of CO2.

Exogenous IL-7 Administration

To further assess the role of IL-7 during TPN administration, IL-7 (Recombinant IL-7, R&D Systems) was given to a separate group of TPN mice (TPN+IL-7 group). Mice received an intravenous injection (500 ng twice daily) (3) starting with the first day of TPN, and treatment continued for 7 days.

IL-7 In Vivo Blockade

To assess the role of a decline in IL-7 on the observed altered IEL phenotype and function with TPN, in vivo blockade of IL-7 action was performed in another group of mice. This group was permitted ad libitum enteral nutrition (IL-7R blocking group) and received anti-IL-7R antibody via intraperitoneal injection, as previously described (32). The control group received a nonspecific isotype IgG at the same dose. The IEL has a slow rate of lymphocyte turnover. Therefore, to allow for a complete assessment of the effect of removing IL-7 on the IEL, anti-IL-7R antibody was administered at a dose of 1 mg of IgG per injection once every 3 days for a total of 4 wk (21). All IL-7R-dependent processes have been shown to be blocked by this anti-IL-7R antibody, which was generated from the hybridoma line A7R34 (kindly provided by Dr. Nishikawa Shinichi, Kyoto University, Kyoto, Japan). Ascites fluid was generated in nude mice, and anti-IL-7R antibody was purified by using a Prosep-G Kit (Millipore).

IEL Isolation and Purification

IEL isolation from small intestine. Small bowel IEL and ECs were isolated as previously described (17). Briefly, the small bowel was placed in tissue culture medium (RPMI 1640 with 10% FCS; Life Technologies). Mesenteric fat and Peyers’s patches were removed. The intestine was then opened longitudinally and agitated to remove mucus and fecal material. The intestine was then cut into 5-mm pieces, washed three times in an IEL extraction buffer (1 mM EDTA, 1 mM DTT in PBS), and incubated in the same buffer with continuous brisk stirring at 37°C for 20 min. The supernatant was then filtered rapidly through a glass wool column. Magnetic beads conjugated with antibody to CD45 (lymphocyte specific) were used to remove non-lymphoid cells (BioMag SelectaPure Anti-Mouse CD 45R antibody particles, Polyscience, Warrington, PA). Cells bound to beads were considered purified IEL; the EC remained in the supernatant. Flow cytometry confirmed purity of IEL, which was >99%, based on a control sample stained with anti-CD45 antibody.

Purification of IEL subpopulations. IEL subpopulations were purified by flow cytometry with cell sorting using an EPICS Elite (Coulter) cytometer, as described previously (42). The following antibodies were used for isolation of specific IEL subpopulations: CD4, CD8β, CD8α, TCR-αβ, and TCR-γδ (PharMingen). Isotype-matched, irrelevant antibodies were used as a negative control.

Reverse Transcriptase Polymerase Chain Reaction

Isolation of total RNA. A guanidinium isothiocyanate-chloroform extraction method was used. Total RNA from isolated EC or IEL, or subtype IEL, was extracted using Trizol reagent (Life Technologies) according to the manufacturer’s directions.

Reverse transcriptase and polymerase chain reaction. Poly-A-tailed mRNA was reversed transcribed into complementary DNA by adding total cellular RNA to the following reaction mixture: PCR nucleotide mix (Invitrogen), M-MLV reverse transcriptase (Invitrogen), Oligo(dT)12–18 primer (Invitrogen), and RNAse inhibitor (40 U/μl, Roche Diagnostics). Diethyl pyrocarbonate-treated H2O was added to yield the appropriate final concentration. Samples were incubated at 40°C for 70 min, and the reaction was then stopped by incubating at 95°C for 3 min. PCR reaction was run using thermal cycler settings that were optimized to ensure that products were in the logarithmic phase of production. PCR products were run out on a 2% agarose gel. Quantification of cDNA product was completed using a Kodak 1D image quantification software (Kodak), and target PCR products were compared by normalizing each sample to the production of β-actin.

Real-time PCR. A Smart Cycler thermal cycler (Cepheid, Sunnyvale, CA) was used for quantification of IL-7 mRNA expression. For this, a mastermix of the reaction components was prepared as described previously (42). The following experimental protocol was used: denaturation program (95°C for 2 min), amplification, and quantification program repeated 43 times (95°C for 15 s, 66°C for 15 s, and 72°C for 40 s). Specificity of real-time PCR products was documented with gel electrophoresis. Additionally, cDNA was extracted by using a centrifugal filter device (Millipore), and sequencing of this product showed that it matched the IL-7 GenBank NM_008371 mRNA sequence. The number of cDNA copies was then calculated by using the standard gene sequence size of each gene ampiclon. Serial dilutions of the amplified gene at known concentrations were tested to make a standard curve. Standard curve extrapolation of gene copy number was performed for the IL-7 gene as well as for β-actin. Normalization of values was performed by dividing the number of copies of the IL-7 gene by the number of copies of β-actin multiplied by 10^-5.

Immunoblot Analysis for EC-Derived IL-7 Expression

Briefly, isolated EC were homogenized on ice in lysis buffer (39). Protein determination was performed using a Micro BCA protein assay kit (Pierce, Rockford, IL). Approximately 60 μg of total protein in loading buffer were loaded per lane in a SDS-polyacrylamide gel (13%) and separated using electrophoresis. Proteins were then transferred to a polyvinylidene difluoride membrane (Bio-Rad Laboratories). The membrane was blocked with blocking solution (Zymed Laboratories) and probed with goat anti-mouse biotinylated-IL-7 antibody (R&D Systems) (0.15 μg/ml in blocking solution) for 1 h. Bound antibodies were exposed to a streptavidin-horseradish peroxidase conjugate (1:10,000, Zymed Laboratories; Invitrogen, Carlsbad, CA), detected on X-ray film. Within a range of 37 kDa no additional bands were detected. A further confirmation of the antibody specificity was performed by the use of recombinant mouse IL-7 (R&D Systems) as a positive control. Blots were then stripped and reprobed with monoclonal mouse anti β-actin antibody (1:8,000 in blocking solution; Sigma, St. Louis, MO) to confirm equal loading of protein. The peroxidase-conjugated second antibody was goat anti-mouse IgG (1:8,000 in blocking solution; Invitrogen). Quantification of results was performed using Kodak 1D image quantification software (Kodak). Thus, results of immunoblots are expressed as the relative expression of IL-7 to beta-actin expression.
proliferation and a rise in EC apoptosis (43, 48). We first assessed whether TPN was associated with changes in the expression of intestinal EC-derived IL-7. Total parenteral nutrition (TPN) administration led to a significant decrease in the rate of EC proliferation compared with a mean of 2.86 ± 0.87% of IEL numbers when compared with controls (8.1 ± 1.7 vs. 5.2 ± 0.8% for control and TPN groups, respectively; number of cells × 10^6 per group, *P < 0.05*).

The change in the number of IEL after TPN administration was also studied. We found that TPN administration led to a significant decrease in IEL numbers when compared with controls (8.1 ± 1.7 vs. 5.2 ± 0.8 for control and TPN groups, respectively; number of cells × 10^6 per group, *P < 0.05*).

**Changes in Intestinal IEL After TPN Administration**

**Changes in IEL phenotype and number.** Several changes were identified in IEL surface phenotypic markers with TPN administration. These changes included a 2.5-fold decline in the CD8αβ⁺ IEL subpopulation (i.e., CD8αβ⁺ heterodimeric portion of the CD8⁺ cells) (Fig. 3). Additional changes included a marked decline in the maturity of IEL, as indicated by a decline in CD44⁺ IEL, and consisted of a 3.6-fold decline in CD8⁺/CD44⁺ IEL and a 3.2-fold decline in CD4⁺/CD44⁺ IEL (Fig. 4). The CD4⁺ population was known to be very responsive to exogenous stimulation (16). After 7 days of TPN administration, both the CD4⁺/CD8⁻ as well as CD4⁺/CD8⁺ IEL subpopulations decreased by 87 and 80%, respectively, compared with controls (Fig. 5).

Flow Cytometric Analysis

**IEL phenotype analysis.** IEL phenotype was studied with flow cytometry. To examine the IEL subsets, the following monoclonal antibodies (BD PharMingen, San Diego, CA) were used: anti-CD4, CD8α, CD8β, αβ-TCR, and γδ-TCR, as well as CD44. Anti-mouse IL-7 receptor alpha antibody (Clone A7R34) was purchased from eBioscience (San Diego, CA). Acquisition and analysis were performed on a FACSCalibur (Becton-Dickinson) using CellQuest software (Becton-Dickinson). The IEL population was gated on the basis of forward and side scatter characteristics. Quantification of each IEL subpopulation was based on its percentage of the gated IEL population.

**IEL cell cycle analysis.** The modified protocol of Geiselhart et al. (9) was used. Briefly, purified IEL were permeabilized by resuspension in saponin buffer for 10 min, followed by centrifugation at 1,500 rpm for 5 min at 4°C. The supernatant was decanted and the pellet was resuspended in a saponin buffer containing propidium iodide (Sigma) and RNase (Roche Diagnostics), followed by incubation for 15 min at 4°C. Labeled cells were analyzed with flow cytometry, and CellQuest software was used for cell cycle analysis. Cells in either the S phase or G2/M were considered proliferating and were expressed as a percent of all gated IEL.

Data Analysis

All data are expressed as means ± SD. Cytokine expression and flow cytometric results were analyzed by ANOVA, and a Bonferroni post hoc analysis was used to detect differences between groups. Statistical significance was defined as *P < 0.05*.

**RESULTS**

**Changes in EC-Derived IL-7 Expression After TPN**

It has been shown that IL-7 is produced by intestinal EC (4, 9, 35, 37) and plays an important role in mucosal immune responses by regulation of growth and function of IEL (11, 35). Our laboratory has previously reported that TPN administration was associated with a significant decline in the rate of EC proliferation and a rise in EC apoptosis (43, 48). We first assessed whether TPN was associated with changes in the expression of intestinal EC-derived IL-7. Real-time PCR results showed that EC-derived IL-7 mRNA expression significantly decreased after TPN administration when compared with controls (Fig. 1). Western immunoblot studies of EC-derived IL-7 protein confirmed these changes. TPN administration led to a significant decrease in EC-derived IL-7 protein expression (Fig. 2).

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**Changes in IEL proliferation.** Because of the profound effect of TPN on IEL phenotypic changes and decline in the number of IEL, the nonstimulated percent of proliferating IEL (i.e., spontaneously proliferating lymphocytes) was determined. IEL proliferation declined to 1.74 ± 0.23% with TPN administration compared with a mean of 2.86 ± 0.33% in the control (enterally fed) group. Thus 7 days of TPN administration led to a 1.6-fold decline in IEL proliferation (Fig. 6).

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**Data Analysis**

All data are expressed as means ± SD. Cytokine expression and flow cytometric results were analyzed by ANOVA, and a Bonferroni post hoc analysis was used to detect differences between groups. Statistical significance was defined as *P < 0.05*.
IL-7R expression. IL-7R expression has been detected on IEL and lamina propria lymphocytes (32, 35). To determine whether local EC-derived IL-7 production affects the neighboring IEL through the IL-7/IL-7R-dependent signaling pathway, the IL-7R expression of various IEL subpopulations was studied.

IL-7R expression on IEL subpopulations. IEL were sorted by flow cytometry according to the phenotypic cell markers: CD4, CD8α, CD8β, αβ-TCR, and γδ-TCR. As shown in Fig. 7, IL-7R mRNA expression was identified on CD4+ and CD8α−, as well as the CD8αβ− IEL subtypes; IL-7R was also identified in both αβ-TCR+ IEL and γδ-TCR+ IEL subpopulations. Flow cytometry was also used to confirm surface protein expression of IL-7R in IEL subpopulations. Results are expressed as both the percent of IL-7R cells for all gated IEL cells, as well as the percent of IL-7R+ cells for each IEL subpopulation. Similar to the mRNA analysis, IL-7R was observed in αβ-TCR+, γδ-TCR+, CD4+, as well as CD8− IEL populations (Fig. 8). CD4+ IEL had a higher number of IL-7R+ positive cells (2.2 ± 0.22% of all gated IEL or 42.3% of CD4+ IEL were IL-7R positive) compared with the CD8− IEL subtype (13.6 ± 2.4% of all gated IEL or 16.2% of CD8− IEL were IL-7R positive). Additionally, αβ-TCR+ IEL had a nearly twofold higher IL-7R expression compared with γδ-TCR+ IEL: 20.4 ± 3.8% of all gated IEL or 35.4% of αβ-TCR+ IEL were IL-7R positive, compared with 7.7 ± 4.7% of all gated IEL or 20.9% of γδ-TCR+ IEL found to be IL-7R positive (Fig. 8).

Changes in IEL IL-7R mRNA expression after TPN administration. Because IL-7 expression signals through IL-7R in the intestinal mucosa (41), we hypothesized that TPN-associated IEL changes may also affect IEL IL-7R expression. IEL IL-7R mRNA expression (normalized to beta-actin expression) sig-

Fig. 3. Representative flow cytometry results of gated intraepithelial lymphocyte (IEL) populations. Cell populations are expressed as the percentage of gated cells with CD8α and CD8β markers based on isotype-matched control antibodies. Data are expressed as the percent of total gated population of lymphocytes. TPN administration led to a loss of the CD8αβ+ IEL subset. IL-7 administration attenuated the loss of CD8αβ+ IEL associated with TPN administration (TPN+IL-7). Using anti-IL-7 receptor antibody (IL-7R blockade) led to a significant decrease in the CD8αβ+ IEL subset compared with controls.

Fig. 4. Distribution of the CD44+ IEL. CD44 was used as a marker of IEL maturation. TPN resulted in a significant decrease in the CD8−CD44+ IEL, P < 0.05 compared with control mice. IL-7 administration in TPN-treated mice attenuated the decrease in CD8−CD44+ and CD4−CD44+ IEL subsets compared with the TPN group (P < 0.05 compared with controls). Blocking IL-7R also resulted in a significant decrease in CD4+CD44+ and CD8−CD44+ subpopulations compared with controls (n = 6 each group, *P < 0.05).

Fig. 5. Distribution of IEL phenotypes. TPN administration led to a loss of CD4+CD8− IEL and CD4+CD8− subsets. IL-7 administration significantly attenuated the loss of CD4+CD8− and CD4+CD8+ IEL subpopulations associated with TPN administration. Blocking IL-7R with anti-IL-7R antibody resulted in a significant decrease in both CD4+CD8− and CD4+CD8+ IEL compared with control (n = 6 each group, *P < 0.05).

Fig. 6. TPN results in a decline in the proportion of IEL that are in active cell proliferation. IEL were surface labeled with fluorochrome-conjugated antibody to CD3 and then treated with propidium iodide for detection of cell cycle status using flow cytometric analysis. Histograms were generated by gating on IEL populations and displaying the cell cycle. TPN administration significantly decreased IEL proliferation compared with controls, P < 0.05. Administration of IL-7 to TPN mice attenuated this decline, and IL-7R antibody replicated the decline in IEL proliferation. Results are the percentage of IEL in a proliferative phase (as determined by cells in the S/G2/M cell cycle phases). Results are the mean of n = 6 mice in each group, *P < 0.05 compared with the control group.
significant decrease (0.48 ± 0.2 in TPN vs. 0.84 ± 0.02 in control) after TPN administration compared with controls (P < 0.05, based on n = 6 per group).

Changes in IEL with IL-7R blockade. To further confirm that a decline in EC-derived IL-7 accounts for the observed changes in the IEL with TPN administration, anti-IL-7R antibody was administered to enterally fed mice. Blockade of IL-7R resulted in several IEL phenotype changes that were similar to those observed with TPN administration. The CD8αβ+ IEL subpopulation was affected most profoundly, decreasing by 55% compared with controls (Fig. 3). IL-7R blockade also caused significant declines in the CD4+CD8−, CD4+CD8+ (Fig. 5), CD4+CD44+, as well as the CD8+CD44+ subsets compared with controls (Fig. 4). Coincident with these phenotypic changes, IL-7R blockade also resulted in a significant decline in total IEL numbers compared with the control group (P < 0.05, 4.2 ± 0.4 vs. 8.1 ± 1.7 × 10^6 IEL cells for the IL-7R blockade and control groups, respectively).

The effect of blocking IL-7R on IEL proliferation was also studied. Similar to mice that were given TPN, the basal proliferation rate decreased to 1.9% compared with 2.8% in the control group. Thus IL-7R blockade led to a 32% decline in the IEL proliferation rate compared with control mice (Fig. 6).

IEL Changes With IL-7 Administration to TPN Mice

We next investigated whether exogenous administration of IL-7 could attenuate the observed IEL changes associated with TPN administration. Exogenous IL-7 was given to TPN-treated mice starting on the first day of TPN administration and continued twice daily for 7 days. IL-7 administration significantly attenuated the loss of the CD8αβ+ IEL subpopulation observed with TPN (Fig. 3). The percent of CD8αβ+ IEL in the IL-7-treated TPN group was similar to the levels in the control (enterally fed) group and was significantly higher than values in the TPN group (P < 0.05). IL-7 administration in TPN mice also resulted in a significant increase in the percentage of CD44+ IEL (P < 0.05). The CD8−CD44+ IEL population was more than twofold higher in IL-7-treated mice compared with TPN mice (Fig. 4). IL-7 administration also attenuated the loss of CD4+CD44+ IEL after TPN administration (Fig. 4). CD4+CD8− subpopulations were also investigated: IL-7 administration significantly attenuated the loss of the CD4+CD8− and CD4+CD8− IEL subpopulations during TPN administration (Fig. 5). IL-7 administration to TPN mice significantly blunted the decline in IEL proliferation. IEL proliferation was 2.95 ± 0.39% in the IL-7 treated group and was not significantly different that the control group (2.44 ± 0.29%; P > 0.05; Fig. 6). Finally, IL-7 administration was also effective in preventing the observed decline in the number of IEL after TPN (P < 0.05). With IL-7 administration, the number of small intestinal IEL rose to 12.9 ± 3.4 × 10^6 per mouse, compared with 5.2 ± 0.8 × 10^6 in the TPN group.

DISCUSSION

In this study, we found that TPN administration led to a significant decrease in EC-derived IL-7 expression, as well as to significant changes in the IEL phenotype, and to a decline in total IEL numbers. Exogenous IL-7 administration was able to attenuate most of these TPN-associated IEL changes. In addition, IL-7R blockade in enterally fed mice led to IEL phenotypic changes that were similar to those seen after TPN. This supports the concept that the decline in EC-derived IL-7 expression with TPN administration may be an important mechanism to account for many of the observed changes to the IEL. The marked changes to the intestinal immune system with administration of TPN are a significant clinical problem. TPN administration in a number of clinical studies has been associated with an increased rate of infections and septic episodes (20, 27). It is also thought that the source of sepsis may be due to an underlying loss in the function of the intestinal mucosal immune system (16, 31, 38). Thus understanding potential
mechanisms responsible for these changes in the IEL may allow for improved strategies to prevent TPN-associated infections.

The predominant IEL subpopulation in mice is CD8$^+$CD4$^-$ (70–85% of the total IEL number) (10). IEL also have a large number of γδ-TCR$^+$ cells, and up to 50% of IEL are γδ-TCR$^+$, compared with less than 2% of peripheral blood lymphocytes, in mice (16). Additionally, IEL are believed to originate from both a thymic-independent (29) and a thymic-dependent source (10). Alterations in the luminal environment produce significant changes in the IEL (16). This study found that there is a significant decrease in the CD4$^+$CD8$^-$, CD4$^+$CD8$^+$ and CD4$^+$CD44$^+$, and CD8$^+$CD44$^+$ IEL subpopulations, as well as a loss of CD8αβ$^+$ IEL after TPN administration in mice. The CD4$^+$ IEL are known to be very responsive to exogenous stimulation, and their loss may explain the observed decrease in IEL proliferation with TPN (16). The reduction in CD44$^+$ cells suggests a shift to a less mature IEL. The precise etiology of these IEL changes is uncertain.

It is well appreciated that IL-7 can promote growth and differentiation of many T cell phenotypes (2), and this is supported by the wide range of IEL subpopulations that express IL-7R. IL-7$^+$ is also essential for early developmental processes such as the differentiation of pre-T cells into mature thymocytes (50). This latter function cannot be performed by any other known cytokine (50). In the absence of IL-7$, homo- static proliferation of naive T-cells is almost completely abolished, and the lifespan of naive T cells is greatly reduced (34).

In vivo, administration of IL-7 has been demonstrated to enhance peripheral T cell functional capacity and expand the peripheral T cell population (19). Geiselhart et al. (9) have reported that IL-7$^+$ administration alters the peripheral T cell CD4$^+$ to CD8$^+$ ratio and results in an increase in peripheral T cell numbers and altered function. Watanabe et al. (35) observed that exogenous IL-7$^+$ administered to mice resulted in a stimulation of lamina propria lymphocytes. Interestingly, in an IL-7 overexpression transgenic mouse model, IL-7R expression on IEL was found to be increased (36). Recently, we reported that IL-7$^+$ administration to healthy wild-type mice led to a significant increase in CD8αβ$^+$ IEL and to a significant relative decline in the percentage of the CD8αε$^+$ IEL subpopulation (46). IL-7$^+$ administration also significantly increased the percentage of CD8$^+$CD44$^+$ IEL and led to a significant change in the IEL cytokine expression function (46). These data suggest that IL-7 may be essential for the ongoing maintenance of the IEL in mature mice. The mechanisms by which the IEL phenotype changes with TPN are not known. TPN results in several phenotypic IEL changes that were quite similar to those observed in TPN mice. These changes include a loss of the CD8αβ$^+$, CD4$^+$CD44$^+$, and CD4$^+$ (including the CD4$^+$CD8$^+$ and CD4$^+$CD8$^-$) IEL subpopulations. The fact that blocking IL-7R$^+$ only partially replicated some of the changes observed in the IEL phenotype after TPN may indicate that other mechanisms are also involved in this process. Additionally, IL-6$^+$ numbers were lower in the IL-7$^+$ blockade group compared with the TPN group. This is most likely due to the fact that, although IL-7 expression is reduced with TPN administration, local levels of IL-7$^+$ were still present in the TPN group, whereas IL-7$^+$ blockade resulted in a much greater decline in this IL-7$^+$IL-7$^+$ pathway. The incomplete replication of the phenotype with receptor blockade may also be due to the fact that the turnover of the IEL population is slower than most lymphocyte populations (28). Nevertheless, exogenous IL-7 administration prevented the development of the majority of TPN-associated IEL phenotype changes and the decline in total IEL numbers, underscoring the importance of this cytokine in the development of TPN-induced changes to the mucosal immune system. A previous study of ours has shown that there is a close physical close communication between EC-derived IL-7$^+$ and the neighboring IEL by the colocalization of these cell populations (46). These data suggest that cell-to-cell interactions between EC and IEL, via IL-7, could be an important model of communication for the maintenance and activation of the IEL. The fact that the highest expression of IL-7$^+$ was found on CD4$^+$ and αβ-TCR$^+$ IEL may be quite important because there are significant functional differences between IEL subpopulations. Namely, the CD4$^+$ and αβ-TCR$^+$ IEL populations are known to have a much greater proliferative capacity compared with CD8$^+$ and γδ-TCR$^+$ IEL subpopulations (5, 18). These major differences in IEL proliferation observed by these investigators may well be explained by our finding of the highest expression of IL-7$^+$ on CD4$^+$ and αβ-TCR$^+$ IEL. The changes in IEL phenotype may play an important role in the EC function. We have previously shown that γδ-TCR$^+$ IEL-derived keratinocyte growth factor has an important role in the EC proliferation (42). We also found that IEL-derived IFN-γ plays an important role in the increase of EC apoptosis (43). All these findings suggested that there is a
strong cross talk between EC and IEL. Future studies will need to focus on how the reported changes in IEL after IL-7 administration affects intestinal EC function.

Our study closely corroborated some of the findings of a recent publication by Fukatsu et al. (6). In their work, the authors noted that exogenous administration of IL-7 increased the number of T cells in Peyer’s patches and IEL in a TPN rodent model. The authors noted that, despite this increase in T cells, they did not find an improvement in the TPN-associated decline in IgA levels. The examination of IEL in their study was, however, quite limited. These authors did not examine the role of endogenous IL-7 in the modulation of the specific IEL phenotype, nor did they examine how IL-7 might affect IEL proliferation, as done in our present study. Furthermore, an assessment of the loss of IL-7 (as performed in our study) was also not investigated by these authors.

In conclusion, the present study demonstrates that TPN administration causes significant changes to both intestinal EC and IEL populations. This study further confirms a close communication between EC-derived IL-7 and IEL. The decline in production of EC-derived IL-7 after TPN administration may present an important mechanism for resultant TPN-associated IEL changes.

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