Effect of a probiotic fermented milk on the thymus in Balb/c mice under non-severe protein–energy malnutrition

Ivanna Novotny Núñez1, Carolina Maldonado Galdeano1,2, Esteban Carmuega3, Ricardo Weill4, Alejandra de Moreno de LeBlanc1 and Gabriela Perdigón1,2*

1Centro de Referencia para Lactobacilos (CERELA-CONICET), Cbaca¿buco 145, San Miguel de Tucumán (T4000ILC) Tucumán, Argentina
2Cátedra de Inmunología, Instituto de Microbiología, Facultad de Bioquímica, Química y Farmacia, Universidad Nacional de Tucumán, Argentina
3Center of Studies on Infant Nutrition (CESNI), Buenos Aires, Argentina
4Departamento de investigación y Desarrollo, DANONE Argentina S.A., Buenos Aires, Argentina

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Abstract

Protein–energy malnutrition (PEM) causes a significant impairment of the immune system, the thymus being one of the most affected organs. It has been demonstrated that the administration of probiotic fermented milk (PFM) recovered the intestinal barrier, histological alterations and mucosal and systemic immune functions in a non-severe malnutrition model using BALB/c mice. The aim of the present study was to evaluate, in the same model of malnutrition, the effect of a PFM added to a re-nutrition diet on the recovery of the thymus, analysing histological and functional alterations caused by malnutrition. Mice were undernourished and divided into three groups according to the dietary supplement received during re-nutrition: milk, PFM or its bacterial-free supernatant (BFS). They were compared with well-nourished and malnourished mice. PFM was the most effective re-nutrition supplement to improve the histology of the thymus, decreasing cellular apoptosis in this organ and recovering the percentage of CD4+/CD8+ single-positive thymocytes. Immature double-positive thymocytes were increased in the malnourished control (MC). The production of different cytokines in the thymus was increased in mice given PFM, compared with the mice that received other dietary supplements and MC. Mice given the BFS presented an improvement in the thymus similar to those that received milk. We demonstrated the importance of the whole PFM supplementation on the histological and functional recovery of the thymus in a non-severe PEM model.

Key words: Fermented milk; Non-severe protein–energy malnutrition; Probiotics; Thymus; Mouse model

The thymus is a primary lymphoid organ in which the differentiation of T lymphocytes occurs(1). It may also be considered as an organ of the endocrine system and, therefore, an endocrine gland that secretes hormones and other soluble factors controlling the production and maturation of T lymphocytes(2,3), and regulating the activity and interactions of T cells in peripheral tissues(3,4,5). It exerts a clear influence on the development and maturation of the lymphatic system and in the mucosal and systemic immune response(4)

It is known that malnutrition causes a significant impairment of the immune system. The severity of the malnutrition depends on the degree of nutrient imbalance and the interaction between nutrients and the age of the host(5). Protein–energy malnutrition (PEM) causes atrophy of lymphoid tissues, the thymus being one of the most affected organs.

Probiotics are defined as live micro-organisms which, when administered in adequate amounts, confer a health benefit to the host(6). The beneficial effect of a probiotic fermented milk (PFM) using a non-severe PEM mouse model was previously reported(7). The administration of the PFM during the re-nutrition period recovered the intestinal barrier, histological alterations and mucosal and systemic immune functions in the mice. It also demonstrated the beneficial effect of the PFM in the protection against Salmonella enterica serovar Typhimurium infection in this same non-severe malnutrition model(7)

The aim of the present study was to evaluate the influence of a PFM, as a complement of a re-nutrition diet in a non-severe PEM model, on the recovery of the thymus,
analysing the histological and functional alterations caused by malnutrition, as a way to determine if the improvement of the immune system observed previously was related to the thymus recovery\(^7\). The effect of the PFM was compared with whole milk or with the bacterial-free supernatant (BFS) obtained from the PFM. BFS was analysed to evaluate the possible effect of biologically active components released during milk fermentation, such as peptides and carbohydrates, or other soluble components present in the fermented milk.

**Experimental methods**

**Malnutrition of animals by food restriction and re-nutrition protocols**

Specific pathogen-free BALB/c mice (for bacteria such as cilia-associated respiratory bacillus *Citrobacter rodentium*, *Salmonella* spp., *Clostridium piliforme* and protozoa such as *Giardia muris*) were obtained from the closed, random-bred colony maintained at Centro de Referencia para Lactobacilos (San Miguel de Tucumán, Argentina).

After weaning (21 d), thirty-five mice (males and females) weighing 10 (SD 2) g, were randomly divided into two control groups (ten mice per group, five males and five females) and three test groups (five mice per group, three males and two females). Well-nourished control (WC) animals received (3–5 g/animal per d) conventional balanced diet (23% proteins, 6% raw fibre, 10% total minerals, 1.3% Ca, 0.8% P, 12% moisture and vitamins). Malnourished control (MC) mice were not renourished and continued with the experimental diet. After that, the malnourishment period started and when the animals lost 25% of their body weight compared with the WC, the restrained ingestion during the entire experiment. Mice of these two control groups received water ad libitum.

The mice of the three test groups received the restricted diet similar to the MC group, and when the animals lost 25% of body weight compared with the WC, the restrained ingestion was continued, but was supplemented with three different dietary supplements: milk, where the WC mice were renourished (rehydrated at 10% (w/v)); PFM, where the animals were renourished with a commercial fermented milk containing yoghurt starter cultures (DN-114-001) (10\(^8\) CFU/ml) or BFS, where the mice received the diet supplemented with a BFS obtained by centrifugation of PFM at 10,000 \(g\) for 20 min. The supernatant obtained was plated on de Mann–Rogosa–Sharpe agar for the bacterial count, which was found to be lower than 10\(^8\) CFU/ml. This CFU number is cleaned by peristalsis. Mice of the three test groups received restricted food containing the specific dietary supplement during the 5 d and water was supplied ad libitum.

A total of five mice were housed in a cage in the control groups and three or two mice per cage in the test groups, maintaining females and males in separate cage. The metallic cages were kept in a climate-controlled room (20 ± 2°C of temperature and 40–70% relative humidity), with a 12 h light–12 h dark cycle. The cages were cleaned every 2 d. A total of five mice from each control group were killed by cervical dislocation after food restriction (25% less than WC, basal sample, day 30 of the experiment, 7 weeks of age) and five mice each of the control and the test groups were killed at 5 d post re-nutrition (day 35 of the experiment, 8 weeks of age). The immune system of the mice is mature at the time of the killing (7–8 weeks of age). The thymus was recovered for histological and immunological studies.

**Histological samples**

The thymus was removed, fixed with 10% formaldehyde in PBS solution (pH 7). After fixation, the tissues were dehydrated and embedded in paraffin using the methods described by Sainte-Marie\(^9\). Serial paraffin sections (4 μm thick) were made and used for immunofluorescence and haematoxylin–eosin staining.

**Apoptosis determination in thymus on histological cuts**

Apoptosis was evaluated for the presence of DNA breaks using the Apoptosis Detection System kit (Promega). The fluorescein-12-dUTP-nick-end-labelled DNA was visualised using fluorescence microscopy. Cells were defined as apoptotic if the entire nuclear area of the cell was stained fluorescent. Apoptosis was expressed as the number of apoptotic cells in ten fields observed at 400× magnification.

**Determination of macrophage and dendritic cells in thymus tissues**

Macrophages and dendritic cells (DC) were determined using indirect immunofluorescence in the thymus tissues, as was previously described\(^10\). The BM8 monoclonal antibody (affinity purified anti-mouse F4/80 antigen – Pan Macrophage Marker, eBioscience) was used to identify mature macrophages and the 33D1 monoclonal antibody to identify DC that are able to present foreign antigens (affinity purified anti-mouse DC marker, e-Bioscience). The results were expressed as the number of positive fluorescent cells per ten fields of vision (1000×).

**Immunohistochemical detection of cytokine-producing cells**

Cytokine-producing cells were detected by indirect immunofluorescence assays following the technique previously described\(^11\). Rabbit anti-mouse TNF-α, interferon (IFN)-γ,
IL-10, IL-6, IL-2 or IL-4 (Peprotech, Inc.) polyclonal antibodies were used. A dilution of fluorescein isothiocyanate-labelled anti-rabbit IgG (Jackson Immuno Research Laboratories, Inc.) was used as secondary antibody. The results were expressed as the number of positive fluorescent cells per ten fields of vision (1000×).

**Determination of CD4⁺ and CD8⁺ T lymphocytes by flow cytometry**

The thymus was aseptically removed, washed with saline solution (0.15M-NaCl) and placed into sterile tubes containing 5 ml of Hank’s balanced saline solution. The samples were immediately homogenised under sterile conditions using a nylon mesh. The samples were centrifuged and resuspended in 2 ml of Hank’s balanced saline solution. Cell concentration was adjusted to 3 × 10⁶ cells/ml using a Neubauer chamber. The cells were incubated with fluorescein isothiocyanate anti-CD8 and phycoerythrin anti-CD4 monoclonal antibodies (both from BD Bioscience Pharmingen) at 4°C for 30 min. The number of CD4⁺ and CD8⁺ cells was determined using a BD FACSCaliburTM Flow Cytometer equipped with 488 and 635 nm laser excitation sources.

**Statistical analysis**

Statistical analysis were performed using MINITAB 14 software (Minitab, Inc.) by general linear model ANOVA, followed by a Tukey’s post hoc test, and P<0.05 was considered significant. Unless otherwise indicated, all values (n=15) were the means of three independent trials (n=5 per trial, and non-significant differences were observed between individual replicates) with their standard deviations. For each immunohistochemical determination and for each mouse, the results were obtained from two individual blind counts (by two different investigators).

**Results**

**Effect of milk, probiotic fermented milk or its bacterial-free supernatant on the histology of the thymus**

The histology of the thymus was affected in the MC group in more than 75% of the animals, where a loss of corticomedullary differentiation with lobule decreases were observed compared with the WC group (Fig. 2(A) and (B)). These changes were reversible after PFM administration during the re-nutrition period (Fig. 2(C)). Mice given milk or BFS also showed a similar effect, but the improvement was lower than in the animals that received PFM (Fig. 2(D) and (E)). No differences were found between male and female animals.

**Effect of the different re-nutrition supplements on thymic cellular apoptosis**

The analysis of apoptosis showed a significant increase in the MC group (78 (sd 6)) compared with the WC group.
Formicethatreceivedmilk(43( SD 6)), PF M(32( SD 7)) or BFS (43 ( SD 6)) the number of apoptotic cells decreased, reaching values even lower than the WC group (Fig. 3).

Influence of different supplements added to diet on marker expression in macrophages and dendritic cells

The number of macrophages and DC in thymus was not affected by malnutrition. Mice re-nourished with milk or PF M maintained the number of these cells similar to the MC group and close to the WC group. In the BFS group, the number of macrophages increased significantly and the number of DC decreased significantly compared with the MC and WC groups (Fig. 4(A) and (B)).

Evaluation of CD4<sup>+</sup> and CD8<sup>+</sup> (single- and double-positive) T lymphocytes

The analysis of T-cell populations in thymus showed a significant decrease of single-positive thymocytes (CD4<sup>+</sup>CD8<sup>-</sup>) in the mice from the MC group (11 (sd 1)%), compared with the mice from the WC group (17 (sd 2)%). In mice that received PF M (17 (sd 2)%), milk (14 (sd 0)%), or BFS (15 (sd 0)) as re-nutrition diet, these values increased significantly compared with the MC group. The effect was remarkable for PF M, where the values were similar to the WC group. The percentage of immature double-positive thymocytes (CD4<sup>+</sup>CD8<sup>+</sup>) increased significantly in mice from the MC group (81 (sd 1)%), compared with the WC control group (76 (sd 2)%), and the test groups, milk (77 (sd 1)%), PF M (74 (sd 2)%), and BFS (78 (sd 2)%). No significant differences were observed in the percentage of single-positive (CD8<sup>-</sup>CD4<sup>-</sup>) cells among all the groups.

Determination of cytokine expression in thymus tissues

The number of IL-2-producing cells decreased in the MC group (14 (sd 5)) compared with the WC group (28 (sd 7)). IL-2<sup>+</sup> cells increased significantly only in the thymus of mice re-nourished with PF M (26 (sd 5)) compared with the MC, with these values being near to those obtained in the WC group. No significant differences were found between the mice that received milk (21 (sd 6)) or BFS (21 (sd 8)); however, the production of this cytokine increased in these groups compared with the MC group.

IFN-γ<sup>+</sup> cells increased in the MC group compared with the WC group. This cytokine also increased significantly in mice that received PF M (49 (sd 6)) compared with the WC group (34 (sd 7)), but it did not show significant differences between the MC group (46 (sd 13)) and the group of mice given milk (47 (sd 9)) as a re-nutrition diet, while BFS administration (22 (sd 8)) showed lower levels than the other groups under study.

![Fig. 3. Apoptosis assays in thymus comparing the different test groups. Cellular apoptosis was evaluated in the slices from thymus for the presence of DNA breaks using the Apoptosis Detection System kit (Promega). Apoptosis was expressed as the number of apoptotic cells in ten fields observed at 400× magnification. Values are means for n 15 mice from each group, with standard deviations represented by vertical bars. * Mean values with unlike letters were significantly different (P<0.05).](https://www.cambridge.org/core/terms).
TNF-α+ cells increased significantly in the MC group (45 (SD 4)) compared with the WC group (34 (SD 6)). No significant differences were observed among the test groups, milk (36 (SD 8)), PFM (35 (SD 7)) and BFS (28 (SD 4)), where the values obtained were similar to the WC group.

The analysis of IL-6+ cells showed a diminution in the MC group (21 (SD 6)) in comparison to the WC group (35 (SD 5)). In mice given milk (33 (SD 7)) or PFM (35 (SD 6)), these cells increased significantly compared with the MC group, reaching values similar to the WC group. For the BFS group (19 (SD 5)), the values were similar to the MC group.

IL-4+ cells in the thymus of the animals that received BFS decreased significantly (15 (SD 6)) compared with the WC group (34 (SD 5)). No significant differences were found among the rest of the groups (MC group: 26 (SD 7), milk: 28 (SD 5) and PFM: 24 (SD 2)). These cells were diminished when compared with the WC group.

IL-10+ cells only increased significantly in mice that received PFM (41 (SD 3)), compared with the WC group (21 (SD 6)) and the MC group (31 (SD 4)). The re-nutrition diets supplemented with milk (21 (SD 6)) did not decrease the number of IL-10+ cells significantly in comparison to the MC group. The values of IL-10+ cells in the BFS group (12 (SD 5)) were lower than the MC group.

The values of IL-12+ cells were similar to the WC group in all the test groups (Table 1).

**Discussion**

The thymus is one of the most affected organs during an undernourished process. In malnourished mice, the thymus undergoes histological atrophy and changes in the lymphoid compartment\(^{(1-4)}\). It was observed that in severe malnourished mice the thymic intralobular extracellular matrix containing fibronectin, laminin and type IV collagen increased. The enhancement of these extracellular matrix was correlated with the degree of thymocyte depletion\(^{(12)}\).

In our non-severe PEM model using BALB/c mice, the administration of PFM or its BFS as a re-nutrition supplementation improved thymic histology faster than milk administration, with this effect being more remarkable for PFM (Fig. 2), and no differences were observed between males and females. This histological improvement induced by the different treatment was accompanied by a diminution in cellular apoptosis (programmed death cells). In this sense, it has been suggested that nutritional imbalances can induce apoptosis in a variety of cell types\(^{(13,14)}\). We observed that malnourished mice increased the number of apoptotic cells in the thymus; this result agrees with the findings of other authors, where malnutrition was associated with a significant increase of spontaneously apoptotic cells in the thymus\(^{(15,16)}\). The thymus samples obtained after the re-nutrition period showed that the three dietary supplements decreased cellular apoptosis in this organ, reaching values similar or slightly lower than the WC group (Fig. 3). These results demonstrated that the physiological equilibrium between cellular mitosis and apoptosis was improved, and that the three dietary supplements assayed would stimulate the activation of the cells instead of cell death.

The improvement observed in the histology of the thymus and the diminution in cellular apoptosis led us to analyse other immune cell populations, such as the antigen-presenting cells, i.e. macrophages and DC. These cells participate in the process of selection and maturation of T lymphocytes that are unable to recognise self-antigens, before they are released from the thymus to peripheral secondary immune tissues (the spleen and the lymph). The analysis of these two antigen-presenting cell populations (Fig. 4(A) and (B)) showed that in the non-severe PEM model used in the present study, no significant differences were observed in the MC group compared with the WC group. The numbers of these cells were also not affected by re-nutrition with the PFM or milk; however, BFS administration induced an increase in the number of macrophages accompanied with a significant decrease in the number of DC in the thymus, compared with the malnourished and well-nourished mice. We believe that the number of mature macrophages increases in detriment of DC to favour the positive selection of T cells to be released from the thymus to the periphery, because the
macrophages are more effective than DC in this function. However, this lack of regulation by the BFS would not be desirable, because it is far from the physiological condition (the WC group). The effect obtained with the PFM and milk groups was more regulated and near to the physiological conditions.

T lymphocyte subsets were also analysed to evaluate if the re-nutrition diets can influence this population that is severely

![Graph](https://chart.example.com/graph.png)

**Fig. 5.** CD4+ and CD8+ (single- and double-positive) T lymphocytes in thymus. The number of CD4+ and CD8+ cells was determined by flow cytometry. (A) Values are means for n 15 obtained from mice of each group, with standard deviations represented by vertical bars. **a,b** Mean values with unlike letters were significantly different for each cell population (P<0.05). **x** Well-nourished control (WC); **y** malnourished control (MC); **z** mice re-nourished with milk (M); **w** probiotic fermented milk (PFM); **e** bacterial-free supernatant (BFS). Representative dot plots of the different groups are shown in (B) WC, (C) MC, (D) M, (E) PFM and (F) BFS. FL-1 is for fluorescein isothiocyanate (FITC) and FL2 is for phycoerythrin (PE). FIFC anti-CD8 and PE anti-CD4 antibodies were used.
affected by malnutrition. In this sense, there are publications showing that increased apoptosis affects CD4⁺CD8⁺ double-positive lymphocytes in severe malnutrition. However, other studies did not show significant differences in this double-positive cell population comparing the thymus of well-nourished and malnourished rats. The present results showed that non-severe PEM decreased the percentage of CD4⁺CD8⁺ single-positive thymocytes, with increases in the percentage of immature double-positive thymocytes (CD4⁺CD8⁻). In WC, this population is able to mature into single-positive (CD4⁺CD8⁻ or CD4⁺CD8⁺) T cells and to be released from the thymus into peripheral tissues. This increase in double-positive thymocytes was not observed in the test groups that received PFM or BFS, where the re-nutrition diet was able to recover the percentage of CD4⁺CD8⁻ single-positive T cells. No significant differences in the immature double-positive thymocytes were observed in the mice that received any supplement diets, with the values being similar to those obtained from the WC group. These observations agree with the improvement in the histology of the thymus and with the diminution of cellular apoptosis results from the mice that received re-nutrition diets. The PFM group showed the best effect on the T-cell population of the thymus and reached the number of CD4⁺CD8⁻ single-positive thymocytes similar to those observed in the well-nourished mice, which agree with the improvement of the thymocyte subsets after re-nutrition.

Another cytokine that can be affected by malnutrition is IFN-γ, which is produced mainly by T lymphocytes and activates thymic epithelial cells. In our model, differing from the results obtained for other authors, we did not observe significant differences after malnutrition in the MC group or in the group that received milk compared with the WC group. The BFS group showed a significant diminution in this cytokine. However, re-nutrition with PFM increased the number of IFN-γ-positive cells significantly in the thymus compared with the well-nourished mice. This cytokine could be produced by T cells, macrophages and also by natural killer T cells and the increase observed could be attributed to increases in the functionality or in the number of these cell populations. The increases observed for IFN-γ showed that PFM administration could activate thymic epithelial cells and it could influence the proliferation and activation of thymocytes.

Other cytokines related to macrophage functionality, such as TNF-α and IL-12, did not have the same profile. The numbers of IL-12 producer cells were similar in the thymus after malnourished or re-nourished periods, which would be in agreement with the results obtained for macrophages. However, the TNF-α-positive cells increased significantly in the MC group and decreased after re-nutrition without differences between the diets. These increases in TNF-α could be related to the increases in the thymic apoptotic cells observed in the mice from the MC group, as was reported in other models.

IL-6 was the other cytokine analysed, which is produced primarily by thymic epithelial cells and can also be produced by CD4⁺ T lymphocytes and macrophages. In the present study, IL-6-positive cells decreased significantly in malnourished mice, which was in accordance with the decrease in the mature T-cell subset. IL-6 producer cells increased in the mice after re-nutrition with milk or PFM, but not with BFS, reaching the values observed in well-nourished mice in the first two groups. This observation could be related to the restoring of the thymocyte proliferation and maturation observed after consumption of the re-nutrition diet.
The analysis of IL-4 showed that this cytokine decreased after malnutrition, with the number of IL-4-positive cells being significantly diminished in the groups that received PFM or BFS compared with both the WC and the MC groups. IL-4 can be produced by thymocytes and also by natural killer T cells. It is known that IL-4 inhibits the growth and differentiation of fetal mouse thymus in organ culture. The lack of increase in this cytokine after PFM administration agrees with the increase in IFN-γ production in the thymus of these mice, where, as was discussed earlier, IFN-γ is involved in the activation of the thymic epithelial cells to release other cytokines and to regulate IL-4 production, which can participate in the activation and proliferation of thymocytes.

Finally, IL-10 was studied and the results showed that the mice given PFM during the re-nutrition period showed significantly increased number of positive cells for this cytokine. These increases can contribute to the decrease in cellular apoptosis observed after re-nutrition. IL-10 can act as an anti-apoptotic factor, increasing Bcl-2 expression (related to mitosis), reducing caspase-3-dependent thymocyte apoptosis and improving survival.

It is important to note that, in the analysis of the cytokines, the best improvement in thymus function was observed with the administration of PFM during the re-nutrition period. Mice given the BFS presented a slight improvement in the thymus, similar to the milk re-nutrition. This fact demonstrated the importance of the whole fermented product to have the best effect in the histology and function of the thymus in this non-severe PEM model.

This is a first report in an animal model of malnutrition using BALB/c mice, demonstrating the effect of probiotic administration on the thymus after a period of malnutrition. The improvement in thymic function would be also responsible for the increase in the mucosal and systemic immune responses observed in previous studies using the same non-severe malnutrition model.

Studies to find out more about the probiotic effect on different cell populations, such as natural killer T cells, and on the components from thymic intralobular extracellular matrix, such as fibronectin, involved in the maturation or in the cellular differentiation in the thymus of malnourished mice are currently underway.

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