Dietary Oils and Phospholipids Containing n-3 Highly Unsaturated Fatty Acids Suppress 2,4-Dinitro-1-fluorobenzene-induced Contact Dermatitis in Mice

Kazuya MORIZAWA*1, Yoko I.TOMOBE*2, Mamoru TSUCHIDA*2, Yoshio NAKANO*2
Hidehiko HIBINO*2 and Yukihisa TANAKA*2

*1 NOF Corporation Food Research Laboratory
(4-18-11, Toshima, Kita-ku, Tokyo 114-0003)
*2 NOF Corporation Tsukuba Research Laboratory
(5-10, Tokodai, Tsukuba-shi, Ibaraki-ken 300-2635)

Abstract: Dietary perilla oil, tuna oil and salmon roe phospholipids (fish roe PL) containing n-3 polyunsaturated fatty acids were compared with n-6 fatty acid-rich corn oil for their anti-inflammatory activities in the contact hypersensitivity reaction in the ears of mice sensitized with 2,4-dinitro-1-fluorobenzene (DNFB). Suppression of ear swelling was most active with fish roe PL followed by tuna oil and Saibokuto, a traditional Chinese medicine, when compared with the corn oil diet group; perilla oil enriched with α-linolenic acid (n-3) tended to suppress the ear swelling, but the effect was not statistically significant. Infiltration of inflammatory CD4-positive T lymphocytes into the ears was suppressed by fish roe PL. Fish roe PL diet suppressed the expression of mRNAs for IFN-γ, IL-6 and IL-1β in the ears. These results suggest that tuna oil and fish roe PL may be effective in suppressing delayed-type hypersensitivity.

Key words: n-3 Highly unsaturated fatty acid, Contact hypersensitivity reaction, Contact dermatitis, Anti-inflammation, Expression of cytokine mRNA

1 Introduction

Many researchers have reported that an intake of polyunsaturated fatty acid (PUFA), especially n-3 PUFA has a beneficial role in the prevention of cardiovascular diseases\(^1\),\(^2\), which is, at least in part, due to reduced atherosclerosis\(^3\). Eicosapentaenoic acid (EPA; 20:5), in particular, has been shown to be a useful agent in the prevention of hypertriglyceridemia\(^4\) and osteoporosis\(^5\). In the last decade, there has been considerable interest in the effects of different types of dietary PUFA on the immune system\(^6\). This interest in the immunomodulatory effects of fatty acids arises from epidemiological studies which showed that populations, such as the Inuit who consume large quantities of fish oil rich in n-3 PUFA EPA and docosahexaenoic acid (DHA; 22:6), have a very low incidence of inflammatory and autoimmune problems\(^6\). Further, many clinical studies indicate that fish oil supplementation of the human diet has beneficial effects in acute and chronic inflammatory conditions\(^7\)–\(^13\). Laboratory animals fed on diets rich in n-3 PUFA (Canola, linseed, fish oil), particularly those containing fish oil, have increased immunosuppression\(^14\)–\(^17\). Arm\(^18\) has reported that fish oil supplementation ameliorated the symptoms of asthmatic subjects. Bjørneboe\(^19\) demonstrated an improvement in the condition of atopic dermatitis patients after treatment with dietary Max-Epa (18% EPA and 12% DHA). The mechanism underlying skin lesion formation, in atopic dermatitis, has remained unclear except that there is a delayed hypersensitivity reaction against environmental substances\(^20\),\(^21\) or an Ig-E mediated hypersensitivity reaction against food or inhaled allergens\(^20\),\(^22\). The eczematous skin lesions contain cytokine-producing, CD-4 positive T cells.

Corresponding author: Yukihisa TANAKA
(T-helper cells, Th cells)\(^{24,25}\) which have been divided into two subclasses depending on their cytokine secretion pattern: the Th1-like subtype, which is characterized by predominant production of interferon (IFN)-γ, and interleukin (IL)-2 and the Th2-like subtype, by its synthesis of IL-4\(^{26}\). In previous studies 85% of skin samples obtained from chronic eczematous lesions of atopic dermatitis patients, were found to contain increased levels of IFN-γ mRNA, whereas increased IL-4 mRNA expression was observed only in 25% to 30% of cases\(^{23}\). In those atopic dermatitis patients whose skin disease responded to treatment, the increased IFN-γ mRNA, but not the increased IL-4 mRNA levels, were down-regulated. These studies indicate that the Th1-like cytokine IFN-γ plays a major role in the maintenance of chronic eczematous lesions in atopic dermatitis patients. It has already been reported that IFN-γ, among the several Th1-types of cytokine, is especially important for the formation of a contact hypersensitivity reaction (CHR)\(^{27}\). Although there are limitations to the use of the CHR in mice as a model for chronic atopic dermatitis, it is one approach for studying the late phase of this disease.

In our study we investigated the immunomodulatory effects of dietary edible oils containing n-3 PUFA, fed at the level of 4.8wt% of the total diets, on the inflammatory response, in the ears of mice sensitized with 2,4-dinitro-1-fluorobenzene (DNFB), in the challenge phase of CHR. Corn oil, 4.8wt%, was used as the control diet. We show that perilla oil, fish oil and the phospholipid from salmon roe (fish roe PL) suppressed the ear swelling and the expression of Th1-type cytokines and decreased the infiltration of inflammatory cells into the challenge phase of CHR.

### Table 1 The Fatty Acid Composition of Test Diets (GC%).

<table>
<thead>
<tr>
<th></th>
<th>Corn oil</th>
<th>Tuna oil</th>
<th>Fish roe PL</th>
<th>Perilla oil</th>
</tr>
</thead>
<tbody>
<tr>
<td>16:0</td>
<td>11.2</td>
<td>10.6</td>
<td>13.1</td>
<td>7.1</td>
</tr>
<tr>
<td>16:1</td>
<td>0.2</td>
<td>4.9</td>
<td>1.4</td>
<td>0.5</td>
</tr>
<tr>
<td>18:0</td>
<td>2.2</td>
<td>2.2</td>
<td>8.7</td>
<td>1.9</td>
</tr>
<tr>
<td>18:1</td>
<td>23.9</td>
<td>14.5</td>
<td>11.9</td>
<td>8.0</td>
</tr>
<tr>
<td>18:2</td>
<td>61.4</td>
<td>16.2</td>
<td>15.5</td>
<td>26.1</td>
</tr>
<tr>
<td>18:3(n-3)</td>
<td>—</td>
<td>0.6</td>
<td>0.2</td>
<td>50.5</td>
</tr>
<tr>
<td>20:4(AA)</td>
<td>—</td>
<td>1.8</td>
<td>2.0</td>
<td>—</td>
</tr>
<tr>
<td>20:5(EPA)</td>
<td>—</td>
<td>8.9</td>
<td>9.0</td>
<td>—</td>
</tr>
<tr>
<td>22:6(DHA)</td>
<td>21.2</td>
<td>25.1</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>others</td>
<td>1.1</td>
<td>19.1</td>
<td>13.1</td>
<td>5.9</td>
</tr>
</tbody>
</table>

Each diet contained 1.2wt% of safflower oil and 4.8wt% of test oil cold acetone. The extract contained mainly triacylglycerol, phosphatidylcholine and phosphatidylethanolamine (in a ratio of 1:8:1 by weight). Saibokuto (Tsumura Inc. Japan) which is an anti-inflammatory Chinese medicine was added to the control diet (1.0 wt%) as a positive control (Saibokuto diet). The fatty acid compositions of the test lipids are shown in Table 1. All diets were divided into small packages and stored at 4°C and, to minimize lipid peroxidation, were provided fresh to the mice every day.

#### 2.2 Feeding Design and Induction of CHR

Female, 4 week old, ddy mice obtained from Charles River Japan Inc. (Tokyo Japan) were held for one week and then divided into four groups (n = 4). Body weight was measured twice a week. After 23 days of feeding the test diets, the back hair of the mice was shaved. On the next day, the shaved backs of the mice were treated with 0.1 ml of 0.5% DNFB diluted with an acetone/olive oil mixture (4:1 v/v) to induce the inflammatory reaction (Induction step). After 5 days, 20 μl of the same DNFB mixture was applied to the sensitized mouse ears (Challenge step). At 6 h and 24 h after the challenge, the thicknesses of the mouse ears were measured using a thickness-gauge (OZAKI MFG. Co. LTD., Tokyo) and an increase in thickness was defined as ear swelling.

#### 2.3 Histopathological Analysis

Twenty four hours after the challenge, the mice were sacrificed and their ears embedded in Tissue-Tek OTC (Miles; Elkhart, IN) and frozen using dry ice/acetone. Serial 6 μ sections were cut on a cryostat and placed on poly-L-lysine-coated slides.
Table 2 Histopathological Findings of Test Animals (mean ± SD).

<table>
<thead>
<tr>
<th></th>
<th>Inflammatory cell infiltration</th>
<th>Edema</th>
</tr>
</thead>
<tbody>
<tr>
<td>Corn oil</td>
<td>4.0 ± 0.0</td>
<td>4.0 ± 0.0</td>
</tr>
<tr>
<td>Saibokuto</td>
<td>3.0 ± 0.0</td>
<td>3.8 ± 0.5</td>
</tr>
<tr>
<td>Fish roe PL</td>
<td>*2.5 ± 0.6</td>
<td>*2.8 ± 0.5</td>
</tr>
</tbody>
</table>

1: very slight, 2: slight, 3: medium, 4: severe, 5: more severe*; p<0.01 Significantly different from corn oil diet group (n=4).

Infiltration of inflammatory cells into the epidermis and the severity of edema were observed and ranked by degree in one of five levels: rank 1, very slight; rank 2, slight; rank 3, medium; rank 4, severe and rank 5, most severe (Table 2).

2.4 Immunohistochemistry

Serial 8μm sections were cut on a cryostat and placed on poly-L-lysine-coated slides (two sections per slide). Slides containing frozen sections were defrosted and dried at room temperature (RT) for 1h, fixed in cold acetone for 10min and allowed to dry at RT for 10 min. After washing in phosphate buffered saline (PBS) (5min x 3), non-specific binding sites were blocked with block ace (Dainippon Seiyaku Inc. Japan). Excess solution was removed, the sections circled with a waterproof pen (Dako; Glostrup, Denmark), and then incubated with the primary monoclonal antibody (MAB), anti-mouse CD4 (L3/T4) (Cedarlane Lab. Ltd., Canada) or anti-mouse CD8α (KT15) (Serotec Ltd., Oxford England) at 37°C for 1 hr. All MABs against mouse antigens were rat immunoglobulin G (IgG) and the diluent was 10% block ace. Negative control slides were incubated in diluent alone. After washing in PBS the slides were incubated with biotinylated rabbit anti rat IgG pre-absorbed with powdered mouse liver, followed by avidin-biotinylated horseradish peroxidase complex (ABC) and diaminobenzidine according to the Vectastain protocol (Vector, USA). For individual slides, the nuclei were counterstained with methyl green (Vector, USA).

2.5 Expression of Cytokine mRNA

Cytokine mRNA expression was measured by the reverse transcription-polymerase chain reaction (RT-PCR). The ears were taken from the mice 24h after the challenge and homogenized with 1.0 ml of 4M-guanidium isothiocyanate buffer. After this, the total RNA was extracted and reverse transcribed to cDNA using Moloney Murine Leukemia Virus (M-MuLV) reverse transcriptase and a random primer [pd (N)6] (Amersham Pharmacia Biotech, Sweden). Identical amounts of cDNA, based on the expression of glyceraldehyde-phosphate dehydrogenase (GAPDH) as the house-keeping gene, were subjected to 35 cycles of PCR with cytokine primers (Clontec Inc.). The products were electrophoresed on a 1.0% agarose gel and visualized by ethidium bromide staining. The samples were screened using the computer software Bioimage (Bioimage Japan Inc. Tokyo Japan, Version 0.81) and the relative amounts of cytokine cDNA of each sample was calculated.

2.6 Data Analysis

The data were expressed as the mean ±SD of four mice fed each diet. The data were analyzed by ANOVA and Bonferroni’s test. A p value of less than 0.05 was considered to be statistically significant.

3 Results

3.1 Mouse Growth

The final body weights, and therefore the total growth, did not differ among the groups fed on the different diets (data not shown) and no side-effects were observed among mice fed on the diets described.

3.2 Ear Swelling

The differences in ear thickness (mean ±SD) before and after the challenge (ear swelling) are shown in Fig. 1 A and 1 B. The ear swelling of mice fed the corn oil diet had increased in a time dependent manner in the early late phase (6h after the challenge). However, the ear swelling of the mice fed tuna oil and fish roe PL were significantly suppressed compared with that of the mice fed the corn oil diet (p<0.05, and p<0.01 respectively) during this period and this continued to be apparent in mice fed both diets in the late phase (24h after the challenge) (p<0.01).

The mean value of the ear swelling of the mice fed perilla oil was also suppressed compared with that of the mice fed on the corn oil diet. This difference was not statistically significant but this is
A

Fig. 1  Ear Swelling of Mice Fed Test Diets 6 Hours (A) and 24 hours (B) after the Challenge. *P<0.05: significantly different from the corn oil diet group. **P<0.01: significantly different from the corn oil diet group. #P<0.01: significantly different from tuna oil diet group. (one-way ANOVA followed by Bonferroni’s test).

The data are expressed as mean ±SD (n=4).

probably due to the large standard deviation (SD) in this group. The ear swelling of the mice fed the fish roe PL was the lowest of all groups and was significantly suppressed compared with that of those fed the tuna oil diet (p<0.01). This latter observation is curious as the fatty acid composition of fish roe PL is similar to that of tuna oil.

3.3 Histopathological Analysis

Figure 2 shows the photomicrographs of the ears, stained with hematoxylin and eosin, of the mice fed corn oil and fish roe PL that exhibited the most suppression of their inflammation. On histological examination, 24 h after the challenge, spongiosis (paracytic edema of the epidermis and corium) and mononuclear cell infiltration into the epidermis were observed in the ear tissues from the mice fed the corn oil diet. Considerable infiltration of the lymphocytes (mostly monocytes) was also observed at the stratum reticular. Typical photographs are shown in Fig. 2 A. The number of infiltrated inflammatory cells in the ear tissues from mice fed the fish roe PL diet was reduced compared with those from mice fed the corn oil diet (Fig. 2 B). The severity of the ear tissue damage sustained by the mice fed each diet is shown in Table 2. Formation of focal granulation tissue in the dermis was not observed in the ear tissues from the mice on the fish roe PL diet 24 h after the challenge.

3.4 Immunohistological Study

Results obtained 24 h after the challenge are shown in Fig 2. Infiltration of CD4-positive cells into the corium was observed in the ear tissues from mice fed the corn oil diet, (Fig. 3 A) whereas this was considerably less in tissues from the mice fed fish roe PL (Fig. 3 B). In the ear tissues from the mice fed the corn oil diet, some infiltration of CD8-positive cells into the corium, which was less than that of CD4-positive cells, was also observed.

3.5 The Expression of Cytokines in the Ear Tissues from Mice in the Challenge Phase of CHR

We examined the cytokine mRNA expressed at the CHR sites (Fig. 4). Twenty four hours after
Fig. 3 Immunohistochemical Analysis. A: The ear of the mouse fed the control diet 24 hours after challenge. B: The ear of the mouse fed fish roe PL diet 24 hours after the challenge. One bar equals 100 $\mu$m.

The expression of IL-1$\beta$, IL-2, IL-6 and IFN-$\gamma$, but not IL-4, IL-3 and TNF-$\alpha$ mRNA were expressed in the ear tissues from the mice fed the control diet. The expression of IL-1$\beta$ and IFN-$\gamma$ mRNA was less in the ear tissues from the mice fed fish roe PL than that observed from mice fed the corn oil significantly. The expression of IL-6 mRNA was tend to be less in the ear tissues from the mice fed fish roe PL than that from mice fed the corn oil, but not significantly. Under these conditions, the level of GAPDH mRNA did not change.

4 Discussion

In this study we have examined the anti-inflammatory and immunosuppressive effects of dietary oils and phospholipids that contained n-3 PUFA such as $\omega$-linolenic acid (ALA), EPA and DHA, in a rodent contact dermatitis model. Generally it is thought that inflammatory mediators are generated from PUFA through an arachidonate cascade and play important roles in the development of the inflammatory process. Arachidonic acid (AA) is converted to 2 series of prostaglandins (PGs) and 4 series of leukotrienes (LTs) which are all mediators of inflammation. In particular the 4 series of LTs intensify neutrophil chemotactic responsiveness in the development of the inflammatory process. However, EPA displaces AA from macrophage phospholipids and effectively competes with it for cyclooxygenase and lipoxygenase binding sites to foster the formation of eicosanoids (e.g. thromboxane (TX) A$_3$, LTB$_5$ and PGE$_1$) which have relatively lower inflammatory properties than those generated.

Fig. 4 Expression of Cytokine mRNAs in the Ears of Mice Fed Corn Oil, Saibokuto and Fish roe PL diet 24 h after the Challenge. Total RNA from each sample was used and subjected to RT-PCR analysis. For estimation of similar amounts of cDNA to be used for the PCR, samples were screened for the expression of GAPDH as a housekeeping gene and relative amounts of cDNA were calculated using computer software from Bioimage. *: P<0.05: significantly different from the corn oil diet group. The data are expressed as mean ±SD (n=4).
from AA (e.g., TXA2 and LTB4). Palombo reported that rats given an EPA enriched diet had lower amounts of TXA2/A3 and LTB4/B5 in the supernatants of the alveolar macrophages exposed to endotoxin in vitro than those in rats given a standard diet enriched with linoleic acid. The suppression of inflammatory responses by EPA might be due to some adverse effect on the arachidonate cascade such that the eicosanoid products are not formed.

Krokan argued that, as substituting DHA for AA in the phospholipids of the cell membrane was more difficult than substituting EPA, the suppressive effect of DHA on the arachidonate cascade was likely to be less than that of EPA. In contrast, Corey showed that DHA did inhibit AA conversion to PGE2 in an in vitro experiment. Thus, these authors concluded that DHA did affect the arachidonate cascade. DHA also suppressed the production of platelet activated factor (PAF) which occurs following stimulation of cultured Eo-1 cells by the calcium ionophore A23187. Matsumoto reported 1-oleic 2-docosahexaenoic phosphatidylcholine inhibited 5-lipoxygenase (5-LO) activity. These various studies indicate that DHA might well suppress the inflammatory response through modulation of the arachidonate cascade in a similar way to EPA and the effects of both EPA and DHA on the inflammatory response could be explained by the inhibition of this cascade.

To elucidate the immunosuppressive effects of n-3 PUFA, we used the mouse CHR model in which inflammation is induced with DNFB. It has been reported that ear swelling is at its worst approximately 24 h after administration of DNFB and is ameliorated 48 h after the challenge, so this reaction in the mouse is based on a chronic delayed type hypersensitivity (DHT or type IV hypersensitivity). It was also reported that 24 h after the challenge IFN-γ, IL-6, IL-1β mRNA were expressed in the ear tissues of mice and 48 h after the challenge IL-2, IL-4 and TNF-α mRNA were additionally expressed. From results of analysis of IFN-γ knockout mice and using anti INF-γ antibody the authors suggested that the expression of IFN-γ among several Th1-like cytokines is especially important for the formation of CHR.

An inflammatory reaction is induced by the adhesion of circulating leukocytes on the endothelium and their subsequent transendothelial migration. The endothelial expression of the endothelial leukocyte adhesion molecules are important in this mechanism. DeCaterin reported that DHA inhibited cytokine-stimulated expression of the endothelial leukocyte adhesion molecule, VCAM-1. Their results suggested that dietary DHA might cause a decrease in the level of VCAM-1 mRNA, which is consistent with the protein expression and, therefore, the infiltration and chemotaxis of T-cell, monocytes and neutrophils into endothelial cells might be decreased. In this study we found that suppression of ear swelling in mice fed fish roe PL compared with those fed corn oil, caused a decrease in the number of infiltrated CD4-positive cells in lesion sites and a decrease in the expression of inflammatory cytokine mRNAs. The results imply that dietary DHA suppresses inflammation in the ear tissues sensitized with DNFB through these continuous reactions and suggest that not only EPA but also DHA, in its own right, plays an important role in inflammatory suppression.

Although the fatty acid composition of tuna oil was very similar to that of fish roe PL, the suppressive ability of fish roe PL was greater than that of tuna oil. The significant difference observed was probably due to the difference in the molecular compositions which in tuna oil was predominantly triacylglycerol and in fish roe was phospholipid. We hypothesize that this result is likely to arise from differences in the efficiency of the lipid absorption in the small bowel and lipid accumulation in cell membranes of skin. However, Masuzawa reported that DHA in phospholipids from squid was absorbed and accumulated in some organs at the same rate as DHA in triacylglycerol, but there was no information on the accumulation in the skin cell membrane. To confirm our hypothesis we are currently analyzing rates of different lipid absorption and accumulation.

Acknowledgment

We thank Dr. Eiji Terada (Kitasato University, Sagamihara Japan) who passed away in the winter of 1997, for his many useful suggestions. We pray for the repose of his soul.

(Received Mar. 31, 1999; Accepted Oct. 19, 1999)
References

[報文] n-3系高度不飽和脂肪酸を含む食用油脂およびリン脂質は2, 4-ジニトロ-1-フルオロベンゼンでマウスに誘導した接触皮膚炎を抑制する

守沢和也*1・友部（入鹿山）容子*2・土田衛*2
中野善郎*2・日比野英彦*2・田中幸久*2

*1 日本油脂株式会社 食品研究所（〒114-0003 東京都北区東京4-18-11）
*2 日本油脂株式会社 研究所（〒300-2635 茨城県つくば市東光台5-10）

n-3系多価不飽和脂肪酸を含有した食用油脂であるしそ油、tuna oil及び鯖の魚卵リン脂質（fish roe PL）はn-6脂肪酸を含有するとうもろこし油に比べて2,4-dinitro-1-fluorobenzene（DNFB）で誘導された耳介の接触皮膚炎モデルの炎症反応を抑制する作用を持っている。Fish roe PLは耳介腫脹をもっとも強く抑制し、tuna oil及び漢方薬である柴胡湯がそれに続き、αリノレン酸を多量に含むしそ油はとうもろこし油に比べて抑制の傾向は見られなかった。炎症性のCD4+T細胞の耳介への浸潤はfish roe PL投与によって抑制され、耳介においてIFN-γ、IL-6、IL-1βのmRNAの発現抑制が観察された。本実験の結果でfish roe PLは遲延型のアレルギーを抑制する効果を持つことが示唆された。

（連絡者：田中幸久）Vol.49, No.1, 59 (2000)

[ノート] C-24 (28) 位に二重結合を有する C28 ステロイドの合成

高津戸秀*1・後藤千春*1・鴻海安久*1
野口貴弘*2・藤岡昭三*3

*1 上越教育大学自然系化学教室（〒943-8512 新潟県上越市山下敷町1）
*2 タマ生化学（〒163-0704 東京都新宿区西新宿2-7-1）
*3 理化学研究所植物機能研究室（〒351-0198 埼玉県和光市広沢2-1）

シロイヌナズナの酸性変異体のジェートに含まれるステロイドを同定するための標品として、24-methylcholesta-4, 24(28)-dien-3-oneおよび24-methyl-5α-cholest-24(28)-en-3β-olを合成した。合成の鍵反応として24-オキソステロイドのTebbe試薬によるオレフィン化を用いた。さらに、それらの関連化合物である24-methylcholesta-4, 24(28)-dien-3β-olおよび24-methyl-5α-cholest-24(28)-en-3-oneの合成も行った。

（連絡者：高津戸秀）Vol.49, No.1, 67 (2000)