Low-molecular-weight fucoidan and high-stability fucoxanthin from brown seaweed exert prebiotics and anti-inflammatory activities in Caco-2 cells

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Abstract

Background: The aim of this study is to investigate the anti-inflammatory effects of low-molecular-weight fucoidan (LMF) and high-stability fucoxanthin (HS-Fucox) in a lipopolysaccharide-induced inflammatory Caco-2 cell line co-culture with B. lactis.

Methods: We used various methods such as transepithelial resistance (TER) assay, cytokine secretion assay, and tight junction protein mRNA expression assay to examine LMF and HS-Fucox anti-inflammatory properties.

Results: LMF and HS-Fucox activated probiotic growth and reduced the inflammation of the intestinal epithelial cells. Moreover, the combination of LMFHS-Fucox dramatically enhanced the intestinal epithelial barrier and immune function against the lipopolysaccharide effect by inhibiting IL-1β and TNF-α and promoting IL-10 and IFN-γ.

Conclusion: These findings suggested that LMF and HS-Fucox, alone or in combination, could be the potential natural compounds to enhance the immune system and have an anti-inflammatory effect on the intestinal cells.

Keywords: low-molecular-weight fucoidan; fucoxanthin; anti-inflammatory; prebiotics

Received: 23 April 2016; Revised: 20 June 2016; Accepted: 20 June 2016; Published: 2 August 2016
Materials and methods

Materials
LMF (Hi-Q Oligo-Fucoidans®) and HS-Fucox were derived from Sargassum hemiphyllum and prepared by Hi-Q Marine Biotech International Ltd. (New Taipei City, Taiwan). LMF was obtained by enzyme hydrolysis of original fucoidan. The characteristics of LMF-LJ were average molecular weight of 0.8 KDa (92.1%), fucose content 210.9±3.3 μmol/g, and sulfate content 38.9±0.4% (w/w). The extraction method followed the method mentioned before with technological modifications (24). HS-Fucox is a mixture of brown seaweed extract containing about 10% of fucoxanthin that is coated directly with polysaccharides of its own. It was dissolved in double-distilled H2O (ddH2O) and completely dissolved by stirring at room temperature for 30 min.

Bifidobacterium lactis cultivation and growth curve
Bifidobacterium lactis BCRC 17394 was purchased from the Bioresource Collection and Research Center, Hsinchu, Taiwan. This strain was subcultured in Man Rogosa and Sharpe medium (Scharlau) at 37°C under anaerobic conditions for 24 h, diluted with sterile saline to have the cell density of 1 × 10^8 cfu/mL, and used as inoculums for the following cultivation experiments. B. lactis had an initial cell count of 4.5 log cfu/ml and cultured with various concentrations (0, 10, 50, 100, and 200 μg/ml) of LMF, HS-Fucox, and LMF+HS-Fucox for 48 h. LMF+HS-Fucox was a mixture with 50% LMF and 50% HS-Fucox. The cell counts of the sample groups were compared with the initial cell counts. The cell count measurement was carried by a serial 10-fold dilution, followed by spreading 0.1 ml of the diluents onto plate count agar plates. The plates were incubated at 37°C for 24 h prior to counting the colony to obtain cfu/ml.

Cell line and culture condition
Caco-2 cell, the human intestinal epithelial cell, was obtained from the American Type Culture Collection. Caco-2 cells were cultured in Dulbecco’s Modified Eagle’s medium (DMEM), supplemented with a 15% fetal bovine serum (FBS) and 1% penicillin-streptomycin at 37°C in an atmosphere of 5% CO₂. The medium was renewed every two days.

Co-culture of Caco-2 cells and B. lactis
Caco-2 cells were grown for 12 days in six-well tissue culture plates to allow full differentiation to occur. At 12 h, prior to the addition of the B. lactis, the medium was aspirated and replaced with antibiotic-free DMEM. Caco-2 cells cultured in six-well plates were previously determined to contain 2 × 10^8 cells/well, and 1 μg/ml LPS was added to each group, except the control group. 100 μl of B. lactis (10^6 cfu/ml) was added with or without a sample onto Caco-2 cells in separate wells. The plates were incubated for 8 h at 37°C in an atmosphere of 5% CO₂. After being incubated, transepithelial resistance (TER) was determined directly by meter, the supernatants were collected for cytokine secretion assay, and the Caco-2 cells were washed twice with 1-ml phosphate buffered saline (PBS), then for tight junction protein mRNA expression assay.

Cell viability
The cell viability of cells was determined by 3-(4,5-Dimethylthiazol-2-yl)-2,5-Diphenyltetrazolium Bromide (MTT) colorimetric assay (25). Cells were reacted with MTT (1 mg/ml) for 4 h, and absorbance was recorded at 570 nm (26). The cell viability (%) was determined as (A1 / A0) × 100%, where A0 and A1 were the absorbance of the control group (meaning, in the absence of sample and LPS treatment) and the sample group (0, 50, and 200 μg/ml with LPS treatment), respectively.
**TER assay**

TER was measured in Ω cm$^2$ using a Millicell ERS-2 Epithelial Volt-Ohm Meter (Millipore, Bedford, MA), by placing separate electrodes in the upper and lower wells according to the manufacturer’s instructions (27). Monolayers showing TER values of 200–300 Ω cm$^2$ were used for the experiments.

**Tight junction protein mRNA expression assay**

Total RNA was isolated by RNAzol B (Amersham Pharmacia Biotech, Sweden), and the concentration of total RNA was detected by spectrophotometer (Hitachi, Japan). The synthesis of cDNA was performed using Improm-II TM Reverse Transcriptase (Promega, WI, USA) according to the manufacturer’s instructions. Polymerase chain reaction (PCR) was performed on the reverse-transcribed cDNA product to determine the expression of occludin, claudin-1, claudin-2, and β-actin (as an internal control) using a thermal cycler (Biometra, UNO-Thermoblock, UK). The initial 1 min of 95°C denaturation was followed by the amplification sequence protocol of occludin (1 min of 55°C annealing and 3 min of 72°C extension), claudin-1 (1 min of 60°C annealing and 1 min of 72°C extension), claudin-2 (1 min of 56°C annealing and 1 min of 72°C extension), and β-actin for 30 cycles. Primers were listed 5′-3′ as follows: Ocludin: F, TCA GGG AAT ATC CAC CTA TCA CTT CAG; R, CAT CAG CAG CAG CCA TGT ACT CCT CAC. Claudin-1: F, GGC CGA TAT TTC TTC TTG CAG G; R, TTC GTA CCT GGC ATT GAC TGG. Claudin-2: F, CTC CCT GGC CTG CAT TAT CTC; R, ACC TGC TAC CGC CAC TCT GT. β-actin: F, GAC TAC CTC ATG AAG ATC CT; R, CCA CAT CTG CTG GAA GGT GG (F: forward primer for sequence, R: reverse primer for sequence). The above primers were purchased from Mission Biotech Co., Ltd. (Taipei, Taiwan). The PCR products were separated by electrophoresis on 1.2% agarose gels and visualized by ethidium bromide staining under UV irradiation. The image of the resulting gel was captured and analyzed by ImageMaster VDS and ImageMaster 1D Elite software (Amersham Pharmacia Biotech, Sweden).

**Cytokine secretion assay**

After cultivation of the Caco-2 cells and B. lactis as mentioned above, the supernatants were collected for IL-1β, IL-10, TNF-α, and IFN-γ. The concentrations of IL-1β, IL-10, TNF-α, and IFN-γ were determined by ELISA kits (R&D Systems Inc., Minneapolis, MN, USA) according to the manufacturer’s instructions.

**Adhesion of B. lactis to Caco-2 cells by direct counting**

The light microscopic observation of B. lactis was carried out by fixing washed cells in 100% methanol for 30 min followed by Gram staining. B. lactis is a Gram-positive cell that remains purple in color.

**Statistical assay**

Numerical data are presented as means ± standard deviation. The data were analyzed by one-way analysis of variance (ANOVA) and followed by the least significant difference test using SPSS ver.10 (Chicago, IL) software.

**Result and discussion**

**LMF-, HS-Fucox-, and LMF + HS-Fucox-activated probiotic growth**

Reports have revealed that B. lactis, as a probiotic, is protective against immune and infectious diseases (28, 29). It is used in probiotic food, particularly fermented milk products intended for medicinal use. Intestinal dysbiosis has increasingly been observed in a variety of intestinal and systemic diseases, and maintaining an adequate bacteria profile may be a key point for healthy intestinal barrier function. Prebiotics are used to selectively stimulate the growth of bifidobacteria and increase the body’s natural resistance to invading pathogens (30). Hence, the effects of LMF, HS-Fucox, and LMF + HS-Fucox on B. lactis were examined. Our result revealed that LMF and LMF + HS-Fucox significantly promoted the growth of B. lactis at 50 µg/ml, whereas HS-Fucox showed significant promotion at 100 µg/ml. LMF and LMF + HS-Fucox had a better effect on the growth of B. lactis than HS-Fucox (Fig. 1). When the Caco-2 cells were cultured alone, their MTT absorbance value was 1.39 ± 0.03. When the Caco-2 cells were co-cultured with B. lactis bacteria, the MTT absorbance value was 1.36 ± 0.05. There were no significant differences between the two methods, and the co-culture showed no significant effect on cell growth (Fig. 1). Most prebiotics are water soluble or highly polar to stimulate the growth of probiotics, such as fucoidan (31) and laminarin derived from *Laminaria digitate* (32). Some reports demonstrated that oligosaccharides are the
most appropriate prebiotics to be used in effective synbiotic formulations (33, 34). LMF, an oligosaccharide hydrolyzed from original fucoidan, appeared as a prebiotic to \textit{B. lactis}. However, HS-Fucox stimulated the growth of \textit{B. lactis} slightly, possibly because that phenolic compound was probably unsuitable as a prebiotic (35), and its prebiotic activity might come from the polysaccharide coating.

\textbf{LMF, HS-Fucox, and LMF\textplus{}HS-Fucox against LPS-induced intestinal epithelial cell damage} 

LPS is produced by Gram-negative bacteria and can induce innate immune responses. The intestinal inflammation may serve as a nidus that can cause local and systemic organ dysfunction. In addition, LPS induced epithelial cell damage and mucosal hyperpermeability in vitro (36). First, we determined the effect of LMF, HS-Fucox, and LMF\textplus{}HS-Fucox on the protective functions of Caco-2 cells. The percentage of cell viability decreased to 67.91\% of the control medium in 1 \textmu g/ml LPS. HS-Fucox significantly inhibited the LPS-induced cell damage at concentrations as low as 50 \textmu g/ml, and LMF and LMF\textplus{}HS-Fucox significantly inhibited the LPS-induced cell damage at a higher concentration of 100 \textmu g/ml (Fig. 2). Sachindra et al. reported that the hydroxyl radical-scavenging activity of fucoxanthin was 13.5 times higher than that of \textalpha-tocopherol and showed the superiority of anti-inflammatory activity, such that HS-Fucox could inhibit the LPS-induced Caco-2 cell damage at a low concentration (37). Moreover, we were the first to use fucoxanthin on an intestinal inflammatory model. We also observed that LMF\textplus{}HS-Fucox (99.34 \pm 3.94\%) showed a stronger effect on increasing cell viability than LMF (94.56 \pm 3.26\%) and HS-Fucox (91.33 \pm 7.99\%) at 100 \textmu g/ml, a result that matched our previous studies showing that HS-Fucox may efficiently scavenge reactive-oxygen species (ROS) (38) and that LMF reduced inflammation through the inhibition of NF-\textkappa B (39). Under the co-culture of Caco-2 cells and samples, cell viability showed no significant effect with or without \textit{B. lactis} (Fig. 3), suggesting that \textit{B. lactis} maintained the homeostasis within the dynamic ecosystem in the human body and had no direct effect on the intestinal cells. Therefore, the new natural bioactive compounds that enhance the probiotic properties and intestinal barrier functions are of prime importance.

\textbf{LMF, HS-Fucox, and LMF\textplus{}HS-Fucox against LPS-induced destruction of intestinal epithelial barrier function} 

The intestinal barrier occurs coincident with increasing the enteral feeding and establishing normal intestinal bacterial colonization. The mechanisms involved in the development of intestinal barrier function are probably multifactorial. Under the co-culture of the Caco-2 cells and \textit{B. lactis} LPS-induced inflammatory system, we first determined the effect of samples on the protective functions of Caco-2 cell monolayers, and the integrity of polarized Caco-2 cell monolayers was determined by measuring the TER, which reflects the tightness of the tight junction between the epithelial cells (40). Through TER assay, it was shown that LPS destroyed the functions of Caco-2 cell monolayers, and the TER (percent of initial) was lower than that of the control group. However, when treating with 100 \textmu g/ml LMF, HS-Fucox, and LMF\textplus{}HS-Fucox, the TER continued to increase for an additional 8 h and remained constant from 4 to 8 h. Among them, LMF\textplus{}HS-Fucox significantly increased the TER (**\textit{p} < 0.01 when compared with LPS group alone) and was followed by LMF and HS-Fucox (*\textit{p} < 0.05 when compared with LPS group alone), indicating an enhancement of the intestinal epithelial barrier function (Fig. 4).
expression of claudin-1 and claudin-2 under H$_2$O$_2$ demonstrated that fucoidan can significantly increase the expressions were recovered (Fig. 5). Iraha et al. also measured by TER assay. The values were expressed as mean

The intestinal epithelial barrier function was induced by LPS. The intestinal epithelial barrier function was measured by TER assay. The values were expressed as mean

B. lactis Caco-2 cells and mRNA expression and its relative fold in the co-culture of Caco-2 cells and LMF, HS-Fucox, and LMF

As expected, LPS inhibited occludin, claudin-1, and claudin-2 mRNA expression. After treating with 100 µg/ml LMF, HS-Fucox, and LMF + HS-Fucox, the mRNA expressions were recovered (Fig. 5). Iraha et al. also demonstrated that fucoidan can significantly increase the expression of claudin-1 and claudin-2 under H$_2$O$_2$ disrupting epithelial barrier function (40). In addition to fucoidan, flavonoid from plant components can enhance barrier function by upregulating claudin-4 expression (43). Moreover, our data showed that the occludin, claudin-1, and claudin-2 expressions were significant greater in the LMF + HS-Fucox (**p < 0.01) group than in the LMF (*p < 0.05) and HS-Fucox (*p < 0.05) groups. This result matched the consequence of Fig. 4. So, it was suggested that LMF + HS-Fucox directly induced the expression of some tight junction proteins and might contribute to the enhancement of the epithelial barrier function. For comparison, by adding LMF with LPS treatment, the increasing trend of TER and tight junction protein expressions was only slightly higher than that of the HS-Fucox group. This result indicated that LMF and HS-Fucox exhibited a protective effect on epithelial cell injury, and HS-Fucox offered scavenging ROS activity (38) strong enough to make LMF + HS-Fucox an appropriate therapy for the treatment of inflammatory bowel diseases.

**LMF, HS-Fucox, and LMF + HS-Fucox modulated LPS-induced immune disorder**

The pro-inflammatory cytokines secreted by the epithelium, such as IL-1β and TNF-α, are hallmarks of inflammatory responses in the intestine. Relatively, the intestine also secretes anti-inflammatory cytokines, namely IL-10 and IFN-γ, that regulate cell inflammation (44). As shown in Fig. 6, LMF, HS-Fucox, and LMF + HS-Fucox were potent inhibitors of IL-1β and TNF-α and promoters of IL-10 and IFN-γ. It seems that LMF, HS-Fucox, and LMF + HS-Fucox were trying to balance the immune disorder under LPS-induced inflammation. To examine the main contributors to regulate the inflammation, LMF, HS-Fucox, and LMF + HS-Fucox were separately used. LMF (*p < 0.05) and HS-Fucox (*p < 0.05) exhibited similar effects on IL-1β, TNF-α, IL-10, and IFN-γ. However, when LMF and HS-Fucox (LMF + HS-Fucox group, **p < 0.01) are combined together, the anti-inflammatory activity became greater. Fucoidan and fucoxanthin both appeared to reduce the level of pro-inflammatory mediators, including IL-1β and TNF-α via the inhibition of NF-kB activation (39, 45). We suggested that LMF + HS-Fucox may provide comprehensive inhibition of the expressions of inflammatory cytokines by NF-kB pathway and probably possess anti-inflammatory properties for other immunity, via similar pathways in the epithelial cell (46). However, the mechanism still needs further study.

**LMF, HS-Fucox, and LMF + HS-Fucox enhanced B. lactis adhesion to intestinal epithelial cells**

Gram staining is a method of staining used to differentiate bacterial species into two large groups, Gram positive and Gram negative, and B. lactis, a kind of Gram-positive cell, appear in purple color. Observation of the Gram-stained cells under a light microscope showed that LPS treatment

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**Fig. 4.** Effects of 100 µg/ml LMF, HS-Fucox, and LMF + HS-Fucox on the intestinal epithelial barrier function in the co-culture of Caco-2 cells and B. lactis for 8 h, induced by LPS. Values were expressed as mean ± SD, n = 5. *p < 0.05, **p < 0.01

**Fig. 5.** Effects of 100 µg/ml LMF, HS-Fucox, and LMF + HS-Fucox on occludin, claudin-1, and claudin-2 mRNA expression and its relative fold in the co-culture of Caco-2 cells and B. lactis for 8 h, induced by LPS. Values were expressed as mean ± SD, n = 5.

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Citation: Food & Nutrition Research 2016, 60:32033 - http://dx.doi.org/10.3402/fnr.v60.32033 (page number not for citation purpose)
Fig. 6. Effects of 100 µg/ml LMF, HS-Fucox, and LMF + HS-Fucox on IL-1β, IL-10, TNF-α, and IFN-γ production in the co-culture of Caco-2 cells and B. lactis for 8 h, induced by LPS. Values were expressed as mean ± SD, n = 5. *p < 0.05, **p < 0.01 when compared with the LPS group alone.
reduced the \textit{B. lactis} counts of the Caco-2 cells the co-culture system, which appeared with a very slight blue/purple color (Fig. 7b). In contrast, a clear purple color appeared when treating with LMF, HS-Fucox, and LMF + HS-Fucox (Fig. 7c, d, and e). The LMF-treated group was inducing more \textit{B. lactis} adhered to Caco-2 cells than the HS-Fucox-treated group (Fig. 7c and d). This result is in agreement with the findings of Fig. 1, in which LMF appeared as a prebiotic to \textit{B. lactis}, and HS-Fucox was probably unsuitable as prebiotic. Furthermore, brown seaweed extracts have been proved to reduce the enterobacteriaceae populations and enhance the IL-6 and IL-8 cytokine expression to an ex vivo LPS challenge (32), and improve the probiotic properties of \textit{Lactobacillus plantarum} (47). The LMF + HS-Fucox-treated group showed stronger activity of \textit{B. lactis} adhesion than the results in the LMF- and HS-Fucox-treated groups, suggesting that LMF + HS-Fucox was important to the \textit{B. lactis} adhesion to Caco-2 cells, not only for their prebiotic effect but also because of many of their functional properties.

In conclusion, LMF and HS-Fucox were proved to enhance the functions of the immune system by inhibiting IL-1\(\beta\) and TNF-\(\alpha\) and promoting IL-10 and IFN-\(\gamma\), and revealed an anti-inflammatory effect in the intestinal cell line. The present findings suggested that LMF and HS-Fucox alone or in combination could be used as potential therapeutic agents in the treatment of intestinal inflammation.

\textbf{Fig. 7.} Gram staining for 100 \(\mu\)g/ml LMF, HS-Fucox, and LMF + HS-Fucox in the co-culture of Caco-2 cells and \textit{B. lactis} for 8 h, induced by LPS. \textit{B. lactis} is a Gram-positive cell that remains purple in color. (a) Control, (b) LPS, (c) LPS+LMF, (d) LPS+HS-Fucox, and (e) LPS+(LMF+HS-Fucox).
Acknowledgements

We thank the National Science Council (NSC) of the Executive Yuan, Taiwan [NSC 104-2320-B-034-003 (to Yen-Chang Lin); NSC 105-2320-B-034-001 (to Yen-Chang Lin)] for their support and the grant funding. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Conflict of interest and funding

Authors declare no conflict of interest in this study.

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