Amaranthus spinosus water extract directly stimulates proliferation of B lymphocytes in vitro

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Abstract

Amaranthus spinosus Linn. (thorny amaranth), a plant that grows in the wild fields of Taiwan, is extensively used in Chinese traditional medicine to treat diabetes. There have been no published studies on the immunological effects of A. spinosus. To determine whether A. spinosus has immuno-modulatory effects and clarify which types of immune effector cells are stimulated in vitro, we investigated the stimulatory effect of wild A. spinosus water extract (WASWE) on spleen cells from female BALB/c mice. We found that WASWE significantly stimulated splenocyte proliferation. However, isolated B lymphocytes, but not T lymphocytes, could be stimulated by WASWE in a dose response manner. After sequentially purifying WASWE, a novel immuno-stimulating protein (GF1) with a molecular weight of 313 kDa was obtained. The immuno-stimulating activity of the purified protein (GF1) was 309 times higher than that of WASWE. These results indicate that WASWE does indeed exhibit immuno-stimulating activity via directly stimulating B lymphocyte activation in vitro. Further, these results suggest that the immuno-stimulating effects of WASWE might lead to B lymphocyte activation and subsequent T cell proliferation in vitro. These results are potentially valuable for future nutraceutical and immuno-pharmacological use of WASWE or its purified fractions.

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Keywords: Amaranthus spinosus; Lymphocyte proliferation; Immuno-stimulating protein

1. Introduction

Increasing the body’s immunity in defending against diseases such as infections and tumors has become a major concern to many people in recent years [1–4]. More research has been focused on exploring the immune-enhancing effect of possible dietary substan-
The growth of herbal medicines or functional foods based on natural medicinal plants has gained rapid development in recent years around the globe. The functional products with specific health benefits from immunological activity have been the fastest-growing sector, moving swiftly into the mainstream. Recently, it was found that active thermally stable proteoglycans with molecular masses exceeding 100 kDa could be extracted from shark cartilage. These proteoglycans exhibit powerful immuno-stimulating activity in vitro [5]. *Ginkgo biloba* extract EGb 761 demonstrated immuno-stimulating activity on the nonspecific and humoral immune responses in a hypothalamic–pituitary–adrenal axis activation model in stressed rats [6]. Folk herb extracts from *Aeginetia indica* Roxb.extensively used in Thailand demonstrate a T cell stimulatory activity in vivo [7]. Neem (*Azadirachta indica*) leaf was found to have a prominent role in the murine Ehrlich carcinoma and B-16 melanoma cellular immune function, causing prophylactic growth inhibition of the tumor cells [8]. *Dioscoreae rhizoma* methanol extract (root of *Dioscorea tokoro* MAKINO) is traditionally used for the treatment of arthritis, muscular pain and urinary diseases in oriental medicine which significantly reduces pro-inflammatory cytokines and is a mediator in the synoviocytes of rheumatoid arthritis [9]. Chinese herbal medicine, Xiao-Qing-Long-Tang has confirmed the immuno-modulatory effect reducing bronchial inflammation in allergen-sensitized mice [10,11]. Most of the Chinese medicinal ingredients promote cellular or humoral immune responses by promoting lymphocyte proliferation and serum antibody titer [12]. However, herbal medicines and their ingredients or bioactive substances in foods, may exert differential immuno-modulation [13,14]. Searching for immuno-modulatory materials and characterizing the immune enhancement effects may have great potential in future practical applications for immuno-pharmacology. *Amaranthus spinosus* Linn. (thorny amaranth), a plant that grows in the wild fields of Taiwan, is extensively served as Chinese traditional medicine or wild vegetables to treat diabetes. However, there have been no published studies on the immunological effects of *A. spinosus* indigenous to Taiwan. This paper studies whether *A. spinosus* has immuno-modulatory effects and clarifies which types of immune effector cells are stimulated by wild *A. spinosus* water extract (WASWE).

It is difficult to wholly evaluate both the cellular and humoral immune responses in vivo. However, feasible in vitro assays to screen the lymphoproliferation activity of the possible daily food items and some of their active constituents are documented [5,15,16]. In this study, wild *A. spinosus* water extract (WASWE) was investigated for its stimulatory responses on immune effector cells using primary splenocyte culture from female BALB/c mice. To further characterize the possible immuno-modulatory components in WASWE, we also report on the isolation, basic physicochemical properties and immuno-stimulating activity of WASWE in vitro.

2. Materials and methods

2.1. Wild *A. spinosus* Linn. water extract (WASWE) preparation

*A. spinosus* leaves were weighed and chopped. After the same weight of deionized water was added, the sample was stirred in a Waring blender to extract the water-soluble constituent. The slurry was then centrifuged at 9000 × g (4 °C) for 30 min. A part of the supernatant was directly lyophilized to obtain a crude WASWE powder and the remainder of the supernatant was heated at 100 °C for 30 min and then lyophilized to harvest the heat-treated WASWE powder. The yield of crude extract was calculated by the recovery of lyophilized powder and expressed as grams of WASWE/100 g of fresh vegetables.

2.2. WASWE chromatographic purification

2.2.1. DEAE sephacel

Ion exchange: DEAE sephacel (Pharmacia Biotech) medium was used to purify crude WASWE. After applying the crude sample (WASWE), the column (1.5 cm×12 cm, Econo-Pac column, Bio-Rad Laboratories) was eluted at a flow rate of 1.0 ml/min with 2 bed volumes of deionized water, 0.1 M NaCl, 0.3 M NaCl, 0.5 M NaCl, 1.0 M NaCl, 2.0 M NaCl, 1.0 N NaOH (washing column), and 1.0 N HCl (washing column), respectively. The eluent was collected using a fraction collector (Gilson, model 203) and detected by absorbance at 280 nm. The material was resolved into seven major peaks:
IF1~IF7. Each fraction was lyophilized and analyzed for immuno-stimulating activity. The yields of ion exchange fractions (IF1~IF7) were calculated by the recovery of lyophilized powder and expressed as grams of IF1~IF7/100 g of WASWE.

2.2.2. Sepharose CL-6B
The starting material (active fraction eluted from ion exchange column) was applied in a Sepharose CL-6B column (Pharmacia Biotech) to further purify the active ion exchange fraction. The column (2.6 cm×100 cm) was eluted with deionized water at a flow rate of 1.3 ml/min. Each tube collected about 240 drops of eluent. Each fraction was lyophilized and tested for the stimulation index of cell proliferation. The molecular weight of each fraction was calibrated with a standard molecular weight kit. The yields of gel filtration fractions (GF1~GF2) were calculated by the recovery of lyophilized powder and expressed as grams of GF1~GF2/100 g of IF4.

2.3. Protein content analysis
The protein content of the purified sample was analyzed using the BCA (bicinchoninic acid) protein assay kit (product No. 23225, Pierce), according to the accompanying instructions, using a 96-well microtiter plate. Briefly, aliquots of 25 µl of each standards (bovine serum albumin, BSA, product No. 23209, Pierce) or unknown samples were pipetted into the appropriate microwell plate wells (Nunc). Aliquots of 200 µl of the working reagent (containing sodium carbonate, sodium bicarbonate, bicinchoninic acid, sodium tartrate in 0.1 M sodium hydroxide and cupric sulfate) were added to each well and mixed the plate well on a plate shaker for 0.5 min. Then, the plate was covered and incubated at 37 ºC for 30 min. After incubation, the plate was cooled to room temperature and measured the absorbance at or near 562 nm on a plate reader (ELISA reader, ASYS Hitech, Austria). Using the standard curve, the protein concentration for each unknown sample was determined.

2.4. In vitro biological assessment

2.4.1. Experimental animals for primary splenocytes
BALB/c mice (female, adult) were purchased from the Laboratory Animal Center, College of Medicine, National Taiwan University and maintained in the Department of Biochemical Science and Technology, College of Life Science, National Taiwan University. The mice were housed and fed a standard Lab diet (normal chow diet) individually. The animal room was kept on a 12-h-light and 12-h-dark cycle. Constant temperature (25±2 ºC) and humidity were maintained. The animals (8- to 10-week old) were sacrificed using CO₂ inhalation to obtain spleens. The abdominal cavities were opened aseptically and the spleens were removed.

2.4.2. Splenocyte proliferation stimulation
The spleen cell proliferation response was assayed as described previously [17,18]. Briefly, spleens were removed aseptically from BALB/c mice. Single spleen cells were prepared by lysing the red blood cells. Splenocytes were isolated from each animal and adjusted to 1×10⁷ cells/ml in TCM medium. Splenocytes (50 µl/well) without or with mitogens (50 µg/ml, respectively), IF1~IF7 (10, 40, 200, 1000, 2500, 5000 µg/ml, respectively) or GF1~GF2 (1.0, 5.0, 10.0, 20.0 µg/ml in TCM medium, respectively) were plated in 96-well microtiter plates. The plates were incubated at 37 ºC in a humidified incubator with 5% CO₂ and 95% air for 48 h. The cells were then pulsed with [3H]-thymidine (1 µCi/well in 10 µl TCM medium). After 24 h, the cells were harvested using a semi automated sample cell harvester (FILTERMATE 196 PACKARD). The radioactivity was determined using a Direct Beta Counter (MATRIX 96, PACKARD). The stimulation index (S.I.) was expressed as S.I.=(mitogencpm−controlcpm)/(controlcpm−blankcpm). Each value was presented as mean±S.E. (n=6).

2.4.3. T and B cell specific proliferation responses in total splenocytes from BALB/c mice
To further define the lymphocyte population stimulated by WASWE, bromodeoxyuridine (BrdU) cell incorporation was performed and analyzed with FACS (flow cytometry) analysis [19,20]. After the spleen cells were stimulated by WASWE (1250 µg/ml), LPS (10 µg/ml) and PHA (10 µg/ml), respectively, for 48 h, 10 µM bromodeoxyuridine (BrdU; Becton Dickinson)
was added and the cells incubated for another 24 h. The cells were then fixed with 1% paraformaldehyde and split into several portions to clarify the proliferation percentage by cell surface markers on specific cell. Parts of the cells were stained with fluorescein-isothiocyanate (FITC)-conjugated anti-BrdU antibody and phycoerythin (PE)-conjugated anti-B220 antibody to detect the proliferation of B cells. The rest of the cells were stained with FITC-conjugated anti-BrdU antibody and PE-conjugated anti-CD3 antibody to detect the proliferation of T cells. After that, the harvested cells were subjected to flow cytometry (FACS) analysis (Becton Deckinson), respectively.

2.4.4. T and B cell isolation from splenocytes

To further confirm which types of lymphocytes were directly stimulated by WASWE, T and B cells were isolated from the spleen cells of female BALB/c mice [19–21]. T cell isolation is based on its less binding activity with nylon wool than that of other cells. T cells (nylon wool nonadherent cells) were separated by passing the spleen cells through a nylon wool column packed in a 10 ml syringe. After incubation of total splenocytes in the column at 37 °C for 1 h, T cells were collected by eluting the column with 10 ml TCM medium. B cells (T depleted spleen cells) were isolated by complement-mediated T cell lysis using anti-mouse Thy1.2 antibody plus rabbit complement (CEDARLANE Laboratory) treatment. The purity of the isolated T and B cells were further analyzed by FACS analysis. Total splenocytes, isolated B cells, and isolated T cells (original cell density: $2 \times 10^6$ cells/ml) were treated with WASWE (final concentration: 1250, 2500 µg/ml) and quantified with the [3H]-thymidine incorporation method to evaluate their proliferation rates as described above. The isolated T cells were also incubated with γ-irradiated (2000 rad) antigen-presenting cells (APC, syngeneic splenocytes).

2.5. Statistical analysis

Data were analyzed by the Windows SAS program (Version 6.12). Data are expressed as mean±S.E. and analyzed statistically using ANOVA followed, if justified by the statistical probability ($P<0.05$), by Duncan’s New Multiple Range test. Differences were considered statistically significant if $P<0.05$.

3. Results

3.1. WASWE effect on cell proliferation of total splenocytes from female BALB/c mice

Splenocyte proliferation from inbred BALB/c mice species cultured in the presence of WASWE was evaluated. The highest stimulation activity was found at the concentration of 1250 µg WASWE/ml (Fig. 1). It was found that WASWE significantly stimulated splenocyte proliferation in a dose response manner. However, WASWE lost its immuno-stimulating activity after it was heated at 100 °C for 30 min (data not shown). To clarify which types of immune effector cells in the spleen were being stimulated by WASWE, the proliferation of lymphocyte subset in total splenocyte culture was studied. The CD3+ and B220+ surface markers on proliferation cells of total splenocyte culture were detected by FACS. Both B (B220+) and T cells (CD3+) in total splenocyte culture were stimulated simultaneously by WASWE at the concentration of 1250 µg/ml (Fig. 2). However, the proliferation rate of B cells (14.85±2.44%) exceeded that for T cells (7.49±3.12%) at the indicated WASWE concentration on the third incubation day. Stimulatory activities by LPS (a B cell mitogen) and PHA (a T cell mitogen) to both B and T cell subsets were higher than that for WASWE. B and T cell stimulation in the presence of LPS revealed significant differences from the Sham (control). B cell stimulation in the presence of WASWE revealed significant differences from the Sham (control). Even though WASWE demonstrated immuno-stimulating activity to total splenocytes including both B and T cells, it is not clear whether WASWE stimulated lymphocyte proliferation of both B and T cells [20].

3.2. WASWE effect on purified populations of B cells and T cells obtained from total splenocytes

To avoid secondary B and T lymphocyte signals being disturbed by each other on the proliferation rate, B and T cells in the total splenocytes were isolated. The cellular purity of B and T cells was assayed by FACS analysis. It was found that the cell constitutes of splenocytes, isolated B and T cells, contained 41.54% B cells and 47.11% T cells, 82.34% B cells and 4.15% T cells, and 88.85% T cells and 2.89% B cells,
Fig. 1. Effects of WASWE on proliferation of total splenocytes from BALB/c mice. Data are assayed using ANOVA. Each bar represents the mean value±S.E. obtained from six separated mice. Within each bars graph, bars not sharing a letter are significantly different (\( P<0.05 \)) according to Duncan’s New Multiple Range test.

Fig. 2. Effects of WASWE, LPS (a B cell mitogen), and PHA (a T cell mitogen) on proliferation of separated populations (B cells or T cells) according to the surface marker distribution on cell proliferation in the entire splenocyte culture from BALB/c mice detected with FACS at the third day of incubation. Original cell density was \( 5.0 \times 10^6 \) cells/ml. Cell proliferation percentages were calculated on the basis of the number of lymphocytes found in each quadrant. Data are assayed using ANOVA. Each bar represents the mean value±S.E. obtained from three separated mice. Within each bars graph of separated populations (B cells or T cells), bars not sharing a letter are significantly different (\( P<0.05 \)) according to Duncan’s New Multiple Range test.
respectively (Fig. 3). The cellular purity of the individual subpopulations in the purified B and T cell populations increased by about 2 folds. The total number of splenocytes, isolated B and T cells (cell density: $2.0 \times 10^6$ cells/ml), respectively, were further exposed to different concentrations of WASWE for 72 h. We determined the stimulation index (S.I.) for the total (unpurified) splenocytes, isolated (purified) B and (purified) T cells and they were 6.00, 2.08, and 0.99 at a concentration of 1250 µg/ml WASWE and 2.50, 4.85, and $-0.41$, respectively, at a concentration of 2500 µg/ml WASWE. The B and T cell subpopulations responded differently to stimulation by WASWE. Higher WASWE concentrations (2500 µg/ml) caused a cytotoxic effect in purified T cells. Purified B cells, but not T cells, could be stimulated with WASWE in a dose-dependent manner. The splenocytes had a much higher proliferation rate than that of individual purified B or T cells at the concentration of 1250 µg/ml WASWE, suggesting some interactions occurred between these two cells. These results suggest that WASWE exerted a direct stimulatory effect on B cells. T cells seemed to have a co-stimulatory effect on B cells. T cells in bulk splenocytes might be activated via secondary signaling from stimulated B cells, leading to T cell proliferation. Since LPS and PHA are powerful B and T lymphocyte stimulants, they were chosen as positive controls for lymphocyte proliferation in this experiment. We found that the LPS (5 µg/ml) stimulation index on isolated B and T cells were 573.83 and 3.47, respectively. The PHA (10 µg/ml) stimulation index on isolated B and T cells were 19.63 and 72.09, respectively. These results were in accordance with our prediction. It was also verified that isolated lymphocyte cultures in this experimental model were workable and believable. Stimulation index of total splenocytes stimulated by LPS or PHA was higher than that of individual purified T or B cells, suggesting some interdependence between these two cells.

3.3. Purification of immuno-stimulating components in WASWE

3.3.1. Ion exchange chromatography

To exploit the bioactive substances that stimulate splenocyte proliferation in vitro, WASWE was first partially purified using DEAE sephacel ion exchange. Seven fractions (IF1~IF7) were obtained (Fig. 4) and conducted to test their immuno-stimulating activity. The tests demonstrated that the fourth fraction, IF4, was the active fraction that stimulated splenocyte proliferation (Fig. 5). Fig. 5 shows that crude
WASWE and IF4 indeed demonstrated immuno-stimulating activity on splenocyte cultures at the 1250 μg/ml WASWE and 5 × 10⁶ cells/ml concentration. The radioactivity uptake (cpm) by the total splenocytes stimulated by crude WASWE and IF4 was 7500 ± 335 and 12,902 ± 382 respectively.

3.3.2. Gel filtration chromatography

IF4 was further purified using a gel filtration column packed with Sepharose CL-6B. Gel-filtrated chromatogram of IF4 was consisted of two major fractions, GF1 and GF2 (Fig. 6a). As compared to the gel-filtrated chromatograms of gel filtration standard,

![Fig. 4. Chromatogram of WASWE by step gradient ion exchange chromatography using DEAE sephacel media. Column size: 1.5 cm×12 cm (20 ml). Elution volume of each gradient exchange was 40 ml at a flow rate of 1.0 ml/min.](image)

![Fig. 5. Effects of different WASWE preparations on total splenocyte proliferation from BALB/c mice. Original cell density was 5 × 10⁶ cells/ml. IF means fractions obtained from ion exchange chromatography with DEAE sephacel. Data are assayed using ANOVA. The results are expressed as the mean value ± S.E. obtained from six separated mice.](image)
molecular weight of GF1 and GF2 were 313 kDa and 1.6 kDa, respectively (Fig. 6b). It was found that GF1 was the major fraction attributing to immuno-stimulating response. The stimulation index (S.I.) for GF1 in the range of 0.5~5.0 \( \mu \)g/ml showed a dose response effect on BALB/c splenocytes (Fig. 7). The S.I. to BALB/c mice stimulated using GF1 at the concentration of 5.0 \( \mu \)g/ml was 4.81±0.07, while S.I. stimulated by crude WASWE at the concentration of 1250 \( \mu \)g/ml was 3.94±0.18 (Fig. 1). The WASWE’s immuno-stimulating activity increased more than 300 times after it was purified from crude-extraction to using gel filtration treatment.

3.3.3. Bioactive substances with immuno-stimulation in WASWE characterization

Table 2 summarized the stimulation index, purification fold, protein content and WASWE yield produced by different purification processes. The S.I. of the crude extract (WASWE), IF4 and GF1 to
splenocytes at the same corresponding concentration of 1250 µg/ml were 3.9, 6.8, and 1203, respectively. The stimulatory activity increased 309 times after WASWE was purified with gel filtration. The protein content of GF1 was 90.8%. Yields of crude extract, ion exchange and gel filtration chromatography were calculated by the recovery of each individual step of the purification process and expressed as recovery in grams of powder per 100 grams of fresh vegetables, per 100 grams of WASWE and per 100 grams of IF4, respectively. The yields of WEWAS, IF4 and GF1 were 4.7%, 1.9% and 24.9%, respectively. The yields seemed variable in different purification processes.

4. Discussion

There has been a rapid development in the use of herbal medicines in recent years around the globe. Many researches have focused on exploring the immuno-stimulating actions of certain foods and dietary herbs. It has been found that Lactobacillus delbrueckii subsp. bulgaricus and Lactobacillus acidophilus mitogenically induced antibody production and spleen cell proliferation [22]. Aqueous extracts of Crinum latifolium and Camellia sinensis, Chinese and Vietnamese traditional medicines, show immunomodulatory properties on human peripheral blood mononuclear cells [23]. In our early studies, we found that dehulled adlay, a food and natural Chinese medicine, demonstrated antiallergic activities [24]. However, some Cruciferous and Brassica vegetables inhibited tumor proliferation and induced apoptosis [25,26]. These plants seemed to have adverse effects on cell-mediated and humoral immunity. We investigated wild A. spinosus which is served as vegetables or herbs in Taiwan, for its immuno-modulatory actions using primary spleen cell cultures from BALB/c mice. There is no known report on immuno-modulation concerning A. spinosus. Our results provide evidence that wild A. spinosus water extract (WASWE) demonstrates significant (P<0.01) immuno-stimulating activity on primary splenocytes from female BALB/c mice (Fig. 1).

Knowledge of which specific cell subpopulations were being stimulated was very important for future use in immuno-pharmacology. Our results indicate that WASWE exerts stimulatory effects on both B and T cells (Fig. 2). However, bulk splenocytes stimulated...
by an adequate concentration (1250 μg/ml) of WASWE exhibited a much higher proliferation rate than that of isolated purified B and T cells (Table 1). This suggests some interactions between these two cells. We assumed that T cells in bulk splenocytes are activated via secondary signaling from stimulated B cells, leading to the T cell proliferation. It has been found that B lymphocytes may act as antigen-presenting cells for CD4+ T cell priming if B cell and T cell coexist [27]. Resting B lymphocytes were also found to act as APC for naive T lymphocytes via CD40 ligand/CD40 interaction [21]. When it is activated, T helper cell-related cytokines may interplay with B cells and direct their differentiations [28]. B cell interactions with activated T cells reduce the threshold for CD40-mediated B cell activation [29]. We provide evidence that WASWE directly stimulates the proliferation of B lymphocytes and suggest the possibility of activated B cells subsequently delivering a second signal to activate T cells. Actually we had found that WASWE stimulated both Th1 and Th2 cytokines production in vitro by splenocytes from female BALB/c mice, however the secretion of Th2 cytokines (IL-5, IL-6, IL-10) was much higher than that of Th1 cytokines (IL-2, IFN-γ) (data not shown). From the profiles of cell proliferation and cytokine production stimulated by WASWE in the splenocyte culture, this presumably occurs that WASWE first activates B cells and the activated B cells subsequently activate T cells especially Th2 cells. After they are activated, both Th1 and Th2 cells, especially Th2 cells (cytokines), can reversely contribute to humoral immunity by inducing B cells to proliferate, differentiate or produce antibodies. However, the extracts from Aeginetia indica Roxburt demonstrate a T cell stimulatory activity in vivo [7]. Our results suggest that different immuno-modulatory principles might exert diverse mechanisms on both humoral and cellular immune responses. Certain herbal medicinal ingredients affect the peripheral lymphocyte proliferation and serum antibody titer [12], so that WASWE bioactive ingredients may have potential for use as a component in new immuno-potentiator drugs.

It is necessary to unravel the basic characterization of the immuno-stimulating constituents in WASWE for future applications in dietary supplements or advanced clinical practice tests. In our preliminary studies on A. spinosus, we found that heat-treated WASWE lost its immuno-stimulating activity, while activated charcoal-treated WASWE increased immuno-stimulating activity (data not shown). The preliminary results indicated that the bioactive constituent in WASWE might be a proteinaceous component with higher molecular weight. Activated charcoal adsorbs colorants (like flavonoids) and some lower molecular weight substances. Most proteinaceous substances are heat-labile. In this study, we found a novel immuno-stimulating substance in WASWE with a molecular weight of 313 kDa (Fig. 6b) containing 90.8% (Table 2) protein. The immuno-stimulating activity of this novel protein increased 309 times as

### Table 1
Stimulation index (S.I.) of WASWE to different lymphocyte proliferation isolated from the spleen of female BALB/c mice

<table>
<thead>
<tr>
<th></th>
<th>Total splenocytes (B cell: 41.54%; T cell: 47.11%)</th>
<th>Isolated B cell (B cell: 82.34%)</th>
<th>Isolated T cell (T cell: 88.85%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WASWE (1250 μg/ml)</td>
<td>6.00</td>
<td>2.08</td>
<td>0.99</td>
</tr>
<tr>
<td>WASWE (2500 μg/ml)</td>
<td>2.50</td>
<td>4.85</td>
<td>−0.41</td>
</tr>
<tr>
<td>LPS (5 μg/ml)</td>
<td>154.93</td>
<td>573.83</td>
<td>3.47</td>
</tr>
<tr>
<td>PHA (10 μg/ml)</td>
<td>344.43</td>
<td>19.63</td>
<td>72.09</td>
</tr>
</tbody>
</table>

1. Stimulation index=(sample cpm−blank cpm)/(control cpm−blank cpm).
2. All values means of triplicates.
3. Original cell density: 2.0×10⁶ cells/ml.
4. Total splenocytes were directly obtained from female BALB/c mice. Isolated B cells were obtained after they were treated using the complement lysis method. Isolated T cells were enriched in a nylon wool column.

### Table 2
Changes in stimulation index, stimulatory activity, protein content and WASWE yield during different purification processes

<table>
<thead>
<tr>
<th>Purified process</th>
<th>Crude extract</th>
<th>Ion exchange (IF4)</th>
<th>Gel filtration (GF1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stimulation indexᵃ</td>
<td>3.9</td>
<td>6.8</td>
<td>1203</td>
</tr>
<tr>
<td>Stimulatory activity</td>
<td>1.0</td>
<td>1.7</td>
<td>309</td>
</tr>
<tr>
<td>Protein contentᵇ (%)</td>
<td>79.0</td>
<td>10.4</td>
<td>90.8</td>
</tr>
<tr>
<td>Yieldᶜ (%)</td>
<td>4.7</td>
<td>1.9</td>
<td>24.9</td>
</tr>
</tbody>
</table>

ᵃ Calculated by the maximum splenocyte proliferation rate of BALB/c mice at the same corresponding concentration of 1250 μg.
b The protein content of lyophilized powder was assayed by the BCA protein kit.
c Calculated by the recovery of each individual step of the purification process.
compared to that of crude WASWE. As we indicated in the part of materials and methods, the protein content was determined by the BCA method in this study. The results from the BCA method seem protein-to-protein variation and are interfered by the constituents, such as reducing sugars and reducing agents, thiols, copper chelating agents, ascorbic acid and uric acid, tyrosine, cysteine and tryptophan. Using bovine serum albumin (BSA) or immunoglobulin (IgG), especially BSA, as standards works well with most protein assay methods. To minimize the confounding of protein type, the BSA was used as a standard protein for quantitation of total protein content in this study. Besides, most interfering substances against protein determination are small molecules. They might not be obtained from gel filtration column in our experimental condition. To further minimize the confounding of water content, the samples used for protein determination were lyophilized to remove water in this study. Because the other constitutes of GF1 should not disturb the protein determination, we assume that the BCA method is a suitable and believable method for protein determination, we assume that the BCA method has also been referred by other reports [5,30]. This study shows that the protein content of GF1 is 90.8%. Most proteins extracted from plants are finally proved to be a glycoprotein. We assume that GF1 is a glycoprotein and the heat-labile property of GF1 might be attributed to its high protein content. The active principles extracted from shark cartilage are thermally stable proteoglycans with molecular masses exceeding 100 kDa. They are potent B cell stimulators but they do not affect T cells [5]. However, basic properties between the novel proteins from WASWE and from shark cartilage are quiet different. The stability of proteoglycans is a reflection of the fact that the protein core is shielded by surrounding glycosaminoglycan chains but this is not the case of GF1. More information on the basic properties concerning the immuno-stimulating protein in WASWE remains to be elucidated.

This study showed that a novel protein with a molecular weight of 313 kDa displayed strong immuno-stimulating activity, and could directly activate primary B cell proliferation. This is a potentially valuable substance for nutraceutical or immunopharmacological use.

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