Immunomodulation Effect of *Nypa fruticans* Palm Vinegar†

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Abstract

Vinegars are aqueous products of fermentation that are composed of acetic acid, water and other trace chemical components. Previous studies have shown that they have several beneficial effects for humans, such as anti-diabetic, anti-bacteria, antioxidant, anti-tumor or the enhancement of immune function. *Nypa fruticans* palm vinegar (NPV) is 1 of the vinegars that is usually used as a food ingredient. However, there are limited studies that have provided results to support its proposed immunomodulatory properties. This study investigated the immunomodulatory effects of NPV. Locally produced NPV from 3 different planting areas (fresh, brackish and saline water) of the Nakhon Si Thammarat province in Thailand were administered to rats, and the enhancement of the in vitro phagocytic activity was assessed. The results showed that NPV potently enhanced antibody production and the phagocytic activity. Moreover, NPV-treated rats showed a reduction of edema in type 4 hypersensitivity reactions. The results indicated that NPV had immunomodulatory effects, and therefore, NPV could be considered a functional food.

Keywords: *Nypa fruticans*, Nipa palm vinegar, NPV, Immunomodulatory effect, Functional food

Introduction

Vinegar is normally used in a variety of food ingredients. Several kinds of fruits or grains are used as the primary starting materials for vinegar production [1]. The natural vinegar product, aside from being used as part of food ingredients, has long been considered. Recently, the potential health benefits of vinegar varieties due to its functional properties and bioactive substances have been reported by many studies [2-4].

*Nypa fruticans* palm is abundant and found along the coasts and rivers from Bangladesh to the Pacific islands. It can grow in soft mud-like mangrove areas with fresh, brackish and saline water. Most tapped palm trees give a of nipa sap very rich in sugar 10 % to 20 % according to species and individual variation. In Asia, the sap is used either as fresh juice or processed into many product types, e.g. wine, alcohol, araq, sugar and vinegar, etc. [5]. Nipa sap contains sucrose, protein, and minerals, including calcium, phosphorous, iron, copper and vitamins [6]. Moreover, there contain several non-essential amino acids, glycine, glutamic acid, alanine, proline, tyrosine and essential amino acids: methionine and leucine [7].

Traditional processes for the production of *Nypa fruticans* palm vinegar (NPV) involve a 2-stage fermentation process in which alcohol is initially formed by fermentation process of nipa palm sap with yeast and mold, involving *Saccharomyces*, *Amylomyces*, *Aspergillus* and *Candida*. The fermentation processes change carbohydrates to alcohols and carbon dioxide, and then, the alcohol is changed to acetic acid by acetic acid bacteria, i.e. *Acetobacter aceti*. NPV has been previously used as a traditional

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medicine because it is believed to have many health benefits. However, there are few studies supporting such assumption. The study of Yusoff et al. [8] reported that NPV could decrease blood glucose levels in mice with induced diabetes mellitus; antioxidant activities were observed in this study.

More empirical studies have identified the immunomodulatory activities of different types of vinegars. Korean persimmon vinegar, for instance, has been confirmed to enhance antibody production [9]. Another study demonstrated that the administration of apple vinegar can potentially improve the expression of immune substances [10]. However, as noted, there are still limited reports on the immunomodulatory effect of NPV. Thus, this study aims to investigate the immunomodulatory effect of NPV from planting areas with different salinities by both in vitro and in vivo analysis.

Materials and methods

Nypa fruticans palm vinegar (NPV)

Nypa fruticans palm vinegar (NPV) samples were selected from locally produced NPV from the Kanapnak sub-district, Pak Phanang district, Nakhon Si Thammarat province, Thailand. The samples were collected from 3 different salinity planning areas, including of fresh, brackish and saline. The electrical conductivity (EC dS/m at 25 °C) was determined in 1:5 soil extract using a conductivity meter and the results were compared according to the soil salinity classification of U.S. Salinity Lab. Staff [11,12].

Animal model

Male Wistar rats (250 - 300 g), which were aged 6 - 8 weeks and obtained from the Nomura Siam International Co., Ltd., were used in this study. All animals were housed and acclimatized in a well-ventilated animal transit room (12:12 h light-dark cycle) at the animal research building, Walailak University, Thailand. Throughout the experiment, the rats were fed food and water by standard methods. The study was authorized for the use of animals in scientific research by the Animal Ethics Committee, Walailak University. The authorization number for the use of animals in scientific research is 006-2017.

Enhancement of the phagocytic activity

One millilitre of 5×10⁵ RAW 264.7 macrophage cells was cultured in a 6-well plate with complete Dulbecco's Modified Eagle Medium (DMEM) (supplemented with 10 % foetal bovine serum, 2 % of 100× penicillin/streptomycin and 1 % of 100× amphotericin B). The plate was incubated for 24 h at 37 °C in 5 % CO₂. Then, 200 μl NPV with final concentrations of 100, 50, 25, 12.5, 6.25, 3.125, 1.56, and 0 μl/mL was added and incubated for 24 h. Cells were scraped, and rabbit IgG-FITC conjugated latex beads (Cayman, USA) were added directly into pre-warmed cultured medium to a final dilution of 1:400 and incubated at 37 °C for 1 h. The RAW 264.7 cells were centrifuged at 400×g for 5 min. The supernatant was decanted, and the cell pellet was re-suspended in 500 µl of assay buffer. The percentage of phagocytic activity was analysed by flow cytometry (BD FACSCalibur™) [13].

Evaluation of the cellular immune response

Fifty-five rats were used for this study. They were divided into 11 groups of five rats each (2 control groups and 9 test groups). The negative control group was fed water, while the positive control group was fed 5 mg/kg cyclosporine A on day 14. Test groups were orally administered NPV from different salinity planting areas including fresh, brackish and saline water. NPV was orally administered to rats in 3 different concentrations (1:1, 1:5 and 1:10) with a dosage of 10 mL/kg body weight/day for 21 days. Rats were intraperitoneally injected with 0.1 mL of 20 % sheep red blood cells (SRBC) in sterile normal saline on days 7 and 14. On day 21, the rats were administered 0.03 mL of 1 % SRBC into the sub-plantar region of the right hind paw, and at 24 h after injection, the edema was assessed using a digital plethysmometer [14]. The results were compared between each group.
Evaluation of the humoral immune response

Wistar rats were divided into 11 groups and orally administered the treatment at different doses to assess the cellular immune effect. However, rats were treated for 14 days, and the positive control group was orally administered 5 mg/kg cyclosporine A on day 7. All rats were subjected to the tail vein blood collection method on day 7 for the determination of the primary antibody response. After blood collection on the same day, 20% SRBC was peritoneally injected. On day 14, tail vein blood was collected for determining the secondary antibody response. Antibody titres against SRBC were measured by the microtitration technique and calculated as the natural log [15]. The results were compared between each group.

Assessment of the spleen index

Wistar rats were divided into 11 groups and treated in the same manner for cellular and humoral immune effect assessment. However, the positive control group in this experiment was orally administered 5 mg/kg cyclosporine A on day 23. At the end of the experiment (day 30), the rats were sacrificed, and the spleens were subjected to weighing. The spleen index was calculated by the following formula for cytotoxicity and immune stimulation assessment [16]:

\[ \text{Spleen index} = \left( \frac{\text{spleen weight (mg)}}{\text{body weight (g)}} \right) \times 100. \]

Statistical analysis

The values are expressed as the mean ± SEM (standard error of the mean). The results were analysed by 1-way analysis of variance (ANOVA), followed by an independent T-test to determine the statistical significance. The level of significance was set at \( p < 0.05 \).

Results and discussion

Previous HPLC analysis of the aqueous extract NPV revealed a mixture of organic acids, mainly lactic acid and acetic acid at percentages of 10.35 and 8.68%, respectively [17]. The anti-inflammatory mechanism showed significant lower expression of liver iNOS (inducible nitric oxide synthase) and NFκB transcriptional factor [18]. From previous and recent studies, there is enough evidence that supported NPV as a functional food. Moreover, the significant points of our present study can remove the controversy surrounding functional foods by using an in vivo animal model and in vitro experiments. Many studies have been demonstrating that a functional food supports the concept that food is not only necessary for living but also a source of mental and physical well-being, contributing to the prevention and reduction of risk factors for several diseases or enhancing certain physiological functions. In reference to the immune system, many studies have indicated that not only probiotics but also single micronutrients incorporated into functional foods contribute to an enhancement of immunocompetence. The effective food components that have been demonstrated to have immunomodulatory effects were probiotics [19-21], trace elements [22,23] and dietary antioxidants [24].

Effect on the phagocytic activity

To test the effect of NPV on the phagocytic activity, which is an important mechanism of innate immunity, NPV from 3 different salinity planting areas was used to directly treat RAW 264.7 macrophage cells, followed by the addition of rabbit IgG-FITC conjugated latex beads. The percentage of active phagocytic cells was determined by flow cytometry. The results showed that the phagocytic activity of macrophage cells that were treated with NPV from the saline planting areas showed significant increases with all of the NPV concentrations when compared to negative untreated cells, while only high concentrations (25 and 12.5 µl/mL) of NPV from the fresh water planting area showed a significant increase in the phagocytic activity. The results of NPV from both saline and fresh water planting areas showed dose-dependent enhancement of the phagocytic activity of macrophage cells. For NPV from brackish water planting areas, only low concentrations (3.125 and 1.56 µl/mL) of NPV-treated cells showed significantly enhanced phagocytic activity, which might be due to the different toxic ingredients.
from different NPV sources. However, the results indicated that NPV could enhance the phagocytic activity of macrophages cells, as shown in Figure 1.

![Figure 1](image_url)

**Figure 1** Percentage of phagocytic activity of macrophage cells treated with different concentrations of NPV from 3 different salinity planting areas. The bars show the mean ± SEM. *; significantly different when compared to the negative control.

Phagocytosis by macrophages is an important nonspecific host-defence (innate immunity) mechanism that occurs immediately after exposure to a foreign antigen. It is also used as an important indicator of the activation of immune function in several studies [25,26]. Our results suggest that NPV can increase the phagocytic potential in macrophages. The phagocytic stimulation activity of NPV has a similar favourable effect to that of most apple cider vinegars (ACVs) that was previously demonstrated by Yagnik and colleagues [27]. The phagocytic activity enhancement might possibly be from the organic acids that are the major constituents of NPV, which is supported by a previous study that demonstrated poultry fed a basal diet supplemented with organic acids had significantly greater phagocytic activity [28].

**Cellular immune response modulation by delay-type hypersensitivity inhibition**

To assess the delay-type hypersensitivity (DTH) that indicates the effect on the cellular response, rats were orally administered 3 different concentrations of NPV obtained from 3 different growing areas in parallel with control rats. On day 21, SRBC was subcutaneously injected into the right hind foot paw, and the edema was evaluated after 24 h. Most of the NPV-treated rats showed significantly lower paw edema when compared with negative untreated rats, except for the rats treated with low doses of NPV from fresh water planting areas, which showed no significant difference. This might be due to the effect
of different concentrations of active ingredients in NPV from different sources. These modulation effects of NPV were equivalent to cyclosporine A-treated rats, as shown in Figure 2.

Delayed type hypersensitivity (DTH) reaction is a protective localized cell-mediated immune-inflammatory response. It is a mechanism primarily against intracellular pathogens. Upon antigen presentation, T-lymphocytes may become sensitized lymphocytes and will generate a regional abnormal reactive inflammation. This inflammation type is delayed and characterized by cell degeneration and necrosis [29].

Figure 2 Edema in the right hind paw 24 h after SRBC injection. The rats were orally administered NPV from different dilutions (1:1, 1:5 and 1:10) and different salinity planting areas (fresh, brackish and sea water). The bars show the mean ± SEM. *; significantly different when compared to the negative control.

The oral administration of NPV in rats showed significantly suppressed hind paw edema compared to the negative control rats. Hind paw edema indicates the strength of the delayed allergic response (DTH), which is mediated by T-cells involved in cellular immunity [30,31]. This inflammation type is delayed and characterized by cell degeneration and necrosis [28]. The inflammation seen in DTH plays a critical protective role against intracellular pathogens. However, such inflammation is often seen against innocuous non-pathogenic antigens leading to severe damage to the tissue, as seen during contact dermatitis against specific allergens. The DTH was considered to be a Th1-driven inflammatory disease; however, treatment with antibodies against IFN-γ cytokine showed that such mice still continue to exhibit DTH signs [32]. In addition, the importance of Th17 cells in DTH disease progression was highlighted in IL-17 knockout mice, which showed significantly reduced ear swelling in an allergic contact dermatitis model [33]. Therefore, it is likely that both Th1 and Th17 cells play an effector role in DTH. A previous study demonstrated that vinegar or acetic acid could suppress the inflammatory response by inhibiting Th1 and Th17 in an animal experiment [34]. Thus, from previous data and our recent results, it is
suggested that NPV has immunomodulatory activity, which could be related to the inhibition of both the Th1 and Th17 pathways.

**Humoral immune response modulation**

To assess the effect of NPV on the humoral immune response in an animal model, rats were orally administered different concentrations and of NPV from different growing areas. The primary and secondary immune responses were evaluated by determining the antibody titre. Blood from the tail veins of rats was collected. It is interesting that low concentrations of NPV (1:10 dilution) showed significantly increased antibody titres for all 3 NPVs from different salinity planting areas when compared to the negative control. Moreover, anti-SRBC from the secondary response of most NPV-treated groups was significantly higher than the primary response, while the response was non-significant in the untreated negative control group. The results are shown in Table 1 and Figure 3. The results clearly suggest that NPV feeding can enhance the antibody response effects.

**Table 1** Anti-SRBC titre of rats fed daily with different concentrations of NPV from different salinity planting areas for the primary (day 7) and secondary (day 14) immune response. The data are the mean ± SEM.

<table>
<thead>
<tr>
<th>Rat groups</th>
<th>Anti-SRBC (mean ± SEM)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Primary response</td>
</tr>
<tr>
<td>Negative control</td>
<td>10.05±3.86</td>
</tr>
<tr>
<td>Positive control</td>
<td>20.42±1.91*</td>
</tr>
<tr>
<td>NPV-Fresh water 1:1</td>
<td>10.48±0.83</td>
</tr>
<tr>
<td>NPV-Fresh water 1:5</td>
<td>13.98±4.43</td>
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<tr>
<td>NPV-Freshwater 1:10</td>
<td>10.76±5.96</td>
</tr>
<tr>
<td>NPV-Brackish water 1:1</td>
<td>6.66±1.60</td>
</tr>
<tr>
<td>NPV-Brackish water 1:5</td>
<td>7.77±2.12</td>
</tr>
<tr>
<td>NPV-Brackish water 1:10</td>
<td>12.02±4.06</td>
</tr>
<tr>
<td>NPV-Saline water 1:1</td>
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</tr>
<tr>
<td>NPV-Saline water 1:5</td>
<td>10.48±0.83</td>
</tr>
<tr>
<td>NPV-Saline water 1:10</td>
<td>9.66±0.96</td>
</tr>
</tbody>
</table>

a; significantly different when compared to the negative group, b; significantly different when compared to the primary response.
Figure 3 Anti-SRBC titre on the primary (day 7) and secondary (day 14) immune responses of rats that were orally administered different concentrations of NPV from 3 different salinity planting areas on a daily basis.

Spleen index
Rats were treated with different NPV concentrations from different salinity planting areas for 30 days. Then, the rats were sacrificed, and the spleen was weighed. Both rat groups that were fed with NPV from fresh and brackish water planting areas at the high concentration (1:1 dilution) displayed significant increases in the spleen index when compared to the negative control. This means that the spleens in these
groups showed an enlarged appearance. However, the medium and low concentrations (1:5 and 1:10) displayed no significant difference in the spleen index when compared to the negative control. In summary, the results indicated that NPV in an appropriate consumption volume did not influence spleen size enlargement. The results are shown in Figure 4.

There are studies that have demonstrated the antibody production enhancement and lymphoid organ effect of vinegar intake in animal models. For example, in evaluating the effects of probiotics and vinegar as dietary supplements on the antibody production of broiler chickens, it was found that chickens had improved immune response against SRBC inoculation, especially in the primary response [35]. The vinegars were known as huge sources of probiotics that can influence microbial communities in the intestine and cease the growth of pathogens [36]. The results of our study were consistent with previous studies since NPV showed the enhancement of specific antibody production; however, there were significant increases, especially in the secondary response. Other evidence that supported vinegar enhancing antibody production is that it is composed of polyphenolics, and a flavonoid was reported to be a component in NPV [8] and play a role in immunoglobulmin stimulation or the immunomodulation effect [37]. Moreover, a previous study also determined the effect of probiotics and vinegar on the spleen size; however, there was no effect on the spleen size of animals [36]. The findings of previous studies are sustained by the present study.

Figure 4 Spleen index of rats that were administered different concentrations of NPV from different salinity planting areas for 30 days. The bars show the mean ± SEM. *, significantly different when compared to the negative control.
Especially in the Pak Phanang River Basin in the sub-district of Kanapnak, Pak Phanang district, Naknon Si Thammarat province, the southern east coast of Thailand, the nipa palm is an important economic field crop. Villagers cultivate the nipa palm as their primary source of income in producing various products including sugar, vinegar, whisky, roofing, and cigarette wrap paper. The salinity of planting areas ranges from fresh to brackish and saline water [38]. Salinity has been considered to be an important requirement for the occurrence of *Nypa fruticans* [39]. It grows well in very low salinity areas (below 5 ppt), as well as in freshwater outflow [32]. The previous studies showed that high salinity is a limiting factor for the growth of *Nypa fruticans*. With higher salinity, fewer new leaves are produced in juvenile and mature plants [40]. From these data, the salinity from different planting areas was the possible factor that affected the different active ingredients of NPV. The different active ingredients of NPV might reflect the immunomodulatory effect, as demonstrated in our results. However, the immunomodulatory effects of NPV from different salinity planting areas of southern Thailand showed the same trend, as clearly shown in the results.

**Conclusions**

Locally produced NPV form Nakhon Si Thammarat province, Thailand, exhibited enhancement of phagocytic activity *in vitro*, modulated cellular immune response of delay-type hypersensitivity inhibition (reduced edema) *in vivo* and modulated humoral immune response by enhanced antibody production *in vivo*. Moreover, it was affected to spleen with appropriated consumption. However, there was still a limited investigation of the mechanism of the *in vitro* and *in vivo* immunomodulatory effect of NPV vinegar with animal testing. Further, mechanistic and cytokine-associated mechanistic investigations are needed to uncover the underlying immunomodulatory activities of different NPV samples. This study concludes that NPV has the ability to modulate either the innate immune response (phagocytic activity) and adaptive immune response (cellular and humoral immunity), and the NPV benefits to human health can be investigated as well.

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**References**


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