FRESH SOY OIL PROTECTS AGAINST VASCULAR CHANGES IN AN ESTROGEN-DEFICIENT RAT MODEL: AN ELECTRON MICROSCOPY STUDY

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OBJECTIVE: To observe the effects of consuming repeatedly heated soy oil on the aortic tissues of estrogen-deficient rats.

METHODS: Thirty female Sprague Dawley rats (200-250 g) were divided equally into five groups. One group served as the normal control (NC) group. The four treated groups were ovariectomized and were fed as follows: 2% cholesterol diet (OVXC); 2% cholesterol diet + fresh soy oil (FSO); 2% cholesterol diet + once-heated soy oil (1HSO); and 2% cholesterol diet + five-times-heated soy oil (5HSO). After four months, the rats were sacrificed, and the aortic tissues were obtained for histological studies.

RESULTS: After four months of feeding, the NC, FSO and 1HSO groups had a lower body weight gain compared to the OVXC and 5HSO groups. The tunica intima/media ratio in the 5HSO group was significantly thicker (p < 0.05) compared to the NC, OVXC and FSO groups. Electron microscopy showed that endothelial cells were normally shaped in the FSO and NC groups but irregular in the 1HSO and 5HSO groups. A greater number of collagen fibers and vacuoles were observed in the 5HSO group compared to the other treatment groups.

CONCLUSIONS: Fresh soy oil offered protection in the estrogen-deficient state, as these rats had similar features to those of the NC group. The damage to the tunica intima and the increase in the ratio of tunica intima/media thickness showed the deleterious effect of consuming repeatedly heated soy oil in castrated female rats.

KEYWORDS: Soy oil; Heating; Aorta; Estrogen deficiency; Atherosclerosis.

INTRODUCTION

According to International Cardiovascular Disease Statistics by the World Health Organization (WHO), cardiovascular diseases are the leading cause of death globally. The WHO estimates that by 2015, almost 20 million people will die from cardiovascular diseases, which is projected to remain the single leading cause of death. Atherosclerosis is a cardiovascular disease that leads to morbidity and mortality in developed countries. The major risk factors of atherosclerosis are hyperlipidemia, hypertension, cigarette smoking and diabetes. It is also associated with increasing age, male gender, family history and genetic abnormalities.

One of the best ways to prevent atherosclerosis in the society is to alter dietary habits, including a reduced consumption of saturated animal fats. Vegetable oils, such as olive and soy, are recommended for human consumption due to their high content of monounsaturated fatty acids (MUFA) and polyunsaturated fatty acids (PUFA). Soy oil, which is produced from soybean (Glycine max), is the most widely marketed edible oil in the world. Soy oil contains approximately 60% of PUFA, 24% of MUFA and 16% of saturated fatty acids. This high level of PUFA dietary intake can improve the blood lipid profile status. In addition, with its high content of tocopherols, soy oil is known to exhibit various antioxidant actions against lipid peroxidation.

Edible oil, however, is often used in deep-frying,
which can lead to a series of chemical reactions involving hydrolysis, oxidation and polymerization. These changes may degrade the oil’s quality.\(^9\) The hydrolysis produces free fatty acids, which subsequently oxidizes and generates peroxide and hydroperoxide compounds, and secondary lipid oxidation products, such as aldehydes, ketones and alcohols.\(^9,10\) The consumption of these oxidation products has various adverse health effects, such as liver dysfunction, hypertension and renal damage.\(^11,12,13\)

We previously reported the adverse effects of consuming repeatedly heated palm oil by observing the ultrastructure of aortic tissues in estrogen-deficient rats.\(^14\) Estrogen-deficient rats were used to simulate an estrogen-deficient state observed postmenopausal women who are more susceptible to developing cardiovascular diseases, such as atherosclerosis.\(^15,16\) In the present study, we observed the effects of consuming repeatedly heated soy oil on the aorta of rats following 16 weeks of consumption.

**MATERIALS AND METHODS**

**Preparation of animal diets**

Commercial soy oil was purchased (Yee Lee Edible Oils, Malaysia) in three forms: fresh, heated once and heated five times.\(^11,17\) A total of 1 kg of ‘keropok lekor’ (fish-flavored chips) was deep fried using 2.5 L soy oil at 180°C for 10 minutes. The oil was then cooled to room temperature and stored in a tightly sealed container before being mixed with animal diets. For five-times-heated soy oil, the same heating procedure was performed. After each round, the oil was cooled for 5 hours before the heating process was repeated with a fresh batch of keropok lekor. No fresh soy oil was added between batches to replace any lost oil absorbed by the frying material.

The normal rat chow was comprised of 20% crude protein, 5% crude fiber, 2.5% crude fat, 13% moisture, 7% ash, 0.7-1.4% calcium, 0.6-1.2% total phosphorus and 51% of nitrogen-free extract. Other additives, such as vitamins A, D3, E, C, K and B12, thiamine, riboflavin, pantothenic acid, niacin, pyridoxine, folic acid, choline, antioxidants and trace minerals, were also present in the normal rat chow.

The experimental soy oil contained 14% saturated fatty acids, 26% MUFA and 60% PUFA. The cholesterol diet (2%) was purchased from MP Biomedicals, Inc. (Seven Hills, Australia). This cholesterol diet had the same composition as the normal rat chow with an addition of 2% cholesterol. The cholesterol diet was ground and mixed with the respective soy oil at a ratio of 85:15. The mixture was then formed into pellets and dried in an oven overnight at 80°C.

**Study design**

Thirty 2- to 4-month-old female Sprague Dawley rats weighing 200 to 250 g were used. Prior ethical approval was obtained from the animal unit of Universiti Kebangsaan Malaysia (UKMAEC: FAR/2003/Kamsiah/25 Jan/090). The rats were randomly divided into five groups: normal control (NC), ovariectomized control (OVXC), fresh soy oil (FSO), once-heated soy oil (1HSO), and five-times-heated soy oil (5HSO). All rats, except the NC group, were ovariectomized after 1 week of acclimatization. After recovery, each group was assigned with their respective diets, i.e., the NC group was fed with normal rat chow, the OVXC group was fed with the 2% cholesterol diet, the FSO group was fed with a 2% cholesterol diet fortified with fresh soy oil, the 1HSO group was fed with a 2% cholesterol diet fortified with once-heated soy oil and the 5HSO group was fed with a 2% cholesterol diet fortified with five-times-heated soy oil. The rationale for the number of heating was based on an earlier documented study in our laboratory.\(^14\) The rats were fed for 4 months before being sacrificed, after which their aortic tissues were taken for histological analyses. Their body weight and food intake were recorded every week during the study period.

**Histomorphometric study**

The proximal portion of the aortic tissues were removed and preserved in 10% formalin. They underwent standard tissue processing techniques, and aortic cross sections were stained with Verhoeff van Gieson for easy identification of the elastic fibers. The structure of the section was observed under light microscope (Eclipse 80i, Nikon Corporation, Tokyo, Japan), and images were taken with a camera (Qimaging MicroPublisher 5.0, Surrey, Canada). The images were analyzed using Image-Pro Plus version 5.0.2.9 (Media Cybernetics, Inc., Bethesda, USA), and the thickness of tunica intima (TI) and tunica media (TM) of the aorta were measured at 0°, 90°, 180° and 270° in each of the cross sections. Their average readings were taken for statistical analysis.

**Electron microscopic study**

The arch of the aortic tissues was cut into 0.5 mm rings, which were fixed with glutaraldehyde at 4°C for 12-16 hours. They were then washed with 0.1 M phosphate buffer, post-fixed with 1% buffered osmium tetroxide for 1-2 hours, and washed with distilled water. The samples were then bulk-stained with uranyl acetate for 30 minutes before being dehydrated in graded concentrations of ethanol, infiltrated in
propylene oxide and embedded in resin at 60°C for 24 hours. They were cut into 50 nm slices using an ultramicrotome. The ultrathin sections were stained with 3% uranyl acetate and Reynold’s lead citrate. The ultrastructure of the TI was viewed under a transmission electron microscope (Philips HMG 400, Philips, Eindhoven, The Netherlands).

**Statistical analysis**

All analyses were conducted using Statistical Product and Service Solutions (SPSS) (Chicago, IL, USA). Data are presented as the mean ± standard error of mean (SEM). Kolmogorov-Smirnov test was used to test the distribution of all data. Normally distributed data were analyzed using analysis of variance (ANOVA) followed by Tukey’s HSD post-hoc test. Data that were not normally distributed were analyzed using Kruskall-Wallis test. A $p$-value less than 0.05 was considered statistically significant.

**RESULTS**

**Average food intake and body weight gain (Table 1)**

<table>
<thead>
<tr>
<th>Group</th>
<th>Average food intake (g/week)</th>
<th>Body weight gain after 4 months (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NC</td>
<td>116 ± 2</td>
<td>79 ± 35</td>
</tr>
<tr>
<td>OVXC</td>
<td>110 ± 4</td>
<td>114 ± 15</td>
</tr>
<tr>
<td>FSO</td>
<td>78 ± 1 ±6</td>
<td>93 ± 42 ±6</td>
</tr>
<tr>
<td>IHSO</td>
<td>79 ± 2 ±6</td>
<td>91 ± 7 ±6</td>
</tr>
<tr>
<td>5HSO</td>
<td>86 ± 3 ±6</td>
<td>122 ± 32 ±2</td>
</tr>
</tbody>
</table>

Results are mean ± SEM (n = 6). a = significant compared to NC ($p < 0.05$), b = significant compared to OVXC ($p < 0.05$), c = significant compared to FSO ($p < 0.05$), d = significant compared to IHSO ($p < 0.05$), NC, normal control; OVXC, ovariectomized control; FSO, fresh soy oil-treated group; IHSO, once-heated soy oil-treated group; 5HSO, five-times-heated soy oil-treated group.

Rats treated with soy oil (i.e., the FSO, IHSO and 5HSO groups) showed a reduction of average food intake compared to the NC and OVXC groups. Interestingly, the food intake was the highest in the NC group. After 16 weeks of feeding, the NC, FSO and IHSO groups had lower body weight gain compared to the OVXC and 5HSO groups ($p < 0.05$).

**Histomorphometric study**

Figure 1 shows the average intimal thickness and medial thickness of each group. There was no significant difference in the intimal thickness among the groups. Conversely, all ovariectomized groups showed significant differences in the medial thickness compared to the NC group. The 5HSO group also showed a significantly smaller medial thickness compared to the FSO group. Additionally, the 5HSO group showed the highest TI/TM ratio compared to the NC, OVXC and FSO groups (Figure 2).

**Electron microscopic findings**

*Rat aorta from the NC group (Fig. 3A)*

This tissue showed the normal endothelial cell shape. Endothelial cells were located near the internal elastic lamina.

*Rat aorta from the OVXC group (Fig. 3B)*

This tissue contained endothelial cells whose shapes were irregular. The endothelial cells were distant from the internal elastic lamina.
**Rat aorta from the FSO group (Fig. 3C)**

The characteristic features of the endothelial cells were very similar to the NC group.

**Rat aorta from the 1HSO group (Fig. 3D)**

Endothelial cells were irregularly shaped. The subendothelial layer was thicker. Abundant collagen fibers were observed in the subendothelial layer, and some endothelial cells contained vacuoles.

**Rat aorta from the 5HSO group (Fig. 3E)**

Endothelial cells were irregularly in shape. The subendothelial layer was thicker. Abundant collagen fibers were observed in the subendothelial layer. More vacuoles were observed in the subendothelial layer, and some endothelial cells contained vacuoles.

*Figure 3 - A) A normal control rat aorta. B) An ovariectomized control rat aorta. C) The aorta of a rat fed with fresh soy oil. D) The aorta of a rat fed with once-heated soy oil. E) The aorta of a rat fed with five-times-heated soy oil. Note the collagen fibrils (c) in the subendothelial layer (in B) and vacuolization (v) of the endothelial cells (in D and E). L, lumen; TI, tunica intima; IEL, internal elastic lamina; TM, tunica media; EC, endothelial cell (7,700x)*
were observed in the endothelial cells, and the endothelial cells were further away from the internal elastic lamina compared to the 1HSO group.

**DISCUSSION**

Here we observed ultrastructural changes in the aorta of estrogen-deficient rats after the consumption of heated soy oil. We have previously shown that ingestion of repeatedly heated edible oils resulted in adverse effects on serum lipid peroxidation and lipid profiles. These changes may be due to the degradation products that are produced after the oils are heated at high temperatures. Staprans et al. showed that oxidized lipids in the diet were absorbed in the gastrointestinal tract and were transported into circulation. The ingestion of oxidized lipids contributes to the development of atherosclerosis, as oxidized lipoproteins promotes the formation of foam cells, which are the predominant cells in the initial lesion of atherosclerosis.

The addition of soy oil, either in fresh or heated forms, reduced the average food intake in experimental rats compared to the control rats. This reduction shows that the rats preferred normal rat chow or the 2% cholesterol diet without soy oil. We assume that the addition of oil, which modified the taste and smell of the diet, may have contributed to the reduction of their food intake. The heating process, however, did not have any significant effect on their food intake. All soy oil-treated groups showed a similar food intake, regardless of the number of times the oil was heated. The OVXC group showed a significant body weight gain after 4 months of feeding compared to the NC group. This increase may be due to the 2% cholesterol diet, which is known to contribute to significant weight gain. Despite their low food intake, the 5HSO group exhibited the highest weight gain compared to the NC, FSO and 1HSO groups. We assume that the weight gain in the 5HSO group is caused by water retention. An earlier study reported that water retention may be due to multiple causes, such as renal failure or hypertension, which affected weight gain. In the present study, however, we did not look into the renal status or hypertension.

The histomorphometric study showed that feeding with soy oil did not result in any significant differences in TI thickness among the soy oil-treated groups. All the ovariecotimized rats, however, showed significant medial thinning compared to the NC group. The reason for this change is not clear. Aging and estrogen deficiency may contribute to such changes. Because ovariecotimized rats simulate a postmenopausal condition, it is likely that the observed medial thinning was due to muscular and connective tissue loss secondary to the estrogen deficiency. Feeding with repeatedly heated soy oil resulted in a significant reduction of medial thickness compared to the FSO group, suggesting that fresh soy oil may have a protective effect in the muscle growth, but such protective effects are lost when the oil is repeatedly heated. Although the TI layer did not show any significant difference, the reduction of the TM layer in ovariecotimized rats contributed to the significantly higher TI/TM ratio in the 5HSO group compared to the NC, OVXC and FSO groups. This higher TI/TM ratio indicated an early development in the process of atherosclerosis.

The aortic arch was chosen for the electron microscopic study because this site is most susceptible to atherosclerosis due to any hemodynamic changes that occurred in the vessel. The NC group showed normal endothelial cells that were elongated or elliptical in shape. Conversely, the OVXC group showed an increase of subendothelial space containing collagen fibrils because the endothelial cell had migrated towards the lumen. The endothelial cell in this group was desquamated from the internal elastic lamina. The observed changes may result in a decrease in vasodilator response and, at the same time, may expose smooth muscle cells to vasoconstrictors, causing artery constriction. We postulate that these changes may be due to the deficiency of estrogen in the ovariecotimized rats. Rats treated with fresh soy oil showed no obvious ultrastructural changes. The endothelial cell was similar to those observed in the NC group and shared tight junctions with the internal elastic lamina, suggesting that feeding fresh soy oil to estrogen-deficient rats had protective benefits against any damage caused by estrogen deficiency coupled with a high cholesterol diet. The high level of antioxidants, primarily tocopherols, in soy oil may be responsible for the protective effects of soy oil on the aortic ultrastructure observed in this study.

The 1HSO and 5HSO groups showed some abnormal ultrastructural changes compared to normal rats. An increase in the subendothelial layer could be observed in both groups with the presence of abundant collagen fibrils in the area. A large vacuole was observed in the endothelial cell of the 1HSO group because the cell had moved towards the lumen. The endothelial cell of the 5HSO group had also moved further from the internal elastic lamina, and many vacuoles were observed in the endothelial cells. The vacuolization and condensation of cytoplasm in the endothelial cells indicated an early stage of apoptosis. Necrosis or apoptosis is known to occur due to an excess of oxidized low-density lipoprotein in macrophages during the progression of atherosclerosis. Based on these findings, soy oil, which showed beneficial effects in the FSO group, had lost its potential when heated. The adverse effects of the heated soy oil became more apparent after the oil was heated repeatedly.
We postulate that the loss of the beneficial effects of soy oil was likely due to the significant reduction in tocopherols because we previously reported that more than 80% of tocopherol is lost after of the oil was heated five times. Repeated heating not only reduces antioxidant contents but also generates free radicals. Soy oil, which contains a high amount of PUFA, is unstable at high temperatures and susceptible to lipid peroxidation. Lipid peroxidation produces volatile and non-volatile compounds. The non-volatile compounds are more important because they remain in the oil and are absorbed by the food. These compounds affect the flavor stability and texture of the fried foods that are later consumed. Heating the same oil repeatedly generates more degradation products, which may promote progression of numerous diseases, such as atherosclerosis, when this oil is consumed. These statements have been already shown in an earlier study.

CONCLUSION

Vegetable oil has some protective effects against lipid peroxidation; however, repeated heating may cause deterioration in its quality and formation of new chemical components that are harmful. The practice of using the same oil repeatedly for cooking to save on the cost should therefore be avoided. We conclude that the consumption of fresh soy oil offers protection against vascular changes in an estrogen-deficient state; however, the protective effect is lost when the oil is heated repeatedly. The consumption of repeatedly heated soy oil resulted in alterations in endothelial cells, an increased in the width of the subendothelial layer, and changes in the TI/TM ratio. These findings are indicators of early atherosclerosis. Hence, we do not advise estrogen-deficient subjects to consume repeatedly heated edible oil.

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