Accerelative Effect of Olive Oil on Adrenal Corticosterone Secretion in Rats Loaded with Single or Repetitive Immersion-Restraint Stress

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Summary The present study was performed to clarify the effects of dietary oils on physiological and metabolic changes induced by a stress, using one-time or repetitive water-immersion of restrained rats (single or repetitive stress) as an experimental stress load. In rats fed any test diets containing 20% of the mixture of tripalmitin, tristearin, and corn oil (PSC), olive oil (OLI), safflower oil (SAF), and linseed oil (LIS) with repetitive stress loading, body weight gains and food intakes were generally reduced. The weights of the thymus and spleen also declined, but the adrenal weights were enhanced. Particularly, the increase in the adrenal weight of rats given the OLI diet was greater than of rats supplied with other diets. When the rats were loaded with the single or repetitive stress, the concentrations of urea, lipid peroxide, and corticosterone in the plasma were increased in rats fed any of dietary oils. The rise of plasma corticosterone level was especially great in rats fed the OLI diet. The concentrations of total cholesterol (T-CHOL) and triglyceride (TG) in the plasma and liver generally tended to be higher in rats fed the OLI diet than in rats given the other diets with and without stress exposure. Plasma corticosterone concentration was correlated to the adrenal weight (r=0.87, p<0.05). This study showed that OLI especially enhanced the adrenal weight in rats exposed to the repetitive stress and further raised the increased secretion of adrenal corticosterone in rats loaded with the single or repetitive stress compared with the other oils. The mechanism explaining these actions of OLI was inferred to be related to the levels of T-CHOL and TG in the plasma and liver generally enhanced by stress.

Key Words olive oil, adrenal corticosterone, water immersion-restraint stress, single and repetitive stresses, total cholesterol

The appearance of physiological and metabolic changes (1–9) showing a decrease in body weight, a shrink of the spleen or thymus, an increase of adrenal weight, and so on (10, 11) are reported as the responses of a living body on acute (single) or chronic (repetitive) stress (5). A stress load to rats enhances the secretion of corticosterone, and this hormone is used as an index of stress in many studies (4, 5, 7). An incorporation of cholesterol (CHOL) into the adrenal is highest, however, in the various organs (12), and oleic acid richly contained in olive oil (OLI) is reported to be best as a substrate of acyl-CoA: cholesterol acyltransferase in the liver (13), which practically exists in all tissues. The feeding of OLI to rats enhances the concentration of plasma total CHOL in the previous reports (14–16). It is considered that in rats loaded with stress, the feeding of OLI increases plasma CHOL level, and as a result the incorporation of CHOL into the adrenal is elevated, by which the secretion of adrenal corticosterone is enhanced. Few investigations, however, are involved in the relationships between stress loading and various oils. The present study was conducted to compare the effects of OLI on stress-induced changes with other oils in experimentally stress-loaded rats, adopting the water immersion of rats individually restrained in cages (17), which is similar to a cold restraint normally regarded as a primary physical-stress (5, 7). The present stress load was divided into single and repetitive stresses, as the responses of a living body on these stresses were conjectured to be different between both ones (5).

MATERIALS AND METHODS

Materials. Casein, OLI, safflower oil (SAF), linseed oil (LIS), tripalmitin, and tristearin (Nacalai Tesque, Kyoto), sugar (Fuji Seito, Co. Ltd., Shimizu), and mineral and vitamin mixtures (AIN-76MA) (18) (Nihon Nosen Kogyo, Ltd., Yokohama) were purchased from respective companies. Corn starch (α) and corn oil were presented by Fuji Seifun, Ltd., and Honen Corporation (Shimizu), respectively.

Animals and diets. Male rats of the Wistar strain (Japan SLC, Hamamatsu), about 150 g and 6 wk of age, were individually housed in suspended stainless-steel wire cages and fed in an air-conditioned room with a temperature of 23±1°C, a humidity of 50±3%, and a 12 h light (06:00–18:00) and dark cycle. They had free access to food and water. The experimental plan was approved by the Experimental Animal Care Committee of the Faculty of Agriculture, Shizuoka University. The

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rats were fed a basal diet prepared on the dietary recommendations of the American Institute of Nutrition (18), except for the variation in dietary protein level, to accustom themselves to the surroundings for 4 to 5 d. The basal diet contained (in weight %) casein (25.0), mineral mixture (3.5), vitamin mixture (1.0), choline chloride (0.2), corn oil (10.0), sucrose (46.2), and α-starch (14.1), which was added 0.1 mL of 20% BHT in alcohol per 100 g of the diet.

In the experiment (Exp.) 1 (single stress), rats having about 170 g of body weight were divided into four groups of 12 rats each, which received any of the test diets containing 20% of the mixture of tripalmitin, tristearin, and corn oil (PSC), OLI, SAF, and LIS (PSC, OLI, SAF, and LIS) for 10 d. The PSC was semisynthesized by mixing 64% of tripalmitin, 16% of tristearin, and 20% of corn oil. The test diets were prepared by increasing the content of oils at the expense of sucrose and α-starch in the basal diet, keeping the constant rate (46.2/14.1) between them. The fatty acid compositions of dietary oils are shown in Table 1. At the end of the feeding period, all groups were deprived of food for 5 h, and the stress groups were loaded with the final stress for the latter 2 h during the term of food deprivation. The other treatments were performed as described in Exp. 1.

| Fatty acid | PSC | OLI | SAF | LIS |
|-----------|-----|-----|-----|-----|
| 16:0      | 66.2| 8.1 | 6.9 | 5.6 |
| 18:0      | 16.4| 2.4 | 2.5 | 3.4 |
| 18:1n-9   | 6.9 | 75.1| 12.9| 19.3|
| 18:3n-6   | 10.0| 12.7| 76.6| 15.8|
| 18:3n-3   | 0.3 | 0.6 | 0.3 | 55.5|
| Others    | 0.2 | 1.1 | 0.8 | 0.4 |

1 PSC, the mixture of tripalmitin (64%), tristearin (16%), and corn oil (20%); OLI, olive oil; SAF, safflower oil; LIS, linseed oil.

All samples were stocked at −20°C until analyzed.

**Analytical procedure.** The fatty acid compositions of oils used were determined by the previous report (16). The concentrations of total (T)-CHOL, triglyceride (TG), and urea in the plasma were determined by using enzymatic kits (Wako Pure Chemical Industries, Osaka). The levels of plasma corticosterone and lipid peroxide were estimated by a fluorescent method (19) and Yagi’s method (20), respectively. The amounts of liver T-CHOL and TG were measured by the methods of Zak-Henley (21) and Fletcher, respectively, after the extraction of liguid with the mixture of chloroform–methanol (2:1, v/v) (23) from the freeze-dried liver.

**Statistical analysis.** Data were presented as the mean±SE for 6 rats of each group and subjected to a one-way variance (ANOVA), followed by the inspection of all differences among means by using Duncan’s multiple range test (24) at p<0.05. The correlation coefficient (r) was determined by the method of least squares.

**RESULTS**

**Body weight gain, food intake, and organ weight**

The changes in final body weight, body weight gain, food intake, and liver weight of rats exposed to the single stress or not after the feeding of test diets for 10 d (Exp. 1) are shown in Table 2. The final body weights were generally different among the groups. The weight gains were higher in stress-unexposed (normal) SAF and LIS groups than in the other groups. The food intakes were not significantly different among dietary groups. The liver weights were hardly different among all groups except the normal LIS group, of which liver weights tended to be increased compared with the other groups. The weights of the spleen, thymus, and adrenal were unchanged among dietary groups, though the data were not shown.

The final body weights, weight gains, food intakes, and organ weights in rats loaded with the repetitive stress or not during the feed of test diets (Exp. 2) are presented in Table 2. The final body weights were generally smaller in normal groups than in stressed ones. The difference in weight gain among normal groups was not observed except in the PSC group, in which weight gain was lower than in the other groups, but stress loading reduced the weight gains of all groups. The change in food intake was nearly in proportion to weight gain in the groups fed any of test diets. Liver weights were not different among the groups. The weights of the thymus and spleen were hardly changed by types of oil, but they were reduced by stress-loading. The adrenal weights were generally smaller in stressed groups except the PSC group; especially the stress-exposed OLI group had the greatest adrenal weight.

| T-CHOL, TG, urea, lipid peroxide, and corticosterone in the plasma and T-CHOL and TG in the liver |
|---------------------------------------------------------------|
| The concentrations of T-CHOL, TG, urea, lipid peroxide and corticosterone in the plasma, and T-CHOL and TG in the liver of rats loaded or not loaded with the single or repetitive stress (Exp. 1) are presented in Table 3 and Fig. 1. As shown in Table 3, the concentrations of |
Table 2. Body weight, food intake, and organ weight of rats exposed to water immersion restraint stress once after being fed test diets for 10 d (Exp. 1) and repetitively every other day (4 times) during feeding test diets for 7 d (Exp. 2).

| Dietary group | PSC | OLI | SAF | LIS |
|---------------|-----|-----|-----|-----|
|                | No stress | Stress | No stress | Stress | No stress | Stress | No stress | Stress |
| Final body weight (g) | 187.5 ± 2.8<sup>a</sup> | 189.5 ± 0.7<sup>a</sup> | 192.5 ± 3.3<sup>a</sup> | 193.3 ± 4.3<sup>a</sup> | 195.1 ± 5.8<sup>a</sup> | 194.5 ± 3.5<sup>a</sup> | 199.2 ± 3.6<sup>a</sup> | 189.9 ± 2.8<sup>a</sup> |
| Body weight gain (g/10 d) | 44.9 ± 1.7<sup>b</sup> | 45.6 ± 0.9<sup>b</sup> | 46.3 ± 3.5<sup>b</sup> | 47.0 ± 2.9<sup>b</sup> | 55.5 ± 1.3<sup>b</sup> | 47.6 ± 1.5<sup>b</sup> | 54.6 ± 3.1<sup>b</sup> | 46.6 ± 2.7<sup>b</sup> |
| Food intake (g/10 d) | 120.6 ± 1.3<sup>bc</sup> | 120.9 ± 1.0<sup>bc</sup> | 115.2 ± 3.3<sup>bc</sup> | 117.5 ± 4.5<sup>bc</sup> | 121.3 ± 3.0<sup>bc</sup> | 119.9 ± 1.7<sup>bc</sup> | 121.9 ± 3.2<sup>bc</sup> | 115.5 ± 1.9<sup>bc</sup> |
| Liver weight (g/100 g body weight) | 4.19 ± 0.09<sup>ab</sup> | 4.21 ± 0.15<sup>ab</sup> | 4.04 ± 0.11<sup>b</sup> | 3.92 ± 0.05<sup>b</sup> | 3.96 ± 0.09<sup>b</sup> | 3.72 ± 0.08<sup>b</sup> | 4.45 ± 0.07<sup>b</sup> | 4.11 ± 0.13<sup>b</sup> |

1. PSC, the mix of tripalmitin (64%), tristearin (16%), and corn oil (20%); OLI, olive oil; SAF, safflower oil; LIS, linseed oil.
2. Mean ± SE for 6 rats. Values in a row not sharing the same superscript letter are significantly different at p<0.05 by Duncan’s multiple range test.
Table 3. The concentrations of plasma total cholesterol (T-CHOL), triacylglycerol (TG), urea, and lipid peroxide, and liver T-CHOL and TG of rats exposed to water immersion restraint stress once after being fed test diets for 10 d (Exp. 1) and repetitively every other day (4 times) during feeding test diets for 7 d (Exp. 2).

| Dietary group | PSC | OLI | SAF | LIS |
|--------------|-----|-----|-----|-----|
|              | No stress | Stress | No stress | Stress | No stress | Stress |
| Concentration of plasma lipids (mmol/L), urea (μmol/L), and lipid peroxide (μmol/L) | | | | | |
| T-CHOL | 2.34±0.18<sup>de</sup> | 2.88±0.13<sup>a</sup> | 2.49±0.09<sup>bc</sup> | 3.00±0.13<sup>a</sup> | 2.06±0.09<sup>bc</sup> | 2.78±0.11<sup>ab</sup> |
| TG | 2.34±0.14<sup>b</sup> | 5.24±0.51<sup>a</sup> | 1.63±0.08<sup>b</sup> | 4.73±0.33<sup>a</sup> | 2.06±0.12<sup>b</sup> | 4.82±0.28<sup>a</sup> |
| Lipid peroxide | 3.81±0.41<sup>ed</sup> | 4.05±0.29<sup>cd</sup> | 3.01±0.23<sup>d</sup> | 3.23±0.38<sup>cd</sup> | 4.18±0.47<sup>d</sup> | 13.3±0.3<sup>b</sup> |
| Concentration of liver lipids (μmol/g wet) | | | | | |
| T-CHOL | 9.47±0.23<sup>d</sup> | 9.83±0.60<sup>d</sup> | 14.3±0.3<sup>a</sup> | 14.5±0.8<sup>a</sup> | 13.5±0.7<sup>ab</sup> | 13.1±0.3<sup>abc</sup> |
| TG | 20.2±1.0<sup>d</sup> | 21.6±4.7<sup>c</sup> | 33.4±1.4<sup>b</sup> | 44.9±4.3<sup>a</sup> | 31.0±2.2<sup>bc</sup> | 34.3±6.0<sup>b</sup> |

Exp. 1

| Concentration of plasma lipids (mmol/L), urea (μmol/L), and lipid peroxide (μmol/L) | | | | | |
| T-CHOL | 2.02±0.13<sup>cd</sup> | 2.12±0.08<sup>bcd</sup> | 2.56±0.25<sup>ab</sup> | 2.64±0.25<sup>a</sup> | 2.33±0.12<sup>abc</sup> | 2.19±0.15<sup>bcd</sup> |
| TG | 3.27±0.19<sup>a</sup> | 2.61±0.35<sup>ab</sup> | 3.10±0.52<sup>a</sup> | 2.38±0.24<sup>abc</sup> | 1.88±0.07<sup>bc</sup> | 1.81±0.16<sup>bcd</sup> |
| Urea | 2.41±0.08<sup>b</sup> | 3.58±0.33<sup>a</sup> | 2.73±0.26<sup>a</sup> | 3.55±0.33<sup>a</sup> | 2.43±0.12<sup>bc</sup> | 3.48±0.22<sup>a</sup> |
| Lipid peroxide | 2.88±0.54<sup>bc</sup> | 2.14±0.17<sup>c</sup> | 3.09±0.45<sup>bc</sup> | 2.13±0.25<sup>c</sup> | 3.56±0.39<sup>b</sup> | 5.88±0.34<sup>a</sup> |
| Concentration of liver lipids (μmol/g wet) | | | | | |
| T-CHOL | 15.5±0.7<sup>bc</sup> | 14.5±1.5<sup>c</sup> | 18.5±0.5<sup>ab</sup> | 20.0±1.2<sup>a</sup> | 15.2±1.7<sup>bc</sup> | 14.6±0.9<sup>c</sup> |
| TG | 20.4±3.4<sup>bc</sup> | 19.9±2.0<sup>c</sup> | 27.5±3.4<sup>b</sup> | 35.9±3.9<sup>a</sup> | 17.5±2.0<sup>c</sup> | 14.5±1.4<sup>c</sup> |

Exp. 2

1 See Table 2.
2 Mean±SE for 6 rats. Values in a row not sharing the same superscript letter are significantly different at p<0.05 by Duncan’s multiple range test.
plasma T-CHOL were generally higher in normal PSC and OLI groups than in the normal SAF and LIS groups, and they were significantly increased by stress exposure in all groups except the LIS group. Plasma TG concentrations were hardly different among normal groups, but were enhanced by stress loading in each group, among which difference was scarcely observed. Plasma urea levels were unchanged by dietary oils, but they were extremely increased by stress in all groups, among which there was no difference in the urea levels. The concentrations of lipid peroxide were highest in the stress-loaded LIS group, next high in the stressed SAF and normal LIS groups, and lowest in the other groups. Plasma corticosterone levels (Fig. 1) were low in normal groups, among which there was no difference in the levels, but they were enhanced by stress load in each group. The rise of this hormone level by stress were especially greater in the OLI group than in the other groups.

The contents of liver T-CHOL and TG had an inclination to be higher in the OLI group than in the other groups with and without stress exposure.

The levels of T-CHOL, TG, urea, lipid peroxide, and corticosterone in the plasma and T-CHOL and TG in the liver of rats exposed or unexposed to the repetitive stress during feeding test diets (Exp. 2) are shown in Table 3 and Fig. 2. The concentrations of plasma T-CHOL were inclined to be highest with feeding of the OLI diet, followed in order by the PSC and OLI diets and lowest in the LIS diet. No difference was found in the concentrations between the load and nonload of stress. Plasma TG levels were generally higher in normal rats fed the PSC and OLI diets than in the other diets, but they were inclined to be reduced by stress exposure in both groups. Urea concentrations in the plasma were hardly different among each normal group, and they similarly increased with stress exposure in any group except LIS, in which the urea level was not increased by stress. The contents of plasma lipid peroxide were scarcely different among normal groups, but they extremely rose in the SAF and LIS groups under stress treatment. The livers generally contained the higher amounts of T-CHOL and TG in rats given the OLI diet than in rats fed the other diets under normal and stress load. No significant difference was noted in plasma corticosterone levels among normal groups, but stress loading extremely raised the levels in any of the groups, and enhancement in the levels was especially great in the stressed OLI group (Fig. 2).

Correlation between corticosterone and adrenal weight

As shown in Fig. 3, a positive correlation was found between plasma corticosterone concentrations and the adrenal weights ($r=0.87, p<0.05$).
tion showed that OLI gave the greatest increase in adrenal weight besides the adaptability to stress as described below, since the secretion of adrenal corticosterone was lower under the repetitive stress than under the single one (Figs. 1 and 2).

The T-CHOL and TG levels in the plasma and liver (Table 3) generally tended to be higher with the feeding of the OLI diet than with the feeding of the other diets in the normal and stressed rats of both experiments, which was similar to the earlier reports (14–16, 29). This result supports OLI having an effect to enhance the contents of T-CHOL and TG in the plasma and liver and suggests that these enhancements by the feeding of OLI are related to a further increase in the raised concentration of plasma corticosterone as again described below.

DISCUSSION

The water immersion of restrained rats in this study is normally regarded as primary physical stress (5, 7), and similar stress has been used to induce ulcers in the digestive organs of rats (17, 25). The ulcers were not visibly found in Exps. 1 and 2, however. This observation suggests that the degree of the present stresses was relatively modest. In both experiments (Table 2), the body weight gains of normal rats fed the PSC diet tended to be reduced compared with those of normal rats fed the other diets, though there was no difference in food intake between the PSC group and the other groups. This decrease in weight gain would be due to stearic acid enriched in the PSC diet, since stearic acid is difficult to be absorbed in the intestine and as a result bring about a smaller energy consumption than in the other groups (26, 27). The greater weight gains of normal rats in Exp. 1 than in Exp. 2 would be because of the longer period of feeding test diets in Exp. 1 compared with Exp. 2. The repetitive stress load caused the decline in food intakes, which was similar to the results of the previous studies (11, 28). The weights of the spleen and thymus in Exp. 2 (Table 2) were decreased by loading with stress in rats given any of the diets, among which there was no difference. This result was identical to the earlier studies (10, 11). Adrenal weights (Table 2) were inversely increased by stress, which was also reported previously (10), but the degree of weight increase was different among dietary oils. This investigation showed that OLI gave the greatest increase in adrenal weight in the repetitive stress-loaded rats. The increase in adrenal weight was also correlative to the enhancement of plasma corticosterone level by the repetitive stress (Fig. 3), but the increase in adrenal weight was not observed by the single stress (Exp. 1). This difference between single and repetitive stresses may show that the raise in plasma corticosterone concentration by stress loading was not essentially related to the increase in adrenal weight, but the repetitive stress with feeding of the diets might have a bad influence on the adrenal weight besides the adaptability to stress as described below, since the secretion of adrenal corticosterone was lower under the repetitive stress than under the single one (Figs. 1 and 2).

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nosis, which is involved in TG synthesis in the liver. A few of the metabolic changes in this study, plasma urea concentrations, lipid peroxide contents in the plasma of rats fed the SAF and LIS diets, and the secretion of adrenal corticosterone, were greater with the single stress treatment than with the repetitive one, which resembled in the previous report (30). This difference might be attributed to the adaptability to stress load.

In conclusion, the OLI rich in oleic acid especially increases the adrenal weight in rats loaded with the repetitive water immersion restrained stress and enhances the secretion of adrenal corticosterone in rats exposed to the single and repetitive ones compared with the other oils. The mechanism explaining the actions of OLI was inferred to be related to the raised levels of T-CHOL and TG in the plasma and liver by the feeding of OLI.

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